Typing of *Neisseria gonorrhoeae* isolates by phenotypic and genotypic techniques in New Delhi, India

Seema Sood, Neeraj Mahajan, Rajendra Singh, Sonu Kumari Agrawal, Trupti Shende, Arti Kapil, Hemanta K. Kar¹, Vinod K. Sharma²

**Abstract:**

**BACKGROUND:** The objective of this study is to investigate gonococcal isolates using phenotypic and genotypic methods.

**METHODOLOGY:** Sixty gonococcal isolates obtained were examined. Strains were divided into 9 resistant phenotypes: Chromosomally mediated penicillin-resistant *Neisseria gonorrhoeae* (CMRNGP), penicillinase-producing NG (PPNG), chromosomally mediated tetracycline-resistant NG (CMRNGT), TRNG, PPNG and TRNG, CMRNGPT, quinolone resistant NG (QRNG), Azithro R, and decreased susceptibility (DS) to ceftriaxone. These isolates were also subjected to auxotyping and NG-multi-antigen sequence typing (MAST).

**RESULTS:** Of 60 isolates, 32 (53.33%) PPNG and only one was CMRNGP; 16 (26.66%) were CMRNGT, while 18 (30%) were TRNG. Both PPNG and TRNG found in 13 (21.66%) and none were CMRNGPT. QRNG was seen in 93.33%, 5% Azithromycin R, and 6.66% were DS to ceftriaxone. Based on auxotyping, 24 (40%) nonrequiring, 16 (26.66%) were proline requiring, 13 (21.66%) arginine requiring while 7 (11.66%) belonged to others. The most common ST was 6058 (32.5%).

The discriminatory indices of antibiogram, auxotyping and NG-MAST were 0.77, 0.72, and 0.95, respectively.

**CONCLUSIONS:** NG-MAST is the method of choice for epidemiological studies.

**Key words:**

Antibiogram, auxotyping, *Neisseria gonorrhoeae* multiantigen sequence typing, *Neisseria gonorrhoeae*

**Introduction**

Gonorrhea, caused by *Neisseria gonorrhoeae* (NG), is a common sexually transmitted infection in both developing and developed countries.¹ These infections are of concern to healthcare workers as they carry increased risk of morbidity particularly among women as well as neonate.² Epidemiological surveillance and strain characterization can provide information regarding the transmission dynamics of this infection, which further influence public health interventions.

Over the past 40 years, various typing schemes have been devised to characterize the isolates of NG. These have been used to identify the population of strains that circulate in different communities, temporal and geographic changes among the strains, transmission patterns and antibiotic resistance outbreaks.³,⁴ Phenotyping and genotyping have been accepted as the two major systems for typing of gonococci. Phenotypic techniques, such as antibiogram, auxotyping and serotyping, have been used in the past either alone or in combination for epidemiological studies and to determine relatedness in NG isolates. However, with new developments in genetics, these are...
now being replaced by molecular techniques. Typing methods can provide information regarding relationship between specific types of gonococci with clinical, diagnostic, and therapeutic problems.[5] In a developing country like India, only few studies have been carried out on typing of NG, and most have focused on phenotypic methods only.[6,7] Therefore, the aim of the present investigation was to analyze the gonococcal isolates in New Delhi by not only phenotypic (antibiogram and auxotyping) methods but also by genotypic methods (NG-multiantigen sequence typing [MAST]). Further, we also examined the association between the typing methods and antimicrobial resistance (AMR). In addition, the ability of different typing methods to differentiate between two isolates has been calculated using the discriminatory index.

**Methodology**

**Neisseria gonorrhoeae isolates**
A total of 60 consecutive NG isolates collected from patients presenting with cervicitis and urethritis to Dermatology OPD of AIIMS and PGIMER and Dr. R.M.L. Hospital, New Delhi, from July 2010 to June 2012 were studied. Out of 60 isolates, 59 were from urethral swabs of male patients and one from endocervical swab of a female patient. All clinical isolates were preserved in 20% glycerol broth by deep freezing (−86°C ultra-low freezers Freezer, Model 991; Thermo Scientific) at −70°C and also lyophilized using lyophilizer (Free Zone Triad Freeze dry system: Model - 7400030 from Labconco). All 60 isolates were revived on chocolate agar plates at 36°C in an atmosphere containing 5% CO₂ before typing.

**Antimicrobial susceptibility testing**
Minimum inhibitory concentrations (MICs) of penicillin, tetracycline, ciprofloxacin, spectinomycin, ceftriaxone and azithromycin for all isolates (n = 60) was determined by E test method (Biomerieux, France) as per manufacturer’s instructions. The results were interpreted as per breakpoint criteria of calibrated dichotomous sensitivity technique.[8] In addition, all isolates were tested for β-lactamase production by chromogenic cephalosporin method using nitrocefin discs (BD, USA). The WHO NG reference strains F, G, K-P were used for quality control.

The isolates were divided into nine-resistant phenotypes: Chromosomally mediated penicillin-resistant NG (CMRNG[5] (CMRNG[i.e., non-penicillinase producing NG [PPNG] with penicillin MIC ≥2.0 μg/ml and tetracycline MIC <2.0 μg/ml], PPNG (PPNG, i.e., β-lactamase positive with tetracycline MIC <16 μg/ml)], chromosomally mediated tetracycline resistant NG (CMRNG[7]) (CMRNG[i.e., non-PPNG with penicillin MIC <2.0 μg/ml and tetracycline MIC 2.0–8.0 μg/ml], TRNG (plasmid-mediated tetracycline-resistant NG, i.e., β-lactamase negative with tetracycline MIC ≥16 μg/ml), PPNG and TRNG, i.e., β-lactamase positive with tetracycline MIC ≥16 μg/ml, CMRNG[^7], i.e., non-PPNG with penicillin MIC ≥2.0 μg/ml and tetracycline MIC 2.0–8.0 μg/ml, quinolone-resistant NG (QRNG) (QRNG, i.e., MIC ≥1 μg/ml), azithromycin resistant (MIC ≥1 μg/ml), and isolates with decreased susceptibility (DS) to ceftriaxone (MIC 0.03–0.25 μg/ml).

**Auxotyping**
Auxotyping was performed as per protocol using gonococcal genetic medium (GGM).[9] Four different types of media were prepared – one without proline, second without arginine, third containing all seven amino acids, and fourth omitting all 7 amino acids. Standard inoculum size of approximately 3 × 10⁵ CFU/ml was used. The medium deficient in substrates was inoculated first, i.e., media lacking all seven amino acids/proline/arginine were inoculated first followed by complete GGM, and finally, the control chocolate agar plates supplemented with yeast extract. The culture plates were then incubated at 37°C in 5%–10% CO₂ and were examined after 24 and 48 h of incubation. Isolates with no special nutritional requirement were classified as non-requiring (NR) and those showing absence of growth in proline and arginine deficit media were classified as proline requiring (PR) and arginine requiring (AR), respectively. The WHO NG reference strains F (AR), K (PR) were used for quality control.

**Neisseria gonorrhoeae multiantigen sequence typing**
**Isolation of genomic DNA**
Genomic DNA was isolated by boiling method and stored at −20°C.

All isolates (n = 60) were analyzed using NG-MAST as described previously.[9] Both forward and reverse sequences of por and tbpB genes were analyzed by Big Dye Terminator v 3.1 Cycle sequencing kit (Applied Biosystem, California) as per manufacturer’s instructions on 3130 × 1 genetic analyzer (Applied Biosystems). The forward and reverse sequences of the por and tbpB genes from each isolate were aligned using online available software LALIGN and trimmed to set length from conserved regions as described. Each unique sequence at por and tbpB was assigned an “allelic number” providing each gonococcal isolate with a two-digit allelic profile and each unique allelic profile was designated a numeric ST from the database of the NG-MAST website (http://www.ng-mast.net). The WHO NG reference strains F, G, K-P were used as control.

**Discriminatory index**
The discriminatory capacity of each typing method was determined by calculation of a numerical index, D,
i.e., the Hunter-Gaston discriminatory index using the following formula:[10]

\[ D = 1 - \frac{1}{N(N-1)} \sum_{i=1}^{S} n_i (n_i - 1) \]

Where \( N \) is the number of isolates in the total population, \( S \) is the total number of types found, and \( n_i \) number of isolates belonging to \( j \)th type.

The study was approved by the Ethics Committee, All India Institute of Medical Sciences, New Delhi, India (Ref. No. IEC/NP-A-52/2006 and OP-18/2013 and RP-01/05.08.2013 and IESC/T-170/04.05.2012).

**Results**

**Antimicrobial susceptibility testing**

Of the 60 isolates, 32 (53.3%) were PPNG and only 1 (1.6%) was CMRNG; 16 (26.6%) were CMRNG, while 18 (30%) were TRNG. Both PPNG and TRNG were found in 13 (21.6%) while none were CMRNG. QRNG was seen in 93.3%, while 5% were azithromycin resistant and 6.6% showed DS to ceftriaxone. All the isolates were susceptible to spectinomycin. MIC\(_{90}\) for penicillin, tetracycline, ciprofloxacin, spectinomycin, ceftriaxone, and azithromycin was 1 μg/ml, 2 μg/ml, 3 μg/ml, 5 μg/ml, 0.012 μg/ml, and 0.25 μg/ml, respectively [Table 1]. There were no multidrug-resistant (MDR)-NG and extensively drug-resistant (XDR)-NG as per recently proposed criteria. Further, isolates showing DS to ceftriaxone were resistant to penicillin, ciprofloxacin and less susceptible to tetracycline [Table 2]. One isolate was also resistant to azithromycin. Antibiogram had a discriminatory index of 0.77.

**Auxotypes**

Out of 60 isolates, 24 (40%) were NR, 16 (26.6%) were PR, 13 (21.6%) AR, and 7 (11.6%) belonged to others. The discriminatory index of auxotyping was 0.72.

**Genotyping (Neisseria gonorrhoeae-multiantigen sequence typing)**

Sixty gonococcal isolates were distributed among 40 different STs, 29 (72.5%) of which were novel. The majority of STs (77.5%, 31 of 40) were represented by a single isolate and remaining STs included between 2 and 13 isolates. The most common ST was 6058 (\( n = 13 \)) followed by ST621, ST1572, ST1751, ST2990, ST6069, ST9774, ST9783, and ST9786 (two each). A total of 31 por and 27 tbpB alleles were found in this study. The discriminatory index of NG MAST was 0.95.

**Association between minimum inhibitory concentrations and auxotypes**

All auxotypes were sensitive to spectinomycin. Isolates exhibiting resistance to azithromycin belonged to auxotypes NR, PR, AR - one each and DS to ceftriaxone belonged to auxotypes NR (\( n = 1 \)), PR (\( n = 2 \), AR (\( n = 1 \)). MIC\(_{90}\) for ceftriaxone was highest in auxotype NR (0.012 μg/ml) followed by AR (0.008 μg/ml) and PR (0.006 μg/ml). However, there was no difference in MIC\(_{90}\) for azithromycin (0.25 μg/ml) in these three auxotypes.

**Association between minimum inhibitory concentrations and Neisseria gonorrhoeae-multi-antigen sequence typing**

Isolates showing DS to ceftriaxone belonged to ST6069 (\( n = 2 \)), ST8156, ST6083 (one each) and the ones resistant to azithromycin belonged to ST9779, ST1572,

**Table 1: Antimicrobial susceptibility of Neisseria gonorrhoeae isolates (n=60) by E-test**

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Susceptible (%)</th>
<th>Less susceptible/decreased susceptibility (%)</th>
<th>Resistant (%)</th>
<th>MIC(_{90}) (μg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penicillin</td>
<td>0</td>
<td>27 (45)</td>
<td>33 (55)*</td>
<td>1</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>9 (15)</td>
<td>17 (28.3)</td>
<td>34 (56.6)*</td>
<td>2</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>2 (3.3)</td>
<td>2 (3.3)</td>
<td>56 (93.3)**</td>
<td>3</td>
</tr>
<tr>
<td>Spectinomycin</td>
<td>60 (100)</td>
<td>0</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>56 (93.3)</td>
<td>4 (6.7)</td>
<td>0</td>
<td>0.012</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>57 (95)</td>
<td>0</td>
<td>3 (5)</td>
<td>0.25</td>
</tr>
</tbody>
</table>

*PPNG = Penicillinase producing N. gonorrhoeae, TRNG = Tetracycline-resistant N. gonorrhoeae, HLR = High level resistant, MIC = Minimum inhibitory concentration. N. gonorrhoeae = Neisseria gonorrhoeae

**Table 2: Susceptibility pattern of ceftriaxone decreased susceptible strains to other antimicrobial agents**

<table>
<thead>
<tr>
<th>Ceftriaxone</th>
<th>Penicillin</th>
<th>Tetracycline</th>
<th>Ciprofloxacin</th>
<th>Azithromycin</th>
<th>β-lactamase</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIC</td>
<td>Category</td>
<td>MIC</td>
<td>Category</td>
<td>MIC</td>
<td>Category</td>
</tr>
<tr>
<td>0.064</td>
<td>DS</td>
<td>1.0</td>
<td>CMR</td>
<td>4.0</td>
<td>CMR</td>
</tr>
<tr>
<td>0.064</td>
<td>DS</td>
<td>1.5</td>
<td>CMR</td>
<td>0.75</td>
<td>LS</td>
</tr>
<tr>
<td>0.064</td>
<td>DS</td>
<td>&gt;32</td>
<td>R</td>
<td>0.5</td>
<td>LS</td>
</tr>
<tr>
<td>0.125</td>
<td>DS</td>
<td>1.5</td>
<td>CMR</td>
<td>1</td>
<td>LS</td>
</tr>
</tbody>
</table>

MIC = Minimum inhibitory concentration, DS = Decreased susceptibility, R = Resistant, LS = Less susceptible, HLR = High-level resistant, CMR = Chromosomally mediated resistance, S = Susceptible, + = β-lactamase producer, - = β-lactamase Non-producer
and ST6083 (one each). Out of 40 identified STs, the most common ST was 6058 (n = 13). Out of 13 ST6058, 12 (92.3%) were PPNG and none were CMRNG; 9 (69.2%) were CMRNG while 4 (30.7%) were TRNG. Both PPNG and TRNG were found in 4 (30.76%). All isolates belonging to ST6058 were resistance to ciprofloxacin and sensitive to azithromycin, ceftriaxone and spectinomycin. There was significant association between ST6058 and PPNG (P = 0.00) and CMRNG (P = 0.003). In all other resistance phenotypes, no association was found.

Discussion

Besides characterization of gonococcal strains, typing is also useful in studying of relationship between NG isolates during outbreaks in sexual networks, investigating transmission of antibiotic-resistant strains, and differentiating re-infection from treatment failure. Typing methods used should be discriminatory; however, at the same time, it is important that they are easy to perform, practical, and inexpensive. Among the various typing methods to characterize NG isolates i.e., phenotypic and genotypic, the widely used phenotypic methods are based on auxotyping, serotyping, and antimicrobial susceptibility testing. The phenotypic methods are less efficient and do not meet the desired discriminatory index.[11] Further, auxotyping being laborious and time-consuming, has been replaced by genotypic methods world over. Among genotypic methods, NG-MAST is a leading method which is based on sequencing and allows international comparison of data and tracing of gonococci of public health importance. In India, very few studies have been carried out on gonococcal typing methods and most have focused on phenotypic methods only.[6,7] No study from India comparing the phenotypic and genotypic methods has been documented so far to the best of our knowledge.

As regards AMR pattern as a typing method, many studies from India only differentiate isolates into PPNG and non-PPNG or into TRNG and non-TRNG. The incidence of PPNG varies from 8% to 88.9% in various parts of India.[12] In the present study, 53.3% strains were PPNG while only 1.6% demonstrated chromosomal resistance, thereby implying that majority of penicillin resistance to ciprofloxacin and sensitive to tetracycline. A study from India showed increasing trends of TRNG from 28.1% to 58.5% between year 2009 and 2011.[12] However, reporting for tetracycline in India is technique based and not all laboratories categorize the non-TRNG strains further.[12] A recent study from Shanghai reports tetracycline resistance rates to be 54.7% of which TRNG was 31% and chromosomal resistance was seen in 23.7%, a finding similar to ours.[17] Overall, we observed both, chromosomal and plasmid-mediated resistance to penicillin and tetracycline. In the present study, 93.3% isolates were resistant to ciprofloxacin. Although no longer recommended for treatment of gonorrhea, ciprofloxacin resistance rates were frightening. Previously, very high rates of resistance to ciprofloxacin have been reported from Delhi. On reviewing literature, we found that considerable differences exist in ciprofloxacin resistance from different parts of India ranging from 49.1% to 100%. All over the world, there is a variation in the rates of resistance to fluoroquinolones and is seen in 10.2% to 27.3% cases in US, 33.6% in Australia and 78.2% in Japan.[15,16,18] Rates of resistance to drug used in syndromic management in our country were low. A total of 6.6% isolates exhibited DS to ceftriaxone while 5% were azithromycin resistant. There appears no threat to the in-use therapies administered in syndromic approach, i.e., cefixime 400 mg plus azithromycin 1 g orally as a single dose. Isolates with DS to ceftriaxone almost always exhibit resistance to quinolones or quinolones and penicillin as observed in our study too. Overall, chromosomal resistance was more often associated with DS to ceftriaxone [Table 2]. Our findings, although on a small number of isolates, are in line with the national as well as international data.[19-23] Our isolates were susceptible to spectinomycin which support the finding of the previous studies from India, where no resistance to this drug has been documented except isolated case report.[24] There were no MDR-NG and XDR-NG.

The discriminatory index of antibiogram in our study was 0.77 and was higher than previous studies reported from India.[11,25] This could possibly be because we have taken into account both, plasmid as well as chromosomal resistance to antimicrobials. The NR and PR auxotypes are most common auxotypes in Delhi and Mumbai and the same was observed by us.[11] Restriction of certain auxotypes to different geographical areas can help in global epidemiological studies. Both PPNG and TRNG were demonstrating NR auxotype as the most common auxotype; however, the association was not significant. This is in line with the previous Indian studies.[11] Isolates exhibiting resistant to azithromycin and DS to ceftriaxone belonged to NR, PR, and AR auxotypes. The discriminatory index of auxotyping was 0.72, which is higher than previous studies reported from India.[11,25] This is surprising as we, in our study,
have not investigated the less common nutritional requirements such as methionine, hypoxanthine, uracil, thiamine, pyrophosphate, and isoleucine. However, some workers in India have combined auxotyping with another phenotypic method to achieve a desirable discriminatory index.[11]

NG-MAST has been used as a tool to enhance surveillance of gonococcal AMR and to study the clonal spread in high-risk population. Our strains were distributed among 40 different STs of which 29 (72.5%) were novel. This reflects the adaptation of gonococci to this geographic region and evolution of STs in India. The most common ST was ST6058 the same has been reported by us in the previous year.[26] ST6058 was significantly associated with resistance phenotypes, PPNG and CMRNG. In all other resistance phenotypes, no association was found. All isolates belonging to ST6058 were sensitive to drugs in use. Further, the DS isolates belonged to different STs (ST6069 [n = 2], ST8156 and ST6063 [one each]), thereby demonstrating heterogeneity which suggests the introduction of strains from multiple sources rather than clonal spread. None of our DS isolates belonged to ST 1407, the internationally documented MDR gonococcal clone.[27,28] Isolates exhibiting DS to ceftriaxone belonged to auxotypes NR, AR (one each), PR (n = 2), and the NR and PR auxotypes surprisingly belonged to the same ST6069 and had similar por and tbp B alleles. Therefore, most probably, these isolates represent infection with the same strain. Isolates resistant to azithromycin belonged to auxotypes NR, PR, and AR (one each), of which the AR strain belonged to a novel ST9779.

The D value of NG-MAST in our study was 0.95, thereby implying that it is a highly efficient technique. Further, molecular epidemiological approaches help to furnish additional information on the origin and spread of AMR.

Conclusions

Gonococci are notorious for high rates and rapid spread of drug-resistant strains and therefore warrant that such facilities be introduced in the routine surveillance programs to achieve better disease control. However, their prohibitive cost preclude use in a setting like ours where it is feasible only when funds are available for such research activities. On the other hand, although phenotypic methods have several limitations, they allow the comparison of recent data with the older data.

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Conflicts of interest

There are no conflicts of interest.

References


