Molecular tests for the detection of antimicrobial resistant *Neisseria gonorrhoeae*: when, where and how to use?

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Purpose of review

Molecular methods for the diagnosis of *Neisseria gonorrhoeae* are replacing bacterial culture in many settings. This review focuses on recent progress in the development of molecular tests to detect resistant *N. gonorrhoeae* both to enhance surveillance and to guide decisions about individual patient management.

Recent findings

Assays that detect determinants of resistance for all antibiotics used as first-line gonorrhoea treatment or to detect specific "superbug" strains have been developed to enhance surveillance, but few have been applied in clinical practice. The strategy of most relevance to individual case management that has advanced furthest is to identify ciprofloxacin-sensitive strains so that unnecessary use of ceftriaxone can be avoided. Cross-reactivity with pharyngeal commensal Neisseria species reduces specificity and is a challenge for many assays.

Summary

Progress with laboratory-based molecular tests to detect gonococcal resistance is being made but substantial challenges remain. No laboratory-based assay has been subjected to a field evaluation and no assay so far can be used as a point-of-care test. Given the threat of antimicrobial resistance, now is the time to exploit the molecular technologies used for diagnosis and to invest in the development of molecular gonococcal resistance tests that can be implemented for public health good.

Keywords

Neisseria gonorrhoeae, antimicrobial resistance, antimicrobial surveillance, point-of-care tests

Abbreviations

MIC, minimum inhibitory concentration NAAT, nucleic acid amplification test PBP2, penicillin binding protein 2 PPNG, penicillinase producing *Neisseria gonorrhoeae* WHO, World Health Organization

Introduction

Antimicrobial resistant *Neisseria gonorrhoeae* is a global public health challenge [1]. New diagnostic strategies and novel antimicrobials are urgently needed to conserve ceftriaxone [2], the last antimicrobial for empirical first-line monotherapy for gonorrhoea in many countries [3,4*]. Gonococcal strains with resistance to extended-spectrum cephalosporins have caused treatment failure with ceftriaxone and/or cefixime in Europe, North America, Asia and Africa [5-9]. In this evolving situation, dual antimicrobial therapy (ceftriaxone plus azithromycin) has been introduced for first-line treatment in Europe, Australia and the USA [10].

Detection of antimicrobial resistant gonococci using molecular methods is a necessary innovation [2,6,11,12] because nucleic acid amplification tests (NAAT) are now the most widely used assays for gonorrhoea diagnosis in many countries [11,13,14**,15]. The World Health Organization (WHO) global action plan to mitigate the spread of multidrug-resistant gonococci calls for increased surveillance of antimicrobial resistant N. gonorrhoeae globally, strengthened capacity for bacterial culture and research into molecular methods to monitor and detect resistance [1]. We searched the US National Library of Medicine (OvidSP, Wolters Kluwer Health, New York) using the medical subject headings 'Neisseria gonorrhoeae' or 'gonorrhea' and 'drug resistance, bacterial' and 'nucleic acid amplification techniques' in September 2015. We selected articles from this search, reference lists of review articles [3,4*,6,14**,16] and abstracts of the World STI & HIV Congress 2015. We found no articles about commercially available molecular assays that detect genetic resistance determinants of N. gonorrhoeae, but the number of reports of laboratory-developed assays is increasing [17*,18-20,21*,22-26,27*,28-38]. This review examines the requirements, recent progress and challenges for the molecular detection of antimicrobial resistant N. gonorrhoeae in clinical specimens from the point of view of the two main functions of resistance testing: to enhance surveillance [14**] and to guide decisions about individual patient management [15]. We focus on

publications since 2014 about assays developed to detect genetic determinants of *N. gonorrhoeae* resistance in clinical specimens and for which results have been compared with MICs for specific antimicrobials.

The need for molecular detection of antimicrobial resistant *Neisseria gonorrhoeae* The molecular technologies have changed patterns of gonorrhoea diagnosis in high income countries. NAAT are recommended because they have high sensitivity, robust specimen handling conditions, can be automated with high throughput and can be used on non-invasively collected samples like first-void urine and vaginal swabs [39]. The rapid adoption of NAAT has, however, raised two main concerns about the ability to detect, monitor and manage antimicrobial resistance [1,6,11,14**,15]. First, commercial NAAT specimens provide no viable organism for antimicrobial susceptibility testing. Second, as the number of specimens for culture falls the microbiological skills required for isolation and antimicrobial susceptibility testing are being lost [1]. Countries that have poor healthcare infrastructure and rely on syndromic management of genitourinary symptoms and/or microscopy of Gram-stained smears have a particularly urgent need for better diagnostics and knowledge of gonococcal antimicrobial resistance patterns [40].

Molecular technology is re-defining expectations about the detection of antimicrobial resistant gonorrhoea. The paradigm for sexually transmitted infection treatment reflects the clinical priority to treat symptomatic patients on presentation [3]. In the absence of results of antimicrobial susceptibility testing, the first-line treatment regimen should cure at least 95% of infections [3]. Bacterial culture and antimicrobial susceptibility testing underpin the success of this strategy and have many advantages [11,14**]. First, culture on selective media directly provides the biological material for assessing a strain's susceptibility to multiple antimicrobials simultaneously [3]. Second, breakpoints for the minimum inhibitory concentration (MIC) that determines resistance correspond with clinical treatment failure for many antimicrobials. Third, organised surveillance programmes can monitor changes in the resistance prevalence for many antimicrobials [41-43]. Fourth, clinicians

can use the antimicrobial susceptibility profile to alter therapy, or choose an antimicrobial for an untreated patient. The use of NAATs alone precludes monitoring of antimicrobial resistance, but has also highlighted the limitations of culture. First, considerable technical skills and time are required to culture the fastidious *N. gonorrhoeae* and test for antimicrobial resistance. Second, culture has poor sensitivity for extragenital sites such as the pharynx [3], where horizontal transfer of resistance determinants from commensal *Neisseria* species is thought to be the origin of resistance to extended-spectrum cephalosporins [4*,6].

Requirements of molecular tests to detect antimicrobial resistant *N. gonorrhoeae* Ideally, molecular resistance testing should reflect the exact MICs of different antimicrobials. Resistance to many antimicrobials in *N. gonorrhoeae* is affected by multiple genes, however. Different mutations and an accumulation of these mutations result in the high MIC of the specific antimicrobial [4*]. This characteristic makes it exceedingly difficult to use molecular assays to predict exact MICs of antimicrobials, but by targeting the main resistance determinants the resistance/susceptibility phenotypes can be predicted. The principal genes associated with gonococcal antimicrobial resistance are shown in Table 1.

Most publications about molecular tests to detect gonococcal resistance determinants report their intended use for surveillance purposes [17*,18–20,21*,22–26,27*,28–31]. Fewer mention the need or potential for guiding individual patient management [18,27*,31-35]. There are many similarities in the requirements for assays for both purposes. For example, the capacity for direct detection in clinical specimens [17*,18,19,21*,24,26,27*,29-32,44*] is critical if the assays are to be used in situations where bacterial culture has not been performed. Table 2 lists similarities and differences between test requirements for surveillance and clinical management.

Enhancing surveillance of antimicrobial resistant *N. gonorrhoeae* Experts agree that molecular tests can enhance but not replace culture-based surveillance [1,4*,6,14**]. The lack of correspondence between genetic resistance determinants, MICs and clinical treatment failure, particularly for evolving resistance in extended-spectrum cephalosporins remains a critical stumbling block [6]. Nevertheless, molecular assays to enhance surveillance can advance more quickly than those for clinical management partly because the assays can be solely laboratory-based, short reaction times are not essential and sensitivity and specificity can be suboptimal [26, 31] (as long as they allow consistent monitoring of trends) because the results will not influence individual patient management (Table 2). As new resistant strains emerge or levels of recognised resistance increase, culture-based surveillance can be broadened to confirm the findings of molecular tests.

Some assays have been used in real life settings. A real-time PCR assay was developed to detect penicillinase producing gonococci [30] in remote regions of Australia where resistance to penicillin remains below 5%. The molecular assay had 100% sensitivity and 98.7% specificity compared with phenotypic tests. When applied subsequently to *N. gonorrhoeae* positive NAAT specimens from Western Australia the assay detected PPNG in 15/915 (1.6%) of specimens from remote areas and 34/303 (11.2%) of specimens from more heavily populated areas [44*]. As a result, the treatment regimen of amoxicillin, probenecid and azithromycin was continued in remote areas and, in populous areas, changed from ceftriaxone alone to ceftriaxone plus azithromycin [44*]. This assay has been further optimised to include additional β -lactamase plasmid targets [19]. Molecular tests for surveillance can also be used to monitor specific strains. For example, real-time PCR assays have been developed to screen clinical specimens for extensively drug resistant gonococcal strains ("superbugs") that were first detected in Japan (H041) [29] and subsequently in Europe (F89) [24].

An antibiogram gives information about susceptibility to multiple antimicrobials. Most molecular assays detect resistance determinants and relate to a single antibiotic (penicillin [19,30,44]*; ciprofloxacin [18,31,32]; azithromycin [21*,45]; extended-spectrum cephalosporins [17*]) or a single strain [24,29]. Molecular assays aiming to detect resistance to several antimicrobials have been run in series [26] or as multiplex reactions [27*,36], but their analytical sensitivity was too low to be

used in clinical specimens [36] or, if assessed on clinical specimens, was not validated against a culture-based MIC reference standard [26,27*].

Laboratory-based molecular tests to guide individual patient management Laboratory-based molecular assays could have advantages if they produce faster results and have greater sensitivity than bacterial culture, particularly when used on extragenital specimens for NAAT testing. Laboratory-based assays will usually not help, however, in the management of patients who present with symptoms or who need same day treatment for other reasons.

The strategy that has advanced furthest aims to spare overuse of ceftriaxone plus azithromycin (the only remaining first-line empirical treatment in many countries), by targeting ciprofloxacin to patients with ciprofloxacin-sensitive gonococcal strains before treatment is initiated. Several laboratory-based molecular assays for use in clinical specimens detect one or more chromosomal mutations in gonococci that are associated with ciprofloxacin resistance [18,26,27*,31,32]. Siedner and colleagues designed a real-time PCR assay to detect resistance using a single target mutation in urine specimens [31]. Single nucleotide polymorphism(s) in amino acid codon S91 in the gyrA gene is found in all ciprofloxacin resistant strains, even though additional mutations in gyrA and other loci (Table 1) contribute to higher levels of resistance $[4^*, 46]$. The prototype assay amplified the target sequence in only 85% of all urine NAAT specimens tested [31]. Further optimisation of this assay has improved sensitivity in cultured isolates, which should be validated in clinical specimens [37]. The suboptimal diagnostic test accuracy of real-time PCR assays to detect ciprofloxacin resistance in clinical specimens presents ongoing challenges. In a study that tested 24 NAAT specimens, real-time PCR assays correctly identified only 4/6 ciprofloxacin-resistant specimens with S91 mutations and 16/18 ciprofloxacin-sensitive specimens [18]. Three of four specimens that failed were from the pharynx or rectum. Cross-reactivity occurred with N. meningitidis, N. lactamica, N. subflava and N. polysaccharea. In another study, real-time PCR failed in 10% (28/290) of clinical specimens, particularly those from non-genital sites [38].

An alternative strategy is to identify wild type *gyrA* sequences that predict ciprofloxacin susceptibility rather than resistance [32]. To exclude cross-reactivity, a *N. gonorrhoeae* specific target (*dcmH*) was also detected in the molecular assay. When used in male urine NAAT specimens in South Africa, the assay correctly identified 15/15 ciprofloxacin-sensitive and 18/18 strains with intermediate susceptibility or resistance [32].

Point-of-care molecular tests to guide individual patient management

Point-of-care tests have a rapid turnaround time that guides clinical decisions and allows results and treatment to be given to the patient at the same visit [47]. None of the molecular assays developed so far to detect antimicrobial resistance in *N. gonorrhoeae* is a "transformative point-of-care diagnostic test that will conserve antibiotics for future generations... [and is] accurate, rapid, affordable, easy-to-use and available to anyone, anywhere in the world" [48]. These criteria, set by the Longitude Prize, aim to stimulate innovation in diagnostic test development for all infectious diseases. Even the GeneXpert[®] (Cepheid, Sunnyvale, WA, USA) tests for gonorrhoea and chlamydia diagnosis are not rapid point-of-care tests because the NAAT takes 90 minutes, rather than a 30 minutes benchmark set by international groups, and their high cost precludes widespread use [40,48,49].

The requirements of a point-of-care test focusing on accurate and cost-effective detection of a limited number of targets without the need for separate DNA extraction methods [50] need to be differentiated from tests for surveillance, which might detect many resistance determinants in multistage multiplex assays [27*] (Table 2). Advances in bioengineering and nanotechnology, such as microfluidics, will help to adapt laboratory-based systems to clinic-based formats [40,51]. Target product profiles that define the user, patient population and point-of-care device requirements are essential [47]. Evaluation trials should use clinically relevant endpoints [40,52], such as prediction of resistant phenotypes and reduced antibiotic prescribing. The ideal system would detect both *N*. *gonorrhoeae* and its main resistance determinants at the point-of-care and allow individualised

antimicrobial treatment to help slow the spread of antimicrobial resistance, particularly in resourcepoor settings.

Molecular detection of extended-spectrum cephalosporin resistance

Resistance to extended-spectrum cephalosporins remains the most urgent threat. The development of molecular assays to detect any resistance to extended-spectrum cephalosporins in clinical specimens is inherently challenging because multiple mechanisms are involved and are still evolving [4ç]. Mosaic alleles of the penA gene encoding penicillin binding protein 2 (PBP2) and non-mosaic penA alleles with A501 mutations are the main determinants of decreased susceptibility and resistance to extended-spectrum cephalosporins [4*]. The gonococcal "superbug" F89 strain is a good example of how resistance to extended-spectrum cephalosporins evolves. This strain contains a penA mosaic allele (type XXXIV), which is common worldwide [4*,5-7], with an additional A501P alteration, resulting in high-level ceftriaxone resistance [9]. Molecular test specificity is a major challenge, particularly for pharyngeal specimens that frequently harbour non-gonococcal Neisseria species with similar penA sequences. Several assays have examined the presence of mosaic penA allele sequences in cultured isolates [25,50,53]. One real-time PCR assay that detects N. gonorrhoeae (porA pseudogene), mosaic penA alleles and mutations in additional resistancedetermining loci (mtrR, porB and ponA) was tested on 24 gonococcal-negative NAAT specimens and 34 gonococcal-positive NAAT specimens [17*]. The assay detected a mosaic *penA* allele in one specimen with the highest ceftriaxone MIC (0.25 μ g/ml), but lacked specificity in the prediction of decreased susceptibility to extended-spectrum cephalosporins (in clinical specimens and culture isolates). This suboptimal specificity will be exceedingly difficult to overcome because this assay and other similar assays detect many different mosaic penA alleles which result in highly divergent MICs of the extended-spectrum cephalosporins [4*,6,8]. Furthermore, cross-reactivity in the penA, mtrR, porB and ponA targets with commensal Neisseria isolates or in clinical specimens was identified. All three clinical cross-reactive specimens were from the pharynx [17*].

Molecular detection of azithromycin resistance

Azithromycin alone is not a first-line treatment for gonorrhoea, but detecting resistance is important because of its use in dual treatment regimens. Two mutations in the 23S ribosomal RNA genes (A2059G and C2611T) are associated with azithromycin resistance with MICs depending on how many of the four alleles are mutated [45,54]. Two real-time PCR assays could characterise the 23S rRNA 2059 and 2611 positions as wild type or mutated in 87% (266/306) of genital and rectal specimens. However, cross-reactivity was observed with both assays testing commensal Neisseria species and in 33% (7/21) of pharyngeal samples [21*]. Furthermore, among 64 samples with MIC results, 3% (2/64) with raised MICs (1 μ g/ml), which indicates resistance, were identified as 23S rRNA wild type.

Challenges for molecular detection of antimicrobial resistant N. gonorrhoeae

The number of assays and techniques used to detect genetic determinants of antimicrobial resistant *N. gonorrhoeae* is increasing and some assays are beginning to be adopted for surveillance [14**]. Progress in molecular assays to guide clinical management is lagging, however. In addition to the challenges to developing usable and commercially viable point-of-care tests [47], there are many remaining roadblocks for the development of diagnostic tools to contain antimicrobial resistance in general [40] and for gonorrhoea in particular. First, molecular tests can only detect known targets and new mutations and resistance mechanisms will develop for both extended-spectrum cephalosporins and other antimicrobials. Adaptable assays and maybe new technologies such as whole genome sequencing might be required to overcome this challenge. Second, incomplete understanding of the relationship between genetic resistance determinants and phenotypic resistance [4*] needs to be overcome to improve the validity of evaluation studies. Third, diagnostic evaluation studies that compare detection of genetic resistance determinants with MICs are small and their methodology is subject to biases that overestimate test performance [52]. Larger field studies with blinded evaluation of a range of specimen types and results are needed. Fourth, test sensitivity and specificity are still suboptimal for clinical specimens, particularly for pharyngeal

specimens. Some cross-reactivity with non-gonococcal Neisseria species that carry the same genetic sequences cannot be overcome but advances in primer and assay design might improve test accuracy. Finally, diagnostic tests can help to implement and monitor treatment strategies to spare first-line empirical therapy with ceftriaxone plus azithromycin. But targeting ciprofloxacin treatment, using azithromycin as co-therapy and other actions are short-term solutions with unknown consequences for the spread of antimicrobial resistant *N. gonorrhoeae*. In the medium term, new therapeutic antimicrobials are crucial. A gonococcal vaccine [55] is probably the only long term solution.

Conclusions

Progress with the development of molecular tests to detect gonococcal resistance is significantly more advanced for enhancing surveillance than for guiding clinical decision making but substantial challenges remain. Gonococcal culture and antimicrobial susceptibility testing remain essential tools for surveillance and as a reference standard for assay validation. The choice of which molecular resistance determinants to prioritise needs to consider both short and longer term antimicrobial resistance threats. Cross-reactivity with non-gonococcal Neisseria species, particularly in the pharynx, is a challenge for all assays and suggests that assay development should focus first on genital and rectal clinical specimens. Now is the time to exploit the molecular technologies used for diagnosis and to invest in the development of molecular gonococcal resistance tests that can be implemented for public health good.

Antimicrobial	Principal gene(s) and position ^a	Other genes with verified impact	
		on resistance in clinical strains ^{a,c}	
Ciprofloxacin	gyrA S91 (D95)	parC	
Ceftriaxone	penA mosaic allele (A501-altered penA alleles)	mtrR, porB	
Azithromycin	23S rRNA gene: C2611 (lower level of resistance) and A2059 (high-level resistance) ^b	mtrR, mefA, erm genes	
Spectinomycin	16S rRNA gene: C1192	rpsE	
Tetracycline	<i>tetM</i> -carrying plasmid (high-level resistance) and <i>rpsJ</i> V57 (lower level of resistance)	mtrR, porB	
Penicillin	β-lactamase plasmid (<i>bla</i> _{TEM} gene; high-level resistance) and <i>penA</i> (D345 insertion or mosaic allele; lower level of resistance)	mtrR, porB, ponA	

Table 1 Main genes associated with antimicrobial resistance in Neisseria gonorrhoeae

^aWith exception of the *tetM*-carrying plasmid and the β -lactamase plasmid, listed genes and positions are involved in

chromosomally-mediated resistance. Wild type positions are listed, which can have many different mutations resulting in resistance.

^bThere are four alleles of the 23S rRNA gene in each gonococcal strain.

^cThere are additional efflux pumps, e.g. MacAB and NorM, in *N. gonorrhoeae* that affect the MICs of several antimicrobials.

Characteristic	Surveillance	Clinical management	
		Asymptomatic patient	Symptomatic patient
Specimen type	Culture or clinical	Culture or clinical	Clinical specimen
	specimen	specimen	
Location of test	Laboratory	Laboratory or	Point-of-care
equipment		point-of-care	
Time from sample	Not critical	Same as or quicker	Minimal
collection to reporting		than culture-based	
result		testing	
Technical skill required	Skilled laboratory	Skilled laboratory	Minimal
	technician	technician or minimal	
Skill required to	Minimal-High	Minimal-intermediate	Minimal
interpret results			
Accuracy	Intermediate-High	High	High
Cost	Low-High	Low-intermediate	Low
Scalable	Not critical	Desirable	Critical
Multiplex	Not critical	Desirable	Critical

Table 2 Requirements of molecular tests to detect antimicrobial resistant Neisseria gonorrhoeae

Key points

- Molecular methods are increasingly used for gonorrhoea diagnosis but, despite the global threat of antimicrobial resistance, commercially available diagnostic assays do not detect gonococcal resistance determinants;
- The requirements of molecular tests to detect gonococcal resistance depend on the purpose of the assay; to enhance surveillance of antimicrobial resistance, or to guide the clinical management of gonorrhoea;
- Molecular detection of resistance determinants in genital and rectal specimens is most accurate; cross-reactivity with non-gonococcal Neisseria species particularly in the pharynx reduces assay specificity;
- Detection of ciprofloxacin resistance or susceptibility to spare the use of extended spectrum cephalosporins is the strategy that has advanced the furthest to date, but is not a long term solution.
- Now is the time to exploit the molecular technologies used for diagnosis and to invest in the development of molecular gonococcal resistance tests that can be implemented for public health good.

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Conflicts of interest

N.L and M.U. are co-investigators in the RaDAR-Go project which aims to develop a point-of-care test to detect antimicrobial resistance in *Neisseria gonorrhoeae*.

References and recommended reading Papers of particular interest have been highlighted as:

* of special interest

** of outstanding interest

[1] World Health Organization. Global action plan to control the spread and impact of antimicrobial resistance in *Neisseria gonorrhoeae* In: Geneva: World Health Organization; 2012.

[2] Laxminarayan R. Antibiotic effectiveness: Balancing conservation against innovation. Science 2014; 345:1299-1301.

[3] Tapsall J. Antimicrobial resistance in *Neisseria gonorrhoeae*. WHO/CDS/CSR/2001.3. In: Geneva: 2001. pp. 1-58.

*[4] Unemo M, Shafer WM. Antimicrobial Resistance in Neisseria gonorrhoeae in the 21st Century: Past, Evolution, and Future. Clin Microbiol Rev 2014; 27:587-613.

This comprehensive review describes *Neisseria gonorrhoeae* resistance mechanisms and determinants. The extensive literature review can be used as a source for potential targets for molecular resistance tests.

[5] Lewis DA, Sriruttan C, Muller EE *et al.* Phenotypic and genetic characterization of the first two cases of extended-spectrum-cephalosporin-resistant *Neisseria gonorrhoeae* infection in South Africa and association with cefixime treatment failure. J Antimicrob Chemotherapy 2013; 68:1267-1270.

[6] Unemo M, Nicholas RA. Emergence of multidrug-resistant, extensively drug-resistant and untreatable gonorrhea. Future Microbiol 2012; 7:1401-1422.

[7] Allen VG, Farrell DJ, Rebbapragada A *et al.* Molecular analysis of antimicrobial resistance
 mechanisms in *Neisseria gonorrhoeae* isolates from Ontario, Canada. Antimicrob Agents Chemother.
 2011; 55:703-712.

[8] Ohnishi M, Golparian D, Shimuta K *et al.* Is *Neisseria gonorrhoeae* initiating a future era of untreatable gonorrhea?: detailed characterization of the first strain with high-level resistance to ceftriaxone. Antimicrob Agents Chemother 2011; 55:3538-3545.

[9] Unemo M, Golparian D, Nicholas R *et al.* High-level cefixime- and ceftriaxone-resistant Neisseria gonorrhoeae in France: novel penA mosaic allele in a successful international clone causes treatment failure. Antimicrob Agents Chemother 2012; 56:1273-1280.

[10] Unemo M. Current and future antimicrobial treatment of gonorrhoea - the rapidly evolving Neisseria gonorrhoeae continues to challenge. BMC Infect Dis 2015; 15:364.

[11] Low N, Unemo M, Skov Jensen J *et al.* Molecular diagnostics for gonorrhoea: implications for antimicrobial resistance and the threat of untreatable gonorrhoea. PLoS Med 2014; 11:e1001598.

[12] Bignell C, Unemo M. 2012 European Guideline on the diagnosis and treatment of gonorrhoea.Int J STD AIDS 2013; 24:85-92.

[13] Dicker LW, Mosure DJ, Steece R, Stone KM. Laboratory tests used in US public health laboratories for sexually transmitted diseases, 2004. Sex Transm Dis 2007; 34:41-46.

**[14] Goire N, Lahra MM, Chen M *et al.* Molecular approaches to enhance surveillance of gonococcal antimicrobial resistance. Nat Rev Microbiol 2014; 12:223-229.

This review explains in detail how molecular tests can be use to enhance surveillance for antimicrobial resistant gonorrhoea.

[15] Buono SA, Watson TD, Borenstein LA *et al.* Stemming the tide of drug-resistant Neisseria
 gonorrhoeae: the need for an individualized approach to treatment. J Antimicrob Chemother 2015;
 70:374-381.

[16] Lewis DA. Global resistance of Neisseria gonorrhoeae: when theory becomes reality. Curr Opin Infect Dis 2014; 27:62-67.

*[17] Peterson SW, Martin I, Demczuk W *et al.* Molecular Assay for Detection of Genetic Markers Associated with Decreased Susceptibility to Cephalosporins in Neisseria gonorrhoeae. J Clin Microbiol 2015; 53:2042-2048.

This study describes the most recent assay for the detection of gonococcal resistance to extended spectrum cephalosporins in clinical specimens. Assay sensitivity was sufficient to detect a raised ceftriaxone MIC but lacked specificity.

[18] Peterson SW, Martin I, Demczuk W *et al.* Molecular assay for the detection of ciprofloxacin resistance in Neisseria gonorrhoeae from cultures and clinical Nucleic Acid Amplification Test (NAAT) specimens. J Clin Microbiol 2015 doi:10.1128/JCM.01632-15.

[19] Buckley C, Trembizki E, Baird RW *et al.* Multitarget PCR Assay for Direct Detection of Penicillinase-Producing Neisseria gonorrhoeae for Enhanced Surveillance of Gonococcal Antimicrobial Resistance. J Clin Microbiol 2015; 53:2706-2708. [20] Lawung R, Cherdtrakulkiat R, Charoenwatanachokchai A *et al.* Antimicrobial resistance markers as a monitoring index of gonorrhoea in Thailand. Acta Microbiol Immunol Hung 2012; 59:157-169.

*[21] Trembizki E, Buckley C, Donovan B *et al.* Direct real-time PCR-based detection of Neisseria gonorrhoeae 23S rRNA mutations associated with azithromycin resistance. J Antimicrob Chemother 2015 doi:10.1093/jac/dkv274.

This study reports on an assay to identify mutations associated with azithromycin resistance. Crossreactivity remained a problem for detection in pharyngeal specimens despite the use of "mismatched" primer sequences.

[22] Fayemiwo SA, Muller EE, Gumede L, Lewis DA. Plasmid-mediated penicillin and tetracycline resistance among Neisseria gonorrhoeae isolates in South Africa: prevalence, detection and typing using a novel molecular assay. Sex Transm Dis 2011; 38:329-333.

[23] Gose S, Nguyen D, Lowenberg D *et al*. Neisseria gonorrhoeae and extended-spectrum cephalosporins in California: surveillance and molecular detection of mosaic penA. BMC Infect Dis 2013; 13:570.

[24] Goire N, Lahra MM, Ohnishi M *et al.* Polymerase chain reaction-based screening for the ceftriaxone-resistant Neisseria gonorrhoeae F89 strain. Euro Surveill 2013; 18:20444.

[25] Ochiai S, Ishiko H, Yasuda M, Deguchi T. Rapid detection of the mosaic structure of the Neisseria gonorrhoeae penA Gene, which is associated with decreased susceptibilities to oral cephalosporins. J Clin Microbiol 2008; 46:1804-1810.

[26] Nicol M, Whiley D, Nulsen M, Bromhead C. Direct detection of markers associated with Neisseria gonorrhoeae antimicrobial resistance in New Zealand using residual DNA from the Cobas 4800 CT/NG NAAT assay. Sex Transm Infect 2014; 91:91-93.

This study describes a multiplex assay applied to clinical specimens, but lacks validation against a culture reference standard.

*[27] Balashov S, Mordechai E, Adelson ME, Gygax SE. Multiplex bead suspension array for screening *Neisseria gonorrhoeae* antibiotic resistance genetic determinants in noncultured clinical samples. J This study describes a multiplex assay applied to clinical specimens, but lacks validation against a culture reference standard.Molec Diagn 2013; 15:116-129.

[28] Lawung R, Cherdtrakulkiat R, Charoenwatanachokchai A *et al.* One-step PCR for the
 identification of multiple antimicrobial resistance in Neisseria gonorrhoeae. J Microbiol Meth 2009;
 77:323-325.

[29] Goire N, Ohnishi M, Limnios AE *et al.* Enhanced gonococcal antimicrobial surveillance in the era of ceftriaxone resistance: a real-time PCR assay for direct detection of the *Neisseria gonorrhoeae* H041 strain. J Antimicrob Chemother 2012; 67:902-905.

[30] Goire N, Freeman K, Tapsall JW *et al.* Enhancing gonococcal antimicrobial resistance surveillance: a real-time PCR assay for detection of penicillinase-producing *Neisseria gonorrhoeae* by use of noncultured clinical samples. J Clin Microbiol 2011; 49:513-518.

[31] Siedner MJ, Pandori M, Castro L *et al.* Real-time PCR assay for detection of quinolone-resistant Neisseria gonorrhoeae in urine samples. J Clin Microbiol 2007; 45:1250-1254.

[32] Magooa MP, Muller EE, Gumede L, Lewis DA. Determination of *Neisseria gonorrhoeae* susceptibility to ciprofloxacin in clinical specimens from men using a real-time PCR assay. Int J Antimicrob Agents 2013; 42:63-67.

[33] Li Z, Yokoi S, Kawamura Y *et al.* Rapid detection of quinolone resistance-associated gyrA mutations in *Neisseria gonorrhoeae* with a LightCycler. J Infect Chemother 2002; 8:145-150.

[34] Zhou W, Du W, Cao H *et al.* Detection of gyrA and parC mutations associated with ciprofloxacin resistance in Neisseria gonorrhoeae by use of oligonucleotide biochip technology. J Clin Microbiol 2004; 42:5819-5824.

[35] Zhao L, Zhao S. TaqMan real-time quantitative PCR assay for detection of fluoroquinoloneresistant Neisseria gonorrhoeae. Curr Microbiol 2012; 65:692-695.

[36] Dona V, Low N, Guilarte YN *et al.* Multiplex real-time pcr with high resolution melting analysis for detecting resistance mechanisms in Neisseria gonorrhoeae. Oral Presentation O05.3 World STI & HIV Congress 2015. Brisbane, Australia, 13-16 September 2015. Sex Transm Infect 2015; 91 Suppl 2:A35-A36.

[37] Hemarajata P, Yang S, Soge OO *et al.* Real-time PCR and melt curve analysis targeting gyrA gene for prediction of ciprofloxacin resistance in clinical *Neisseria gonorrhoeae* isolates. Oral Presentation O05.1 World STI & HIV Congress. Brisbane, Australia, 13-16 September 2015. Sex Transm Infect 2015; 91 Suppl 2:A35.

[38] Pond MJ, Hall C, Cole M *et al.* Diagnostic and clinical implications of genotypic fluoroquinolone susceptibility detection for *Neisseria gonorrhoeae*. Oral presentation O05.2. World STI & HIV Congress 2015. Brisbane, Australia, 13-16 September 2015. Sex Transm Infect 2015; 91 Suppl 2:A35.

[39] British Association of Sexual Health and HIV Clinical Effectiveness Group, Bignell C, Ison C, FitzGerald M. United Kingdom National Guideline for Gonorrhoea Testing 2012. London: British Association of Sexual Health and HIV; 2012.

[40] Okeke IN, Peeling RW, Goossens H *et al.* Diagnostics as essential tools for containing antibacterial resistance. Drug Resist Updat 2011; 14:95-106.

[41] Cole M, Spiteri G, Chisholm S *et al.* Emerging cephalosporin and multidrug-resistant gonorrhoea in Europe. Euro Surveill 2014; 19. pii: 20955.

[42] Public Health England. GRASP 2013 Report. The Gonococcal Resistance to Antimicrobials Surveillance Programme (England and Wales). London: 2014; Public Health England.

[43] Kirkcaldy RD, Kidd S, Weinstock HS *et al.* Trends in antimicrobial resistance in Neisseria
 gonorrhoeae in the USA: the Gonococcal Isolate Surveillance Project (GISP), January 2006-June 2012.
 Sex Transm Infect 2013; 89 Suppl 4:iv5-10.

*[44] Speers DJ, Fisk RE, Goire N, Mak DB. Non-culture Neisseria gonorrhoeae molecular penicillinase production surveillance demonstrates the long-term success of empirical dual therapy and informs gonorrhoea management guidelines in a highly endemic setting. J Antimicrob Chemother 2014; 69:1243-1247.

This study is the first to show how an assay to detect antimicrobial resistance in *N. gonorrhoeae* was used in surveillance to change treatment guidelines.

[45] Ng LK, Martin I, Liu G, Bryden L. Mutation in 23S rRNA associated with macrolide resistance in Neisseria gonorrhoeae. Antimicrob Agents Chemother 2002; 46:3020-3025.

[46] Yang Y, Liao M, Gu WM *et al.* Antimicrobial susceptibility and molecular determinants of quinolone resistance in Neisseria gonorrhoeae isolates from Shanghai. J Antimicrob Chemother 2006; 58:868-872.

[47] Pai NP, Vadnais C, Denkinger C *et al.* Point-of-care testing for infectious diseases: diversity, complexity, and barriers in low- and middle-income countries. PLoS Med 2012; 9:e1001306.

[48] NESTA. Longitude Prize. https://longitudeprize.org/. London: 2014. (Accessed 19.10.2015)

[49] Gaydos CA. Review of use of a new rapid real-time PCR, the Cepheid GeneXpert(R) (Xpert) CT/NG assay, for Chlamydia trachomatis and Neisseria gonorrhoeae: results for patients while in a clinical setting. Exp Rev Molec Diagn 2014; 14:135-137.

[50] Kugelman G, Tapsall JW, Goire N *et al.* Simple, Rapid, and Inexpensive Detection of Neisseria gonorrhoeae Resistance Mechanisms Using Heat-Denatured Isolates and SYBR Green-Based Real-Time PCR. Antimicrob Agents Chemother 2009; 53:4211-4216.

[51] Chin CD, Laksanasopin T, Cheung YK *et al.* Microfluidics-based diagnostics of infectious diseases in the developing world. Nat Med 2011; 17:1015-1019.

[52] Peeling RW, Smith PG, Bossuyt PM. A guide for diagnostic evaluations. Nat Rev Microbiol. 2008;6:S2-S6.

[53] Pandori M, Barry PM, Wu A *et al.* Mosaic penicillin-binding protein 2 in Neisseria gonorrhoeae
isolates collected in 2008 in San Francisco, California. Antimicrob Agents Chemother 2009; 53:40324034.

[54] Chisholm SA, Dave J, Ison CA. High-level azithromycin resistance occurs in Neisseria gonorrhoeae as a result of a single point mutation in the 23S rRNA genes. Antimicrob Agents Chemother 2010; 54:3812-3816.

[55] Jerse AE, Bash MC, Russell MW. Vaccines against gonorrhea: current status and future challenges. Vaccine 2014; 32:1579-1587.