Genotypic and Phenotypic Characterization of Antimicrobial Resistance in \textit{Neisseria gonorrhoeae}: a Cross-Sectional Study of Isolates Recovered from Routine Urine Cultures in a High-Incidence Setting

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\textbf{ABSTRACT} The objectives of this study were to perform genomic and phenotypic characterization of antimicrobial resistance in \textit{Neisseria gonorrhoeae} isolates recovered from urine samples from patients in St. Louis, MO, USA. Sixty-four clinical isolates were banked over a 2-year period and subjected to antimicrobial susceptibility testing (AST) by Kirby-Bauer disk diffusion (penicillin, tetracycline, cefuroxime, and ciprofloxacin) and gradient diffusion (tetracycline, doxycycline, azithromycin, ceftriaxone, cefixime, ciprofloxacin, gemifloxacin, and delafloxacin). The medical records for the patients were evaluated to determine the demographics, location, and prescribed treatment regimen. Isolate draft genomes were assembled from Illumina shotgun sequencing data, and resistance determinants were identified by ResFinder and PointFinder. Of the 64 isolates, 97\% were nonsusceptible to penicillin, with resistant isolates all containing the \textit{bla}_{TEM-1b} gene; 78 and 81\% of isolates were nonsusceptible to tetracycline and doxycycline, respectively, with resistant isolates all containing the \textit{tet(M)} gene. One isolate was classified as non-wild-type to azithromycin, and all isolates were susceptible to ceftriaxone; 89\% of patients received this combination of drugs as first-line therapy. Six percent of isolates were resistant to ciprofloxacin, with most resistant isolates containing multiple \textit{gyrA} and \textit{parC} mutations. Correlation between disk and gradient diffusion AST devices was high for tetracycline and ciprofloxacin ($R^2 > 99\%$ for both). The rates of \textit{N. gonorrhoeae} antibiotic resistance in St. Louis are comparable to current rates reported nationally, except ciprofloxacin resistance was less common in our cohort. Strong associations between specific genetic markers and phenotypic susceptibility testing hold promise for the utility of genotype-based diagnostic assays to guide directed antibiotic therapy.

\textbf{IMPORTANCE} \textit{Neisseria gonorrhoeae} causes the sexually transmitted infection gonorrhea, which is most commonly diagnosed using a DNA-based detection method that does not require growth and isolation of \textit{N. gonorrhoeae} in the laboratory. This is problematic because the rates of antibiotic resistance in \textit{N. gonorrhoeae} are increasing, but without isolating the organism in the clinical laboratory, antibiotic susceptibility testing cannot be performed on strains recovered from clinical specimens. We observed an increase in the frequency of urine cultures growing \textit{N. gonorrhoeae} after we implemented a total laboratory automation system for culture in our clinical laboratory. Here, we report on the rates of resistance to multiple historically used, first-line, and potential future-use antibiotics for 64 \textit{N. gonorrhoeae} isolates. We
found that the rates of antibiotic resistance in our isolates were comparable to national rates. Additionally, resistance to specific antibiotics correlated closely with the presence of genetic resistance genes, suggesting that DNA-based tests could also be designed to guide antibiotic therapy for treating gonorrhea.

**KEYWORDS** Neisseria gonorrhoeae, gonorrhea, urine culture

*Neisseria gonorrhoeae*, the microorganism that causes the sexually transmitted disease gonorrhea, infects approximately one million people in the United States annually (1, 2). Historically, penicillin and doxycycline have been used to treat gonorrhea; however, the emergence of *N. gonorrhoeae* strains resistant to these antibiotics in the 1980s prompted the implementation of alternative treatment strategies. While the antimicrobial regimen of choice recommended in clinical guidelines has varied over the past decades, today, single-dose ceftriaxone plus azithromycin combination therapy is the standard of care for the treatment of gonorrhea in the United States and much of the world (3–7). With the advent of direct-from-specimen nucleic acid amplification testing (NAAT), gonorrhea can now be diagnosed and treated in a single clinic or emergency room encounter. NAAT is rapid, highly sensitive, and specific; as such, it has become the standard of care for diagnosing gonorrhea and has largely replaced culture-based diagnosis of *N. gonorrhoeae*, which is slow and suffers from poor sensitivity (8, 9). Importantly however, culture is a prerequisite for performing antimicrobial susceptibility testing (AST) on an organism, and recent trends suggest that *N. gonorrhoeae* is becoming increasingly resistant to antibiotics, including azithromycin and ceftriaxone (7, 10–14) (for a review, see reference 15).

Within the past 3 years, the clinical microbiology laboratory at Barnes-Jewish Hospital (affiliated with Washington University in St. Louis)—a central laboratory for 5 hospitals in a 13-hospital health care system—implemented a total laboratory automation system for culture-based microbiology (BD KiestraTLA; Becton-Dickinson). After these changes, the laboratory observed a dramatic increase in the number of routine urine cultures from which fastidious microorganisms were recovered (16). This included a large increase in routine urine cultures growing *N. gonorrhoeae*, which typically does not grow on blood agar medium under atmospheric oxygen tension commonly used for conventional urine culture (Fig. 1A). Here, we use whole-genome sequencing and AST to characterize the epidemiology and antimicrobial resistance patterns and establish genotype-phenotype correlations for 64 *N. gonorrhoeae* isolates from patients in the St. Louis area.

**RESULTS**

**Sample set and patient demographics.** Of 74 isolates initially identified as *N. gonorrhoeae* during routine workup by matrix-assisted laser desorption ionization—time of flight mass spectrometry (MALDI-TOF MS) (BioTyper) in the clinical microbiology laboratory at Barnes-Jewish Hospital between November 2015 and October 2017, 64 met all inclusion criteria for this study, including recovery from cryopreservation, confirmed identification as *N. gonorrhoeae* by MALDI-TOF MS (Vitek MS) after subculture, and complete patient records. Of note, there was 96% agreement between the two MALDI-TOF MS methods.

The majority of the isolates in our sample set came from young black men living in North St. Louis, MO. All were symptomatic with some degree of urogenital discomfort (e.g., “urethral discharge” or “dysuria”); urine cultures were performed as part of a general urogenital tract infection workup at the discretion of the treating physician. The mean age of patients was 27 years old (standard deviation [SD], 9.6 years), but ages ranged from 15 to 58 years of age (Table 1). Of 64 (93.8%) patients, 60 were male, and 59/64 (92.2%) of patients were black. Two black women, two white women, and three white men were also represented in our data set. Of 64 isolates, 54 (84.4%) isolates came from patients residing in zip codes from the predominantly African American community of North St. Louis (Fig. 1B). Isolates from other locations were rare and originated from patients scattered throughout more demographically heterogeneous
neighborhoods in South St. Louis (3/64 [4.7%]) or St. Louis suburbs in Illinois (5/64 [7.8%]).

Over the time frame when this study was conducted, *N. gonorrhoeae* was isolated from 0.064% of all urine cultures performed, representing 0.3% of all uropathogens recovered (16). NAAT was the predominant method of gonorrhea diagnosis, with 60/64 (93.8%) isolates having a corresponding NAAT test at the time of specimen collection; all were positive except for one that was "indeterminate." The converse was difficult to assess, given the high prevalence of gonorrhea within our laboratory’s catchment area and the superior sensitivity of NAAT over culture as a primary method for diagnosis. Therefore, we were unable to estimate the percentage of specimens that were positive by NAAT but negative by urine culture. Most patients (37/64 [57.8%]) had no concurrent sexually transmitted infections (STIs). However, concurrent infections (as documented via review of the electronic medical record) with *Chlamydia trachomatis* (18/64 [28.1%]), *Trichomonas vaginalis* (2/64 [3.1%]), HIV (2/64 [3.1%]), and hepatitis C virus (HCV) (1/64 [1.6%]) were identified in a subset of patients. At the time of presentation,

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**FIG 1** Recovery of *N. gonorrhoeae* from urine cultures in the St. Louis area. (A) Blood agar plate showing growth of *N. gonorrhoeae* from routine urine culture performed on a total laboratory automation system (the BD KiestraTLA). Colonies appear nonhemolytic, pin-point, and clear/gray/white in color. (B) Map of the St. Louis metropolitan area showing the primary residence of the patient from which each isolate was obtained at the zip code level or with an arrow indicating the direction of zip codes that are not on the map. Heat mapping highlights the number of patients per zip code, with darker red indicating more patients per zip code. BJH, Barnes-Jewish Hospital; SLCH, St. Louis Children’s Hospital.
57/64 (89%) individuals received standard-of-care antimicrobial therapy for gonorrhea: single-dose azithromycin (1 g) plus ceftriaxone (250 mg) (Table 1).

**Antimicrobial susceptibility testing of *N. gonorrhoeae* isolates.** Given recent reports of increasing antibiotic resistance among *N. gonorrhoeae* isolates, we evaluated resistance to several antibiotics in our sample set. We assessed antibiotics from the tetracycline, quinolone, beta-lactam, and macrolide classes that were either used historically to treat gonorrhea or are used currently as first-line agents for treating gonorrhea or have been proposed as future therapies for treating multidrug-resistant *N. gonorrhoeae* (17).

We first assessed the distribution of MICs (or zone diameters) displayed by *N. gonorrhoeae* isolates against each antibiotic. For penicillin, tetracycline, and doxycy-
cline, isolates formed a multimodal distribution of MICs (or zone diameters) that roughly corresponded to the susceptible (S), intermediate (I), and resistant (R) categories of antimicrobial susceptibility (Fig. 2A and B). A bimodal distribution of MICs was also observed for azithromycin, with a group of isolates straddling the lower bound of the clinical breakpoint. In contrast, the susceptibility of isolates to cephalosporins was distributed about a mean that resided deep within the susceptible category. This was also true for the quinolone class of antibiotics, with the exception of four outliers that displayed increased MICs to ciprofloxacin, gemifloxacin, and delafloxacin.

The greatest percentage of isolates with nonsusceptible phenotypes (i.e., resistant or intermediate) were found for penicillin (96.9%: 7.8% R, 89.0% I), doxycycline (81.0%), and tetracycline (40.6% by disk diffusion; 78.1% by gradient diffusion) (Fig. 2C). Isolates were highly susceptible to first-line therapy: only 1 isolate of 64 isolates (1.6%) was non-wild-type to azithromycin, and none of the 64 isolates (0%) were resistant to ceftriaxone (or any of the other cephalosporins tested). Among quinolone antibiotics, the highest rate of nonsusceptibility was seen for ciprofloxacin (5/64 [7.8%]), with four of the ciprofloxacin-resistant isolates also testing nonsusceptible to gemifloxacin (4/64 [6.25%]) and displaying increased MICs for delafloxacin. These four isolates also tested nonsusceptible to penicillin, doxycycline, and tetracycline, indicative of a multidrug-resistant phenotype (Fig. 2D).

We also examined categorical agreement between the disk and gradient diffusion methods for tetracycline and ciprofloxacin. Isolates displayed a wide range of susceptibilities to tetracycline, with zone diameter correlating strongly with MIC ($R^2 = 99\%$ using a least-squares nonlinear fit model; Fig. 3). However, this corresponded to only 40/64 isolates (62.5%) having categorical agreement, with the remaining 24/64 isolates (37.5%) testing susceptible by disk diffusion but intermediate by gradient diffusion. Nonetheless, there were zero major or very major errors for tetracycline disk versus gradient diffusion methods. Isolates with resistance to ciprofloxacin were uncommon; nonetheless, the correlation between zone diameter and MIC was very high ($R^2 = 99\%$), as was categorical agreement (100%).

**Genetic determinants of antimicrobial resistance in *N. gonorrhoeae* isolates.**

For most of the antibiotics we tested, the zone sizes (or MICs) were not normally distributed (as measured by the D’Agostino-Pearson omnibus normality test; Fig. 2A and B)—a potential indication of acquired resistance genes and/or mutations in a subset of isolates. To explore this observation further, we performed whole-genome sequencing on each isolate. The ResFinder and PointFinder databases were used to identify known genes and mutations (collectively referred to as “resistance determinants”), respectively. We detected several penicillin resistance determinants; however, only acquisition of the blaTEM-1b beta-lactamase gene was consistently associated with penicillin resistance and was found in 100% of penicillin-resistant isolates (Fig. 4A). As expected, the presence of blaTEM-1b was not associated with increased cephalosporin MICs (or reduced zone sizes), nor were any of the other resistance determinants we examined (Fig. 4B).

Within the tetracycline class, acquisition of tet(M) was highly associated with resistance to both tetracycline and doxycycline, with all five (100%) resistant isolates containing tet(M) (Fig. 4C). All five isolates resistant to tetracycline also had the rpsJ V57M mutation, but isolates containing the rpsJ mutation yet lacking tet(M) were not resistant.

Only one of five isolates with increased MICs to azithromycin contained an established multidrug resistance determinant (an mtrR promoter mutation), although this particular isolate still met the definition of “wild type.”

The genotype-phenotype correlation for the quinolone antibiotics was less straightforward. All four isolates with known resistance mutations in gyrA and parC were resistant to ciprofloxacin, with the exception of one isolate with a gyrA S91F mutation which was categorized as “intermediate” to ciprofloxacin (MIC = 0.125 μg/ml). This particular isolate contained only the gyrA S91F mutation whereas the other three isolates contained a gyrA D95G mutation in addition to the gyrA S91F mutation
FIG 2 Antimicrobial susceptibility of *N. gonorrhoeae* isolates. (A and B) Clinical isolates (*n* = 64) were subjected to AST via the disk diffusion method (A) or gradient diffusion (i.e., “Etest”) method (B). Zone-of-inhibition diameters (A) and MICs (B) are shown. Each black circle represents the value for one isolate, and the geometric mean (horizontal black bar) with 95% confidence interval (error bars) for the different treatment groups are shown. For isolates with MICs that were off the measurable scale, a value of 65 mm (disk) or the lowest value (gradient strip) was assigned. Throughout the figure, the breakpoints for each antibiotic are shown by colors: green, susceptible; yellow, intermediate; red, resistant (except for azithromycin, where green represents “wild-type” and red represents “non-wild type.” CLSI breakpoints were used with the following exceptions: doxycycline breakpoints were inferred from tetracycline; gemifloxacin breakpoints were inferred from ciprofloxacin; delafloxacin breakpoints were inferred from ciprofloxacin; and EUCAST (Continued on next page)
Accordingly, the isolate with the lone gyrA S91F mutation was also susceptible to gemifloxacin and delafloxacin, whereas the other isolates containing multiple quinolone resistance determinants tested "intermediate" to gemifloxacin and had increased MICs to delafloxacin. Finally, one isolate contained no known quinolone resistance determinants but repeatedly tested as resistant to ciprofloxacin with increased MICs to gemifloxacin and delafloxacin.

Phylogenetic analysis of \textit{N. gonorrhoeae} isolates. To examine relationships between isolates, we constructed a core genome phylogeny using PRANK and raxML on the 1,563 genes shared by the genomes collected in our investigation and the WHO reference strains (18, 19) (Fig. 5). Midpoint rooting of the phylogenetic tree showed that the collected isolates fell within three major clades. WHOF/WHOW strains were in the first cluster, WHOO strain was in the second cluster, and the remaining WHO strains were in the third cluster. To identify putative transmission clusters, we labeled each isolate with its corresponding zip code; however, transmission clusters were not identified. However, we observed associations between resistance determinants and position of the isolates on the phylogenetic tree. \textit{bla}\textsubscript{TEM-1b} was located only in isolates within the first cluster, and \textit{tet}(M) was found only in the second cluster.

DISCUSSION

Because NAAT is the standard of care for diagnosis of \textit{N. gonorrhoeae}, clinical isolates for detailed phenotypic and genotypic characterization are scarce. New culture methods facilitated enhanced recovery of \textit{N. gonorrhoeae} isolates in our laboratory, and subsequently, we characterized 64 patient isolates, including analysis of demography, geography, AST, and whole-genome sequencing.

As expected, the rates of antibiotic resistance to historically used antibiotics such as penicillin and doxycycline were high and were correlated with the presence of the \textit{bla}\textsubscript{TEM-1b} and \textit{tet}(M) genes, respectively. However, almost no resistance to current first-line antibiotics, including azithromycin or drugs in the cephalosporin class was detected, consistent with low rates of resistance to first-line antibiotics reported (Fig. 4D). Accordingly, the isolate with the lone gyrA S91F mutation was also susceptible to gemifloxacin and delafloxacin, whereas the other isolates containing multiple quinolone resistance determinants tested "intermediate" to gemifloxacin and had increased MICs to delafloxacin. Finally, one isolate contained no known quinolone resistance determinants but repeatedly tested as resistant to ciprofloxacin with increased MICs to gemifloxacin and delafloxacin.
throughout the United States (1). We also found a relatively low rate of ciprofloxacin resistance (6%) in the *N. gonorrhoeae* isolates in our study, which is in contrast to the rapid increase in resistance to quinolone antibiotics seen nationwide (approximately 20 to 40% of isolates in 2016, depending on location) (1, 20). However, several of the ciprofloxacin-resistant isolates we identified contained a combination of mutations in the *gyrA* and *parC* genes known to confer high-level resistance to ciprofloxacin (15, 21). These isolates also displayed increased MICs to quinolones that have been proposed as next-generation agents to combat multidrug-resistant *N. gonorrhoeae*—gemifloxacin and delafloxacin—suggesting that determinants of resistance to ciprofloxacin may be poised to rapidly acquire resistance and/or exhibit cross-resistance to other antibiotics in the quinolone class (17, 21). Furthermore, all isolates containing quinolone resistance determinants were also resistant to antibiotics from at least two other classes, suggesting that quinolone resistance may be a marker of multidrug resistance and that this association could be a useful tool as a surrogate for multidrug resistance to incorporate into future molecular diagnostic methods. One limitation of the antimicrobial resistance characterization in our study is that opportunistically available isolates from routine urine cultures could have biased our recovery of antibiotic-resistant organisms.

Given the AST profiles of the *N. gonorrhoeae* isolates in our study and considering that the majority of patients (57/64 [89%]) received single-dose azithromycin (1 g) plus ceftriaxone (250 mg) at the time of presentation (Table 1), we predict that the cure rate for gonorrhea using empirical treatment in our patient population is high.
FIG 5  Core genome phylogenetic analysis with geographic information and antibiotic resistance determinants. A maximum likelihood phylogenetic tree was constructed based upon the 1,563 core gene alignment created in PRANK. Bootstrap support values below 90% are shown (Continued on next page)
percentage of *N. gonorrhoeae*-positive cultures that were concurrently diagnosed by NAAT (Table 1) also implies that few additional cases are diagnosed by urine culture. Taken together, these findings suggest that the current standard of care for *N. gonorrhoeae* diagnosis, NAAT, continues to be appropriate in our patient population, except in cases of suspected treatment failure when cultured isolates are necessary for AST.

Nevertheless, global trends in *N. gonorrhoeae* antimicrobial resistance portend the need for AST methods that are more accessible than the agar dilution method, which is labor-intensive and performed only in reference laboratories. Studies comparing agar dilution to gradient diffusion devices for several antibiotics have demonstrated acceptable performance characteristics of gradient diffusion devices compared to agar dilution for several antibiotics (22–24). We found a high degree of correlation between gradient diffusion and disk diffusion AST results for tetracycline and ciprofloxacin, as reflected by the observed $R^2$ values, suggesting that disk diffusion may also be an acceptable and accessible method of tetracycline and ciprofloxacin resistance testing in *N. gonorrhoeae*; however, this may require future reassessment of the tetracycline breakpoints. Although no major or very major errors occurred, 24/64 isolates (37.5%) tested susceptible by disk diffusion but intermediate by gradient diffusion.

Given the ongoing challenges of performing AST on *N. gonorrhoeae* isolates in the clinical setting, direct-from-specimen genotyping, especially when multiplexed with NAAT, could have a major impact on guiding the choice of antimicrobial therapy (25). Indeed, a recent proof-of-concept study showed that ciprofloxacin therapy was 100% effective in curing gonorrhea caused by strains lacking resistance-conferring *gyrA* mutations (26). Our study adds to a growing body of literature demonstrating strong associations between specific genotypic markers and resistance to several antibiotics [i.e., *bla*TEM-1b for penicillin, *tet(M)* for tetracyclines, and *gyrA* with or without *parC* for quinolones], suggesting that additional genetic targets—via a targeted or whole-genome sequencing approach—may enable the effective use of antimicrobials that are no longer appropriate for use as empirical therapy (27–29). However, our identification of one ciprofloxacin-resistant isolate lacking any known quinolone resistance determinants highlights the current limitations of this approach and the need for ongoing efforts to interrogate the resistome of *N. gonorrhoeae* strains and expand the repository of genetic determinants of phenotypic resistance.

**MATERIALS AND METHODS**

**Ethics statement.** Prior to study initiation, this investigation was reviewed by the institutional review board at the Washington University School of Medicine and approved with a waiver of consent.

**Clinical isolates.** Growth of bacterial isolates from clinical specimens identified as *N. gonorrhoeae* by matrix-assisted laser desorption ionization—time of flight mass spectrometry (MALDI-TOF MS) using the BioTyper (Bruker Corporation) were frozen at −80°C in reconstituted powdered milk. Prior to antimicrobial susceptibility testing (AST) and sequencing, isolates were thawed and subcultured twice on chocolate agar (Hardy Diagnostics) incubated at 35°C ± 2°C with 5% CO₂, and the organism identification was confirmed with Vitek MS MALDI-TOF MS (bioMérieux). Patient characteristics such as age, sex, coinfections, and zip code of residence, were obtained from retrospective review of the electronic medical record.

**Antimicrobial susceptibility testing.** After two subcultures, plates containing GC II agar with IsoViteX enrichment (Becton-Dickinson) were inoculated with a 0.5 McFarland direct-colony suspension of each isolate. The plates were incubated at 35°C ± 2°C with 5% CO₂ and read by two independent readers after 24 h of incubation. Quality control was performed each day of testing (*N. gonorrhoeae* ATCC 49226 and *Escherichia coli* ATCC 25922).

For AST results, results from two independent readers were averaged and then rounded up to the nearest doubling dilution (gradient diffusion method only). For isolates with AST values that were off the measurable scale, a value of 65 mm (disk) or the lowest value on the gradient strip was assigned as the value for the purpose of analysis. Categorical results were determined by comparing these results to established clinical breakpoint criteria from the Clinical and Laboratory Standards Institute (CLSI) M100-S29 document (30) with the following exceptions. Doxycycline breakpoints were inferred from

**FIG 5 Legend (Continued)**

at the corresponding branch-points. The geographic origin of each isolate (zip code) is denoted by color adjacent to the isolate name. The presence or absence of an antibiotic resistance determinant in a particular isolate is denoted by a colored or blank square, respectively. If resistance determinants in the ResFinder or PointFinder databases were not found in any of the isolates, they are not shown in the figure.
tetracycline (30). Gemifloxacin and delafloxacin breakpoints were inferred from ciprofloxacin (30). The recently published CLSI M100-S29 breakpoint was used for azithromycin (30). The European Union Committee on Antimicrobial Susceptibility Testing (EUCAST) breakpoints (31) were used for ceftriaxone. Additional information on AST devices, including manufacturer and breakpoint criteria, can be found in Table S1 in the supplemental material.

**Whole-genome sequencing and analysis.** Sample preparation, sequencing, and sequence analysis were performed as described previously (32) using 12 World Health Organization (WHO) reference genomes (18) (Table S2). Genomic DNA was obtained from pure *N. gonorrhoeae* cultures using the QiAamp BIOTEST Bacteremia DNA kit (Qiagen, Germantown, MD, USA), and 0.5 ng of genomic DNA was used to create sequencing libraries with the Nextera kit (Illumina, San Diego, CA, USA) (33). Samples were pooled and sequenced on an Illumina NextSeq to obtain 2 × 150 bp reads. The raw reads and Illumina adapters were removed with trimmomatic v.38, and contaminating DNA was removed with DeconSeq v.4.3 (34, 35). Processed reads were used to de novo assemble scaffolds with SPAdes v3.13.0 (36). Quality statistics were obtained with QUAST v.4.5, and protein-coding sequences were identified with prokka v1.12 (Table S3) (37, 38). Of 64 genomes, 61 passed in silico quality assessment and were used in genomic analysis. We additionally gathered World Health Organization (WHO) reference *N. gonorrhoeae* genomes and identified all protein-coding sequences with prokka (18). Coding sequences were clustered at 95% identity with Roary v.3.12.0, and the 1,563 core genes were used to create a core genome alignment with PRANK v1.0 (19, 39). The core genome alignment was converted to a maximum likelihood phylogenetic tree with RAxML and viewed in iTol with an overlay showing zip code and identified resistance determinant (40, 41). Vertically transmitted mutations associated with antimicrobial resistance were identified with PointFinder v4.0 on the scaffolds.fasta file from SPAdes. Acquired antibiotic resistance genes were identified with ResFinder v4.0 (42).

**Data analysis.** Data were tabulated in Excel (Microsoft), where demographic and summary statistics were calculated. Data were then imported into Prism version 7 (GraphPad) for graphing and group comparisons. All statistical methods are described in the corresponding figure legends. Color overlays and final figures were prepared in Illustrator CC 2017 (Adobe).

**Data availability.** All genomes used in this study have been deposited into the NCBI Whole Genome Shotgun Database associated with BioProject accession no. PRJNA504667.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00373-19.

**TABLE S1**, XLSX file, 0.01 MB.

**TABLE S2**, XLSX file, 0.1 MB.

**TABLE S3**, XLSX file, 0.03 MB.

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We declare that we have no conflicts of interest.

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