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Antimicrobial Resistance in *Neisseria gonorrhoeae*: Proceedings of the STAR Sexually Transmitted Infection—Clinical Trial Group Programmatic Meeting

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Abstract: The goal of the Sexually Transmitted Infection Clinical Trial Group’s Antimicrobial Resistance (AMR) in *Neisseria gonorrhoeae* (NG) meeting was to assemble experts from academia, government, nonprofit and industry to discuss the current state of research, gaps and challenges in research and technology and priorities and new directions to address the continued emergence of multidrug-resistant NG infections. Topics discussed at the meeting, which will be the focus of this article, include AMR NG global surveillance initiatives, the use of whole genome sequencing and bioinformatics to understand mutations associated with AMR, mechanisms of AMR, and novel antibiotics, vaccines and other methods to treat AMR NG. Key points highlighted during the meeting include: (i) US and International surveillance programs to understand AMR in NG; (ii) the US National Strategy for combating antimicrobial-resistant bacteria; (iii) surveillance needs, challenges, and novel technologies; (iv) plasmid-mediated and chromosomally mediated mechanisms of AMR in NG; (v) novel therapeutic (eg, sialic acid analogs, factor H [FH]/Fc fusion molecule, monoclonal antibodies, topoisomerase inhibitors, fluoroquinolones, LpxC inhibitors) and preventative (eg, peptide mimic) strategies to combat infection. The way forward will require renewed political will, new funding initiatives, and collaborations across academic and commercial research and public health programs.

Antimicrobial resistance (AMR) in *Neisseria gonorrhoeae* (NG) continues to be a serious threat to global public health. Although the use of dual antimicrobial therapy is highly effective, increasing reports of NG infections with cephalosporin- and azithromycin (AZI)-reduced susceptibility raise serious concerns regarding the durability of current treatment recommendations. Although nearly 400,000 NG cases were reported in 2015, the United States Centers for Disease Control and Prevention (CDC) estimates about 820,000 total infections occur annually due to the underreporting of asymptomatic undetected cases. Although
AMR in NG continues to be a concern both in the United States and globally, current nucleic acid-based amplification testing methods cannot measure antimicrobial susceptibility. Therefore, enhanced molecular diagnostics that distinguish among NG infections with antimicrobial resistance versus reduced susceptibility versus susceptibility are needed to help guide antibiotic treatment. The development and use of new bioinformatic tools, in conjunction with new technologies like whole genome sequencing (WGS) methods to identify AMR NG-associated mutations may resolve this issue at a global level. Understanding the mechanisms of AMR in NG may also help guide the development of new treatment and preventative modalities. The STI Treatment and Research (STAR) Sexually Transmitted Infection Clinical Trial Group (STI-CTG) held a programmatic meeting in Silver Spring, Maryland on April 13, 2017, titled: “Antimicrobial Resistance (AMR) in Neisseria gonorrhoeae (NG).” Experts from academia, government, nonprofit, and industry reviewed the current state of research, gaps and challenges in research and technology and future research and public health directions.

### SURVEILLANCE PROGRAMS TO UNDERSTAND AMR

#### The Sexually Transmitted Disease Surveillance Network

To complement routine notification data, CDC established the sexually transmitted disease (STD) Surveillance Network (SSuN) in 2005. In select jurisdictions, laboratory test results are collected from STD clinic attendees along with epidemiological data from a random sample of persons with gonorrhea. Data are representative of NG testing of the rectum, urethra, cervix, and pharynx. In some SSuN jurisdictions, more than 50% of reported gonorrhea cases occurred among men who have sex with men (MSM) in 2015. In other jurisdictions, cases in women and heterosexual men were more common, suggesting epidemic differences that may require different prevention and control approaches. In STD clinics participating in SSuN, the NG positivity rate among MSM tested for gonorrhea was over 5% and was elevated among HIV-infected MSM (eg, ~17% of HIV-infected MSM tested had rectal gonorrhea).

#### Gonococcal Isolate Surveillance Project

Established in 1986 to monitor N. gonorrhoeae antimicrobial susceptibility and inform treatment guidelines, Gonococcal Isolate Surveillance Project (GISP) is a collaboration between the CDC, clinical sites, and regional laboratories. Urthral specimens for culture and antimicrobial susceptibility testing are systematically collected from consecutive men with urethritis each month at participating STD clinics according to a standardized protocol; limited epidemiological data are locally abstracted from medical records and later merged, by CDC, with antimicrobial susceptibility data. Gonococcal Isolate Surveillance Project was designed for long-term surveillance of susceptibility trends; data are not available in a timely manner to inform clinical management and public health response. Although GISP is aimed at surveillance of NG in men, the Enhanced GISP (eGISP) was later created to strengthen surveillance of gonorrhea susceptibility and increase state and local capacity to detect and monitor NG in women and from extragenital sites.

During 2006 to 2016, the proportion of GISP isolates with reduced susceptibility (minimum inhibitory concentration [MIC], \( \geq 0.125 \mu g/mL \)) to ceftriaxone remained low (less than 0.5%). The proportion of isolates with reduced AZI susceptibility (MIC \( \geq 2.0 \mu g/mL \)) increased from 0.6% in 2013 to 3.6% in 2016. Recently, of particular concern, there were 4 GISP isolates collected in Hawaii that had elevated MICs to both AZI (MICs, \( \geq 16.0 \mu g/mL \)) and ceftriaxone (MICs, 0.125 \( \mu g/mL \)). Isolates collected through GISP continue to show reduced susceptibility to antimicrobials no longer recommended as first-line regimens; preliminary data for 2016 indicate that approximately 40% isolates had some resistance to penicillin, tetracycline, and ciprofloxacin.

Based on the approximately 820,000 gonococcal infections that occur each year in the United States, it was predicted that in 2011 about 246,000 infections either were resistant or had decreased susceptibility to at least 1 antibiotic; 11,480 had reduced susceptibility to ceftizime (MIC, 0.25 \( \mu g/mL \)), 2460 reduced susceptibility to AZI (MIC \( \geq 2.0 \mu g/mL \)), and 3280 reduced susceptibility to ceftriaxone. NG isolates with decreased susceptibility to cephalosporins are often resistant to other classes of antibiotics as well. Although these susceptibility trends are concerning, it is important to note that there have been no clinical treatment failures in the United States with the current recommended therapy of 250-mg ceftriaxone and 1-g AZI.

#### International Gonococcal Antimicrobial Surveillance Program

To support international surveillance of gonococcal resistance, the World Health Organization (WHO) founded Gonococcal Antimicrobial Surveillance Program (GASP) in 1990. Gonococcal Antimicrobial Surveillance Program currently has participating countries in Africa, the Americas, the Eastern Mediterranean, Europe, South East Asia, and Western Pacific. Different countries have different approaches to AMR in NG.

From 2009 to 2014, the total number countries reporting to GASP increased from 56 to 77, but there was considerable variation between WHO regions reporting. Of the 77 countries reporting to GASP, 66% reported isolates with any resistance/decreased susceptibility of NG to ceftriaxone (cefixime or ceftriaxone), 81% with any resistance/decreased susceptibility of NG to AZI, and 97% with any resistance/decreased susceptibility of NG to ciprofloxacin, for at least 1 year from 2009 to 2014. Notably, there are large gaps in data on AMR NG in Africa, Central America (extending up to Mexico), and the Middle East with adjacent countries in Asia. Currently, differences in the US and European guidelines for MIC interpretation create challenges to combining disparate country reports. Hence, as countries continue to develop robust surveillance programs and report MIC values, strategies to combine and compare such data need to be further examined and standardized.

Multidrug resistant (MDR) and extensively drug-resistant (XDR) forms of NG have been identified globally, including isolates from Japan, Hawaii, and England. The WHO defines MDR-NG as isolates with reduced susceptibility or resistance to either extended spectrum cephalosporins (ESC) or spectinomycin (ie, category I antibiotics), plus 2 or more of macrolides, fluoroquinolones, penicillins, tetracycline, aminoglycosides, and carbapenems (ie, category II antibiotics). XDR-NG are defined as isolates with decreased susceptibility or resistance to category I antibiotics and 3 or more category II antibiotics. Resistance or reduced susceptibility to cephalosporins (ie, ceftriaxone) define to the emergence of strains with mosaic penA alleles has been noted in the aforementioned countries. Reduced susceptibility to macrolides, such as AZI, has also been noted.

Questions persist about how to either implement or enhance surveillance, especially in low- and middle-income countries, how best to report AMR in NG, what the cost/benefit is for validating treatment failures in low- and middle-income countries, whether older antibiotics can be used again with new molecular diagnostics to predict susceptibility and how to validate various treatment guidelines from around the world. Treatment guidelines.
for NG must also be updated based on in-country surveillance data because many countries continue to use ciprofloxacin as a recommended first-line therapy.22 One success of the GASP program is the implementation of the National Strategy for Combating Antibiotic-Resistant Bacteria (NSAR). 

**US National Strategy for Combating Antimicrobial-resistant Bacteria**

The US National Strategy for Combating Antibiotic-Resistant Bacteria, released in September of 2014, sets forth 5 overarching goals: slowing the development of resistant bacteria and preventing spread of resistant infections; strengthening surveillance; advancing the development and use of rapid and innovative diagnostics; accelerating research and development for new antibiotics, therapeutics, and vaccines; and improving international collaboration.

In fiscal year 2017, congress appropriated $167 million to support implementation of the National Strategy through CDC’s Antibiotic Resistance Solutions Initiative. Although the initiative is broad-based, multiple activities focusing on NG are included, selected activities are described below. To strengthen surveillance, the Antibiotic Resistance Laboratory Network was created. Seven state public health laboratories serve as regional laboratories to conduct AMR testing of multiple pathogens and specialized testing of clinical specimens. Four of the laboratories conduct agar dilution testing of NG for GISP and other enhanced surveillance platforms. Integration of WGS of NG is planned.

Using Antibiotic Resistance Solutions Initiative funding, CDC also implemented the Strengthening US Response to Resistant Gonorrhea (SURRG), a collaboration between CDC and participating jurisdictions to establish local capacity to rapidly detect and respond to AMR in selected local jurisdictions. Jurisdictions participating in SURRG collect specimens for NG culture in STD clinics and other health care settings conduct rapid susceptibility testing on all isolates, interview patients infected with strains with reduced antimicrobial susceptibility and their recent contacts, and expand data collection to facilitate epidemicological and network analyses. The Antibiotic Resistance Solutions Initiative funding is also strengthening surveillance of NG isolates for drug susceptibility patterns in GISP and monitoring of trends of gonorrhea in SSuN.

**World Health Organization**

In response to the increasing threat of AMR, the World Health Organization (WHO) adopted a global action plan on antimicrobial resistance in May of 2015. The WHO’s 5 objectives are: (i) to improve awareness and understanding of antimicrobial resistance through effective communication, education, and training; (ii) to strengthen the knowledge and evidence base through surveillance and research; (iii) to reduce the incidence of infection through effective sanitation, hygiene, and infection prevention measures; (iv) to optimize the use of antimicrobial medicines in human and animal health; (iv) to develop the economic case for sustainable investment that takes account of the needs of all countries; and (v) to increase investment in new medicines, diagnostic tools, vaccines, and other interventions. This action plan emphasizes the need for a coordinated approach leveraging international stakeholders from different disciplines and sectors.

**SURVEILLANCE NEEDS, CHALLENGES, AND NOVEL TECHNOLOGIES**

**Surveillance Needs and Challenges**

From 2000 to 2010, global antibiotic usage increased by 36% and use of cephalosporins doubled, particularly in China and India. Such increases in antibiotic use likely drive the selective pressure for AMR. Although global surveillance efforts (eg, GISP/GASP) strive to detect AMR in NG isolates, one must consider whether those data are being collected with sufficient timeliness to mitigate the risks. Through current surveillance programs, there is a lag in identifying AMR NG for clinical decision making, thereby potentially enabling continued transmission of AMR strains. The process is limited by the constraints of current methods and technologies in growing NG isolates, identifying AMR and its mechanisms, and identifying isolates implicated in AMR outbreaks through phenotypic characterization. Greater use of molecular tools for timely and accurate detection of AMR as are applied on other fields of infectious disease surveillance, including PCR and DNA sequencing, is urgently needed.

Cases of ceftriaxone-reduced susceptible NG have been found in pharyngeal specimens.21,22 Pharyngeal gonorrhea poses multiple challenges due to its asymptomatic nature, ease of transmission and difficulty of treatment. The pharynx may also serve as an NG reservoir and incubator of reduced susceptibility because of the frequent presence of commensal nonpathogenic Neisseria species. Given that Neisseria species are known for DNA uptake and exchange, it is likely that the horizontal transfer of genetic material, including antibiotic resistance genes, in the pharynx leads to AMR NG infections.

In many regions globally, antibiotics are readily available without a prescription, and those regions are historically known for high levels of antibiotic resistance and have groups of people with high rates of oropharyngeal STIs. Given that environment, the NIH-funded (Fogarty Center) ICON Study in northern Vietnam, enrolled MSM to address the frequency of antibiotic usage and any association with antibiotic-resistant or -susceptible NG. Preliminary results of the ICON Study found 62% of current participants reported antibiotic usage in the prior 6 months, often without a prescription and some stopped antibiotic usage as soon as symptoms abated. Nonpathogenic Neisseria were found in 38 (100%) of 38 clinical pharyngeal specimens, with some samples growing up to 4 different Neisseria species, including N. gonorrhoeae and N. meningitidis. Next steps of the ICON Study include determining whether different Neisseria species have different capacities for acquiring resistance, determining the prevalence of similar genetic components in different resistant strains, and whether Neisseria commensals can be used in surveillance to predict trends in NG AMR.

**Novel Technologies**

Advances in genomics might help address AMR through informing the development of molecular diagnostics, identifying outbreaks, advancing the understanding of disease transmission, and through epidemiological/evolutionary inference to guide antibiotic selection. Important AMR-related questions that genomics can help address include the following: (i) How much resistance...
is due to clonal spread and de novo emergence? (ii) To what extent do known genetic resistance mechanisms explain observed pheno-
type resistance? (iii) How can the scientific community identify novel mechanisms of resistance?

Reports have shown that increased MICs to extended-
spectrum cephalosporins (ceftriaxome MIC, ≥0.25 μg/mL; ceftriaxone MIC, ≥0.125 μg/mL) in the United States is predominantly associ-
ated with the mosaic penA XXXIV allele with or without additional
specific point mutations in penA.39-31 Quinolone-resistant NG has
widely spread through predominantly spread of mutations in gyrA
and parC (gyrA-S91F/I, gyrA/D59G, parC-S88P). Reduced AZI
susceptibility has arisen through multiple mechanisms, with the
most common in the United States being 23S RNA mutations
(C251T, and A2059G) and mosaic mtr mutations in the mtrR lo-
cus.33,34 However, about a third of reduced AZI susceptibility
(MIC ≥2 μg/mL) is not clearly explained by 23S RNA mutations,
by a mosaic mtr locus, by a single basepair deletion in the mtrR
promoter or generation of a new promoter for transcription of
mtrCDE.34 Those findings indicate the utility of WGS in develop-
ing nucleotide-based molecular diagnostics. However, several lim-
itations are worth noting. First, not all phenotypic resistance is
explained by known mechanisms of resistance. Further, the fre-
cuency with which novel mechanisms of resistance arise, mixed
strain infections occur or how to best screen for such mechanisms,
or determine the clinical impact of mixed infections is unclear.

Genomic epidemiology can help understand patterns of
spread of gonococcal strains and identify local transmission and
outbreaks. Examples include tracking the transmission of resistant
lineages across geographic and demographic boundaries33-36 and
reconstructing local transmission networks.35,36

Development of point-of-care (POC) diagnostics to iden-
tify drug susceptibility profiles has the potential to impact overall
levels of AMR and, as 60% of gonococcal isolates in the United
States are pan-susceptible, permit reintroduction of older antibio-
tics into treatment regimens.37-42 Although a rapid test for sus-
cceptibility is expected to aid in reducing the overall burden of
AMR as compared with one that does not detect susceptibility,43
questions remain about how best to deploy these strategies.

**MECHANISMS OF ANTIMICROBIAL RESISTANCE
(AMR) IN NG, AND NOVEL ANTIBIOTICS AND
VACCINES TO TREAT AMR NG**

**Mechanisms of β-lactam Antibiotic Resistance
in NG**

There are 2 genetic sources of antibiotic resistance in NG:
plasmid-mediated and chromosomally mediated. In plasmid-
mediated resistance, β-lactam antibiotic resistance occurs due to
the expression of an antibiotic modifying protein (eg, TEM-1–
like β-lactamase for penicillin and amino-penicillin (eg, amoxicil-
in) resistance [PenK]) or a ribosome-protected protein (TetM
ribosomal-binding protein that confers tetracycline resistance).
β-lactamase does not hydrolyze cephalosporins, so it does not
contribute to cephalosporin resistance. However, 1 amino acid
change in the bla gene could convert it to produce extended-
spectrum β-lactamase.44 In chromosomal-mediated resistance,
bacterial resistance occurs due to de novo spontaneous mutations
or due to the acquisition of chromosomal mutations via homolo-
gous recombination commonly thought to occur from Neisseria
commensal species. In stepwise resistance, each step is a relatively
small increase in resistance, but when multiplied overall, it leads to
a large increase in the MIC of a given antimicrobial.

The main difference between PenK NG strains and
cephalosporin-resistant (CephK) cephalosporin intermediate-resistant
strains (CephK) is due to the type of mosaic penA allele arising from
interspecies recombination. It appears that the origin and rapid
emergence of CephK strains was due to a single transformation
event of a mosaic penA allele into existing CephK/PenK strains,
which to this day persist even though penicillin has not been used
for NG treatment in decades.45

In addition to the penA allele, the mtrR and penB determin-
ants contribute additional resistance to β-lactam antibiotics and
provide a general permeability barrier for antibiotics. The mtrR
determinant, caused by mutations either in the promoter region
or coding sequence, increases transcription of the divergently tran-
scribed mtrCDE operon that encodes the MtrC-MtrD-MtrE efflux pump.47-49
The increased expression of the pump causes increased efflux of antibiotics from the cytoplasm and periplasm
of NG. The mutated penB may produce altered forms of the
PorB Porin, the major porin of NG,50,51 resulting in a decrease
in the influx of antimicrobials through the porin channels. It is in-
teresting that the increase in resistance conferred by penB requires
the presence of an mtrR mutation.52

**Novel Therapeutic and Vaccine Approaches**

Novel, nontraditional therapeutic, and vaccine approaches
to combat MDR NG infection are currently being investigated.
Therapeutic approaches include sialic acid analogs (eg, chemical
therapies),53-55 FH/Fc fusion molecules43-56 and monoclonal anti-
bodys (eg, immunotherapeutic molecules). Vaccine approaches
include widely expressed antigens that are immunogenic (eg,
common lipooligosaccharide epitopes represented by peptide
mimics),57-59 and the use of vaccines developed for other Neisseria
species that may cross-protect against NG infections.

Nongonococcal sialic acids can be used to disrupt the
natural protection on most gonococcal organisms. More specif-
ically, endogenous, host mammalian sialic acids are taken up by
gonococci in vivo and result in protection of the organism from
complement-mediated killing whereas nonhost sialic acids, de-
derived from alternative sources, do not possess this protective
function (complement resistance). With respect to mechanism
of action, when alternative sialic acids are administered locally
to infected mice, they replace host sialic acid, can be taken up
preferentially by gonococci, and hasten clearance of bacteria
by removing resistance to complement-mediated killing.53-55
Natural and synthetic sialic acids can be mined for candidates
that are optimal in eliminating complement resistance and hasten-
ing clearance of infecting bacteria.

A fusion protein has been engineered that on the one hand
binds to a complement regulator binding site, present on all gono-
cocci, called factor H, and, on the other hand, possesses an Fc
domain that engages complement and kills the organism; thereby,
enhancing clearance in the animal model. The FH portion has
been altered so as not to bind to human cells thereby avoiding tox-
icity. The FH/Fc fusion protein constructs have been shown to
bind to 12 of 15 different gonococcal isolates, kill 10 of 15 of these
in vitro and hasten clearance of 3 different isolates infecting the
animal model.54-56 Production of FH/Fc, a fully humanized im-
munotherapeutic, is being scaled up in tobacco plants and config-
ured for use parenterally and in intravaginal release devices.

Another immunotherapeutic molecule being developed for
gonorrahoea treatment is the chimeric (mouse/human) 2C7 antibody.
The 2C7 antibody has been tested, intravaginally and parenterally,
in the mouse animal model.57 The 2C7 antibody is being fully hu-
manized and like FH/Fc, production is being scaled up in tobacco
plants and configured for parenteral and intravaginal administra-
tion. Because 2C7 antibody and FH/Fc target different sites on
the organism, combining their use may be additive.
The 2C7 epitope, against which the 2C7 antibody was developed, forms the basis for a novel gonococcal vaccine. The 2C7 epitope is displayed by greater than 95% of clinical isolates; antibodies against the 2C7 epitope are elicited uniformly by women with infection. A 2C7 peptide mimic vaccine was constructed by screening of randomly generated peptides (using a peptide library consisting of > 10^7 peptides) and identifying peptide(s) recognized by 2C7 monoclonal antibody. A multiantigenic peptide (MAP; octameric/tetrameric) was fashioned that elicited antibodies directed against the nominal (2C7) epitope, possessed complement-dependent killing against all gonococcal isolates tested and hastened clearance of infection in vaccinated animals. Stabilization and scale-up of homogenous peptide (>95% pure) has already been accomplished and current work is aimed at optimizing responses to the peptide mimic vaccine with human-approved adjuvants. Although meningococcal group B outer membrane vesicle vaccines have been shown to be immunogenic and efficacious against homologous strains, more recently they have also been found to protect partially against NG infection. A retrospective case-control study of patients seen in New Zealand sexual health clinics revealed that exposure to the outer membrane vesicle meningococcal B vaccine was associated with about 30% reduction in gonorrhea diagnoses.

Although those novel therapeutic and preventive approaches provide hope in curtailing gonococcal infections, they will require more research and development to deliver an approved, affordable treatment for AMR NG that can be brought to the clinic. Meanwhile, there are few novel, more traditional antibiotic approaches that are in development.

**Novel Therapeutic for Uncomplicated NG:**

Zoliflodacin/ETX0914

The standard CDC and WHO treatment recommendation for gonorrhea requires a minimal efficacy of greater than 95% at any mucosal site (cervix, urine, rectum, pharynx). An optimal treatment would be effective against resistant isolates for both urogenital and extragenital infection and would be well tolerated. While a single-dose therapy would be ideal, single-dose therapy versus multidose therapy is less a priority than a safe and well-tolerated antimicrobial regimen with efficacy across resistant isolates and all anatomic sites. Zoliflodacin (Entasis Therapeutics) was developed for the treatment of uncomplicated gonorrhea and is the first drug in a novel class of topoisomerase inhibitors. Zoliflodacin has shown potent in vitro activity against 100 gonococcal isolates and shows a lack of cross-resistance to other antibiotic classes. Because its mechanism of action is distinct from fluoroquinolones, it is hypothesized that zoliflodacin will be effective in treating fluoroquinolone-resistant infections. In phase 1 studies, a single dose of zoliflodacin was well tolerated in healthy adult males, and all adverse events were mild/nonserious. No adverse events lead to study discontinuation. Zoliflodacin was conducted to assess safety and microbiological cure among 180 subjects with gonorrhea. Of the 180 subjects enrolled, 131 were analyzed as microbiological-intent-to-treat evaluable subjects. The number of participants with microbiological cure at urethral or cervical sites in the 2000-mg zoliflodacin, 3000-mg zoliflodacin, and the 500-mg intramuscular ceftriaxone group were 55 of 57, 54 of 56, and 28 of 28, respectively. Among 15 patients across the 3 groups with rectal infections, all were cured. The number of patients with microbiological cure at pharyngeal site was slightly higher in the patients treated with intramuscular ceftriaxone (4 of 4) compared with the 3000 mg zoliflodacin group (9 of 11). Overall, zoliflodacin was well tolerated. Phase 3 studies of zoliflodacin are currently being planned with support from the Global Antibiotic Research Development Program.

**Solithromycin**

Solithromycin (Cempra, Inc.) is a 4th-generation macrolide and the first fluoketolide. It exhibits in vitro activity against a number of urogenital pathogens including NG, Chlamydia trachomatis, Mycoplasma spp., and Ureaplasma spp. Solithromycin was tested in a phase 2 urethritis study to assess the eradication of urogenital NG and in a phase 3 study to assess its noninferiority versus intramuscular ceftriaxone plus oral AZI. Solithromycin was associated with gastrointestinal-related adverse events. A follow-up phase 3 study of Solithromycin patients with a positive baseline culture who returned for evaluation at the test of cure visit, treatment success was 91.3% (95 of 104) for solithromycin recipients, versus 100% (107 of 107) for CTX/AZI patients. Among the 9 solithromycin patients with a positive TOC culture result, there was no correlation between outcome and solithromycin MIC range, 0.004–0.25 μg/mL; all baseline isolates were susceptible to solithromycin using CDC criteria for AZI (MIC <2.0 μg/mL). Emergence of solithromycin resistance in TOC isolates was not observed. Genotyping of pretreatment and posttreatment isolates did not demonstrate reinfection with novel strains. Given the absence of baseline or acquired solithromycin resistance and the absence of evidence of reinfection with a novel strain, the investigators surmised that the most likely cause of treatment failure was pharmacokinetic-related, with presumed insufficient duration of drug exposure at the site of infection. It is hypothesized that solithromycin dose adjustment (for instance, a 2-dose strategy, over 24 hours) and combination treatment strategy with a second antibiotic would result in desired treatment success rates. Although other novel therapeutics for NG are currently being investigated, including the triazacenaphthylene antibiotic agent, gepotidacin, these were not discussed at the programmatic meeting.

**OTHER MEANS TO TREAT GONOCOCCAL INFECTION**

**Crippling Selective Gene Expression**

Understanding the mechanisms of AMR in NG can assist in the design of newer antimicrobials and vaccines and provide insights as to the development of compensatory mutations that reverse fitness defects yet maintain resistance. The MtrCDE drug efflux contributes significantly to such resistance and transcriptional control systems modulate levels of efflux pump gene expressions and, as a consequence, levels of antibiotic resistance. Mutations that increase efflux pump gene expression can adversely impact clinical efficacy of antibiotics. Finally, dampening efflux pump gene expression might allow for return of an old antibiotic (eg, penicillin) or allow for continued use of current antibiotics.
Strategy to Alter Bacterial Membranes—Lipid A Enzymes

Lipid A is a component of bacterial outer membranes and is essential for cell viability of nearly all Gram-negative bacteria. Current investigations are aimed at evaluating whether small molecule inhibitors (eg, TU-514, CHIR-090, LPC-067) of LpxC, an essential gene for NG, can be used to target NG infections. LpxC Inhibitors (eg, LPC-169, LPC-174, LPC-201, LPC-211) have been shown to overcome existing antibiotic resistance (unpublished data). The investigators also looked at the efficacy of LPC-211 in mouse models against a specific ceftriaxone-resistant strain of NG. Although the investigators have found such inhibitors to work well to treat NG, improvements are still needed to file an IND application.

NEXT STEPS: RESEARCH AND TECHNOLOGY GAPS AND CHALLENGES

AMR NG Research Gaps and Challenges

In 2016, 7 patients in Hawaii were found to be infected with strains demonstrating high-level AZI resistance (MICs, ≥16.0 μg/mL) and elevated MICs to ceftriaxone (MICs, 0.125 μg/mL). Although those are rare, the chances of combined AZI and ceftriaxone-reduced susceptibility are growing. A recent report from China found about 3% of NG isolates with dual ceftriaxone decreased susceptibility and AZI resistance. Molecular studies have found that there is considerable variability in the mutations associated with AZI-reduced susceptibility. An important question to consider is how the scientific community can best monitor reduced susceptibility to AZI in regions across the globe. One strategy may be to increase AMR surveillance programs, like GISP/GASP, globally and expand the collection of nonurogenital specimens. Previously, gonorrhea was treated using antimicrobial monotherapy; specific antimicrobials were recommended based on clinical trial results and subsequent antimicrobial susceptibility trends. The use of dual therapy potentially introduces more complexity into decisions about treatment recommendations. The value of dual therapy to prevent AMR is only a theoretical argument at present; investigations of whether using 2 or more antibiotics at one time slows the development of resistance to either drug would advance the field. To that end, murine modeling studies may play an important role in addressing such questions in addition to understanding host-microbe interactions. Creating antimicrobial susceptibility testing matrices that include different doses for each drug may help to determine if the combination of drugs are synergistic or antagonistic and may help to address the aforementioned question of resistance. Although several synergy studies of drugs against NG have been published, little to no antagonism or synergy has been noted. As new antimicrobial agents, such as those discussed previously, and diagnostics become commercially available in coming years, questions about how to select the most effective drug combinations, weighing both clinical efficacy and impact on resistance, should be addressed with additional synergy studies.

Syndromic Management and AMR NG

Syndromic management continues to be the principal approach for STI treatment in low- to middle-income countries because of its simplicity and affordability. Syndromic management is based on the identification of clinical symptoms (or signs) with resultant indications for treatment rather than making an etiological diagnosis using laboratory methods. Although inexpensive and fast, the shortcomings of syndromic management include a lack of specificity and substantial overuse of antibiotics. Syndromic management may greatly contribute to AMR in NG. Another problem with syndromic management is that it does not address those with asymptomatic infections and is therefore unlikely to impact the burden of infection. Implementing rapid POC detection of NG as a first step in the diagnosis of gonorrhoea and potentially even more valuable the POC detection of NG with specific antimicrobial susceptibility could profoundly impact and slow the emergence of AMR in NG.

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