Gram-positive uropathogens: Empirical treatment and emerging antimicrobial resistance

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Abstract

Urinary tract infection (UTI) is a global problem. Most UTI research focuses on gram-negative etiology. Enterobacteriaceae was found to be the most prevalent UTI infection constituting more than 80% of all the reported cases. The major gram-positive bacteria in UTI cases are Staphylococcus saprophyticus, Enterococcus faecalis, Streptococcus agalactiae. Gram-positive pathogens were reported in multiple countries in both uncomplicated and complicated UTI. Antibiotic therapy of gram-positive bacteria is completely different than that of gram-negative UTI pathogens. However, symptoms associated with UTI caused by gram-positive and gram-negative are very similar. Without proper diagnosis, there is a high possibility of getting the wrong diagnosis and subsequent antibiotic therapy. Very limited studies are available focusing etiology and their antibiotic susceptibilities in Bangladesh perspective. We aimed to seek in this gap-filling research area.

Methods and materials

A cross-sectional study was conducted between July 2018 and December 2018 to seek gram-positive bacterial isolate from patients UTI infection constituting more than 80% of all the reported cases. The major gram-positive bacteria in UTI cases are Staphylococcus saprophyticus, Enterococcus faecalis, Streptococcus faecalis [4-6]. Gram-positive uropathogens were reported in multiple countries in both uncomplicated and complicated UTI [7-9]. There are no such exclusive studies have been done in Bangladeshi population to seek gram-positive UTI etiology. Without proper diagnosis, there is a high possibility of getting wrong and inappropriate medications and subsequent adverse side-effects of the associated treatment. Gram-positive bacteria are found more often as etiologic agents of UTI among pregnant women and the elderly [10]. Further, complicated UTIs often occur in nosocomial and/or institutional settings and these individuals are at greater risk of gram-positive and polymicrobial UTI [4,5]. It accounts for a large proportion of antibiotic consumption and has a large socioeconomic impact and may contribute to bacterial resistance [11,12]. Clinicians often face problems in choosing appropriate antibiotic for empirical therapy for treating UTIs [13]. In Bangladesh, indiscriminate use of antibiotic use brings many adverse outcomes to UTI patients clinically with an extended economic burden. This study aimed to assess the susceptibility profile of gram-positive uropathogens against different prescribe antibiotics. Our study had rationalized the necessity of continued monitoring of urinary tract infected patients for a better understanding of their current epidemiology by gram-positive bacterial strains in the Bangladeshi population.

Methods and materials

Data collection

Patient's socio-demographic information, treatment history (whether the patients were treated with an antimicrobial therapy), and any relevant clinical information (kidney stones, pregnancy, recent catheter usage) were recorded by a structured questionnaire.

Study design and specimen collection

A cross-sectional study was conducted between July 2018 and December 2018 to seek gram-positive bacterial isolate from patients

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having urinary tract infection. This study was designed based on laboratory investigations of urine samples collected from UTI patients attending in different community hospitals in the Savar area. To actuate the planning, probable UTI patients who sought medical advice at Gonosastha Hospital and Enam Medical College in Savar, Dhaka, were requested to participate in this study. The study includes consented patients having clinical evidence of UTI with symptoms less than seven days presenting at the out-patient department of hospitals and those admitted for less than 48 h.

**Bacterial isolation and identification**

To isolate UTI pathogen, clean-catch midstream urine samples were collected in sterile glass tubes and inoculated on a differential culture medium, cysteine-, lactose-, and electrolyte-deficient (CLED) agar (Lypophilchem, Italy). Patients were explained about the standard clean-catch midstream sampling procedure beforehand to avoid contamination with commensal bacteria. To avoid false-positive results, urine specimens were plated within 2 h after collection. Using CLED agar provides advantages of growth-supports of most potential uropathogens including gram-positive cocci such as *Staphylococcus aureus*, *Staphylococcus saprophyticus*, and *Enterococcus spp*. In parallel, Mannitol Salt Agar (MSA) was inoculated for targeting gram-positive UTI growth. Urine cultures were incubated overnight at 35°C–37°C in ambient air. Quantitative urine cultures were done, and colony counts 10^2 or 10^3 CFU/mL were considered to define probable UTI in ambient air. Colony counts of less than 10^6 CFU/mL were assumed as potential contamination. The purified bacterial colonies were identified by gram staining, conventional biochemical procedures followed by a rapid biochemical-test kit (API 20E, BioMe’rieux, Durham, NC) consisting of a set of chromogenic panels, carbohydrate batteries, and enzymatic substrates. Bacterial identification was validated further by genotyping using 16S rRNA of the isolates in part.

**Antimicrobial sensitivity assessment**

Isolates were subjected to in vitro antibiotic sensitivity test by standard Kirby-Bauer disc diffusion method and zone of inhibition were interpreted following CLSI (Clinical amp; Laboratory Standards Institute-2010) recommendations. Six antibiotic discs from beta-lactam, nitrofurantoin, and fluoroquinolone were placed on the agar (Lyophilchem, Italy). Patients were explained about the standard culture medium, cysteine-, lactose-, and electrolyte-deficient (CLED) agar (Lyophilchem, Italy). Patients were explained about the standard clean-catch midstream sampling procedure beforehand to avoid contamination with commensal bacteria. To avoid false-positive results, urine specimens were plated within 2 h after collection. Using CLED agar provides advantages of growth-supports of most potential uropathogens including gram-positive cocci such as *Staphylococcus aureus*, *Staphylococcus saprophyticus*, and *Enterococcus spp*. In parallel, Mannitol Salt Agar (MSA) was inoculated for targeting gram-positive UTI growth. Urine cultures were incubated overnight at 35°C–37°C in ambient air. Quantitative urine cultures were done, and colony counts 10^2 or 10^3 CFU/mL were considered to define probable UTI in ambient air. Colony counts of less than 10^6 CFU/mL were assumed as potential contamination. The purified bacterial colonies were identified by gram staining, conventional biochemical procedures followed by a rapid biochemical-test kit (API 20E, BioMe’rieux, Durham, NC) consisting of a set of chromogenic panels, carbohydrate batteries, and enzymatic substrates. Bacterial identification was validated further by genotyping using 16S rRNA of the isolates in part.

**Detection of β-lactamase and quinolone-resistant genes**

Isolates that were showing resistant phenotypically were subjected to PCR for the detection of *bla*TEM and *qnrS* gene. A total of six bacterial lysates were prepared. PCR was carried out in PCR tube containing bacterial lysate as DNA template (1 µl) with a final volume 12 µl containing DNA 2x premix (6 µl), 5 pmol of each primer (1 µl) and deionized water (4 µl). PCR was carried out in a thermal cycler and the cycling condition for *bla*TEM and *qnrS* were: initial denaturation at 94°C for 5 min followed by 32 cycles of amplification with denaturation at 94°C for 30 s, annealing at 50°C for 30 s (*bla*TEM) and 55°C for 30 s (*qnrS*) and extension at 72°C for 1 min, ending with a final extension at 72°C for 7 min. PCR products of the expected size were purified and sequenced using the same primers used to generate the PCR products.

**Statistical analyses**

As described, the validated Bangla of the questionnaire was used for data collection. Completed data collection forms were rechecked by data collection supervisors. Randomly selected 10% laboratory analyses were done twice by different research person to ensure the authenticity of the laboratory data and to assess inter-researcher biases. Verified data were entered and subsequently analyzed using IBM SPSS statistics data editor (version 21). Missing data were excluded from the bivariate analysis. Descriptive statistics were reported as frequency, percentage, and means when applicable along with standard deviation. The antimicrobial susceptibility of test bacteria was evaluated by a 3-type categorical rating scale as susceptible (antibiotic indicated as fully effective), intermediate (indicated as partial effective), and resistant (indicated as non-effective). For the testing association between categorical data, Pearson’s chi-square test was used, and Yate’s correction for continuity was applied where appropriate. As such the exposure of antimicrobials to the acquisition of antibiotic-resistant gene was calculated. The correlation coefficient was calculated to check the strength of the association. Independent sample t-tests were performed when comparing the means of continuous variables. A two-tailed p-value smaller than 0.05, was considered statistically significant.

**Results**

**Demographics of the study population**

We approached patients who attended medical doctors of Gonosastha Medical College with their UTI associated complications. We included 147 patients in this study who had severe clinical symptoms. Of them 36 (24.5%) were male and the remaining 111 (75.5%) were female. Therefore, almost three-fold UTI patients were found female in comparison to their opposite gender (Figure 1). Chi-square test showed female gender associated with a significantly higher number of UTI events (*p* = 0.04). Age distribution of our study cases lies from below one year to above 70 years. However, most frequent UTI infection was in the age group between 21-30 years. Overall age-wise case distribution was described in table 1. A total of five out of 12 *Staphylococcus spp* (42%) was identified in the age group 21-30 years. No other trend was observed in age-distribution infection by gram-positive UTI infection.

**Isolation and identification of gram-positive UTI pathogens**

We examined 147 urine samples from clinically symptomatic patients. CLED culture medium produced different color colonies for different UTI pathogens. Gram-positive *Staphylococcus aureus* produce yellow colonies and *Staphylococcus saprophyticus* generates white to yellowish colonies, and *Enterococcus spp* produces colorless to yellow colonies there. Gram-negative bacteria produce different other colors. Therefore, identifying UTI bacteria by observing colony colors on CLED may be presumptive, but difficult to conclude. We sub-cultured those probable Gram-positive colonies from CLED culture medium on to Mannitol Salt Agar (MSA) and incubated overnight at 37°C. Growth patterns of different UTI bacteria were: initial denaturation at 94°C for 5 min followed by 32 cycles of amplification with denaturation at 94°C for 30 s, annealing at 50°C for 30 s (*bla*TEM) and 55°C for 30 s (*qnrS*) and extension at 72°C for 1 min, ending with a final extension at 72°C for 7 min. PCR products of the expected size were purified and sequenced using the same primers used to generate the PCR products.

**Table 1. Age-distribution of UTI study patients**

<table>
<thead>
<tr>
<th>Age in years</th>
<th>Number of UTI patients</th>
<th>Frequency %</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-0.5</td>
<td>1</td>
<td>0.7</td>
</tr>
<tr>
<td>1-10</td>
<td>5</td>
<td>3.4</td>
</tr>
<tr>
<td>11-20</td>
<td>23</td>
<td>15.6</td>
</tr>
<tr>
<td>21-30</td>
<td>49</td>
<td>33.3</td>
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<tr>
<td>31-40</td>
<td>21</td>
<td>14.3</td>
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<td>41-50</td>
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<td>18.4</td>
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<tr>
<td>51-60</td>
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<td>6.8</td>
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<tr>
<td>61-70</td>
<td>9</td>
<td>6.1</td>
</tr>
<tr>
<td>71-80</td>
<td>2</td>
<td>1.4</td>
</tr>
<tr>
<td>Total</td>
<td>147</td>
<td>100</td>
</tr>
</tbody>
</table>
colonies on MSA showed white-to-yellowish and pink. Bacteria from MSA were subjected to Gram’s staining and found Gram-positive nature. Preliminary gram-positive UTI bacteria have been identified by this study are Staphylococcus species. Analytical profiling index (API) analysis and/or 16sRNA DNA sequencing can be done to identify the etiology into species level. We have detected 12 gram-positive UTI pathogens, which was 8.2% of the total pathogen population.

**Antimicrobial resistance pattern of the UTI gram-positive isolates**

Disc diffusion experiment produced clear zones of bacterial lawn on Muller-Hinton (MH) agar, indicating susceptibility of the respective antibiotics impregnated there. In contrast, resistant isolates did not allow clear zone even in the presence of the antibiotic disc. A mixed trend of the susceptibility against the tested antibiotics was observed in our assessment. The highest resistance was found against nitrofurantoin (91.7%), followed by amoxicillin (83%), ofloxacillin (50%). Both gentamicin and amikacin were found very effective drugs, where only 8.3% was found resistant (Figure 2).

**Antibiotic resistant genes in gram-positive uropathogens**

We prepared DNA samples from all 12 gram-positive isolates by liberating DNA in heat-boiling methods. In the current analysis, we examined beta-lactam ring containing antibiotic-resistant gene bla-TEM and found two isolates (2/6, 33%) isolates positive. Similarly, we checked for fluoroquinolone group antibiotic-resistant gene qnrS and found three isolates positive (3/6, 50%).

**Discussion**

Urinary tract infections represent one of the most common diseases encountered in medical practice, causing significant associated morbidity and occurring from neonates to the elderly [5,15,16]. Prevalence of infections may differ with age, sex, and certain predisposing factors. Most of the reported UTI infection was caused by gram-negative bacteria mainly Enterobacteriaceae. There were also strong evidences against gram-positive UTI infection [8,17]. We sought to examine the prevalence of gram-positive UTI pathogens in the Bangladeshi population. Empirical treatments for urinary infection-based severity criteria and clinical symptoms carry great challenge [18]. As antibiotic therapy of gram-positive bacteria is completely different than that of gram-negative UTI pathogens, there is a high possibility of getting the wrong treatment by an empirical approach. Therefore, correct diagnosis with specific antibiotic susceptibility testing is imperative for setting any treatment stewardship. As UTI is a common community-acquired disease in Bangladesh, therefore, a significant percentage of the population are getting infected with this disease each year. The study was designed to check antimicrobial susceptibilities of gram-positive UTI pathogens as well. We detected about 8.2% gram-positive UTI etiologies, notably, Staphylococcus aureus, Staphylococcus saprophyticus, and Enterococcus spp. which attached earlier published reports [19].

Commonly, UTI infection was recorded much higher in females than in males [20]. Unlike, we found half of the gram-positive infection was reported in males. The preliminary results showed males are more vulnerable by gram-positive in comparison to the overall situation. Age distribution of our study cases lies from below one year to above 70 years. However, most frequent gram-positive UTI infection was in the age group between 21-30 years as supported elsewhere [21].

Antibiotic resistance among uropathogens has become a public health concern in Bangladesh. In this study, antibiotic susceptibility profiles of gram-positive UTI pathogens were determined with six different antibiotic discs a by disc diffusion test. The highest percentage of resistance was observed against nitrofurantoin, whereas the lowest percentage of resistance was found against amikacin and gentamicin. Less than 10% of the gram-positive uropathogens appeared resistant against those two aminoglycoside group antibiotics. The results support aminoglycoside as a drug of choice for gram-positive UTI pathogens. Among the quinolone group of antibiotics, the highest percentage of resistance had been found by 2nd generations lomefloxacin (75%). Our antibiogram findings showed nitrofurantoin was not a very effective drug for gram-positive UTI bacteria. The old beta-lactam antibiotic, amoxicillin was weakly functional against gram-positive UTI pathogens. Fluoroquinolone antibiotics, ofloxacin and lomefloxacin were found moderate level effective.

The isolates were tested for selected AMR genes by polymerase chain reaction (PCR), using primers specific for the detection of bla-TEM and qnrS. About 33% isolates were positive for bla-TEM and 50% was positive for qnrS. Antibiotic-resistant genes were detected and showed harmony to the phenotypic susceptibility pattern. Although utmost sincerity and dedication were investigated to carry out the study it could not go beyond limitations as the sample size was not

**Figure 1. Distribution of gram-positive and gram-negative bacteria among male and female. Female had a significant higher number of UTI than males which was caused by gram-negative bacteria. In the case of gram-positive urinary tract infection, male and female showed equal vulnerability**

**Figure 2. Prevalence of antibiotic resistance among gram-positive UTI bacteria.** AMC: Amoxicillin; LOM: Lomefloxacin; OFX: Ofloxacin; CN: Gentamicin; AK: Amikacin; F: Nitrofurantoin. Amikacin and Gentamicin appeared most effective antibiotics against Gram-positive UTI bacteria. Nitrofurantoin was found the least effective among the tested antibiotics.
large enough. Despite this limitation hopefully, this finding of the study will help clinicians deciding to prescribe antibiotics from the evidence-based study. The clinician should take care of the specific diagnosis before prescribing empirical antibiotic therapy, particularly for male UTI.

Ethics statement

This study was approved by the Ethics and Research Review Committee of the Jahangirnagar University Faculty of Biological Sciences and verbal consent was obtained from UTI patients and doctors.

Acknowledgments

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Disclosure

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