Tigecycline in-vitro susceptibility and antibiotics’ fitness for gram-negative pathogens

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ABSTRACT

Objectives: To determine the tigecycline in-vitro susceptibility of naïve gram-negative pathogens from serious infections in Delhi, India.

Methods: During July to October 2007 investigations were carried out to determine the tigecycline in vitro susceptibility of 50 consecutive gram-negative pathogens from serious infections at the Sant Parmanand Hospital, Delhi, India. Minimum tigecycline inhibitory concentrations were determined employing the E test method (AB Biodisk).

Results: Twenty-four percent of isolates were found to be tigecycline resistant or partly susceptible. Susceptibility of the isolates was lower than meropenem but similar to piperacillin-tazobactam, amoxicillin-clavulanic acid, and amikacin.

Conclusion: Tigecycline resistance was prevalent in the gram-negative isolates from serious infections prior to its marketing in India. The choice of any recently marketed antibiotic for a pilot treatment against serious gram-negative infections should not be automatic. In the initial phase of its marketing, it should be evaluated in parallel with the antibiotics with excellent local susceptibility profiles.

Tigecycline, a glycyclcline is a structural analogue of minocycline designed to avoid tetracycline resistance mediated by ribosomal protection and drug efflux. It is indicated for the treatment of complicated skin and skin-structure infections and complicated intra-abdominal infections and is available for intravenous administration only. Tigecycline, with a broad spectrum of activity against multidrug-resistant gram-positive and gram-negative pathogens is expected to be useful in the treatment of conditions caused by these pathogens. The intravenous formulation exhibits linear pharmacokinetics with a rapid distribution along with a large volume of distribution and extensive tissue penetration. The long terminal elimination half-life of around 40 hours allows a twice-daily administration. Tigecycline undergoes very limited metabolism and the unchanged drug is eliminated through the feces, with glucuronidation and renal routes. Recently, tigecycline was reported as a promising recipe for gram-negative infections. In Philadelphia, the clinical, and microbiological response to tigecycline therapy in 18 patients with serious multidrug-resistant gram-negative
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organisms was far from ideal. During 2007, tigecycline marketing was initiated in India. We investigated the susceptibility or otherwise of local gram-negative isolates during the pre-introduction phase. In-vitro susceptibility assays were carried out at the Sant Parmanand Hospital, Delhi, India, a 140-bedded, multi-speciality, tertiary care, hospital in Delhi. Susceptibility profiles to gram negative pathogens associated with serious infections were not that optimistic.

Methods. During July and October 2007, 50 successive gram-negative isolates from patients with intense infection, including those in the intensive care units, medical, surgical, neonatal, were evaluated. Prior approval from the Ethics Committee was not required. Isolates were drawn from urine (25 isolates), blood and purulent material (10 isolates each), and pulmonary tissues (5 isolates). Tigecycline susceptibility was measured as the isolates' minimum inhibitory concentration (MIC) values. Isolate susceptibility was simultaneously evaluated against the locally favored antibiotics in such cases. Identification was based on gram staining, biochemical reactions with sugars, including indole, urea, citrate, lysine iron agar, triple sugar iron, and phytyl pyruvic acid reactivity. Isolates included Escherichia coli (E. coli) (17 isolates), Klebsiella species (30 isolates), Proteus species (1 isolate), and Pseudomonas species (2 isolates). They were drawn from urine (28 isolates) purulent material (12 isolates), blood (4 isolates) and pulmonary tissues (6 isolates). Tigecycline susceptibility was determined using the E test method (AB Biodisk, Solna, Sweden). For other antibiotics, disk diffusion method, employing disks (Difco or Oxoid), was employed. The interpretations, depending on diameter of the zone of inhibition, were according to the clinical Laboratory Standards Institute (CLSI) criteria. Tigecycline isolates with MIC values of ≤2 μg/ml were regarded as susceptible, 2-≤8 μg/ml partially resistant, and ≥8 μg/ml resistant. Isolates were labeled to be susceptible or resistant to others, in accordance with the guidelines of the CLSI.

Eight isolates including E. coli (3 isolates) and Klebsiella (5 isolates) were tested for extended-spectrum beta-lactamase (ESBL) production. Extended-spectrum beta-lactamase production was determined according to CLSI guidelines employing a disk combination of ceftazidine, ceftazidine-clavulanic acid, cefotaxime, and cefotaxime-clavulanic acid (Becton Dickinson, Sparks, Maryland).

Results. Tigecycline MIC was ≤2μg/ml in E. coli (11 isolates), Klebsiella (20 isolates) and there was only one Proteus (isolate). An MIC of 2- ≤8 μg/ml was observed in E. coli, 2, Klebsiella, 4 and Pseudomonas, 1. An MIC ≥ 8 μg/ml was recorded in E. coli (2 isolates), Klebsiella (4 isolates), and Pseudomonas (1 isolate), shown in Table 1. Of the 4 tigecycline resistant/partly susceptible E. coli, all were susceptible to meropenem, 3 to piperacillin-tazobactam, 2 to amoxicillin-clavulanic acid and amikacin. Of the 6 tigecycline resistant/partly susceptible Klebsiella, 5 were susceptible to meropenem, and piperacillin-tazobactam each, and 3 to amoxicillin-clavulanic acid and amikacin each. Concurrently, in the 2 Klebsiella or E. coli resistant either to meropenem, piperacillin-tazobactam, and amoxicillin-clavulanic acid or amikacin, only one was tigecycline susceptible. The 5 ESBL-producer E. coli and Klebsiella were susceptible to tigecycline. Extended-Spectrum beta-Lactamase negatives included one tigecycline susceptible E. coli and Klebsiella, while one Klebsiella was partially susceptible/partially resistant. All the Klebsiella isolates, which were resistant to piperacillin-tazobactam, amoxicillin-clavulanic acid, and amikacin, shown in Table 1 were resistant to tigecycline as well.

Discussion. Marketing of new antibiotics in developing countries is accompanied by extensive dissemination of information regarding its superiority over the antibiotics in common usage locally. Investigations on tigecycline naïve pathogens in the

Table 1 - In-vitro susceptibility to tigecycline and other popular antibiotics at Sant Parmanand Hospital, Delhi.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Tigecycline MIC</th>
<th>Meropenem</th>
<th>Piperacillin-tazobactam</th>
<th>Amoxicillin-clavulanic acid</th>
<th>Amikacin</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt;2μg/ml</td>
<td>2-≤8μg/ml</td>
<td>≥8μg/ml</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>11</td>
<td>2</td>
<td>2</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>Klebsiella</td>
<td>20</td>
<td>4</td>
<td>2</td>
<td>24</td>
<td>2</td>
</tr>
<tr>
<td>Proteus</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Pseudomonas</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>32</td>
<td>7</td>
<td>5</td>
<td>40</td>
<td>2</td>
</tr>
</tbody>
</table>

S = number susceptible, R = number resistant.
Indian capital metropolis of New Delhi were carried out before its local marketing. A comparison was aimed with the therapeutic options in serious infections by gram negatives, including ESBL-producers, in the hospital. The current choice has been of meropenem, piperacillin-tazobactam, amoxicillin-clavulanic acid, and amikacin. Tigecycline resistant or partially susceptible isolates among the above 50 consecutive gram-negatives were 24%: 95% confidence interval 10.6-42.6%. The total isolates susceptibility to tigecycline was lower than meropenem \( (p=0.0079) \), however, resembled piperacillin-tazobactam, amoxicillin-clavulanic acid, and amikacin. Furthermore, there was not a single isolate that was exclusively susceptible to tigecycline.

The above investigations have been confined to a solitary hospital but the isolates had been tigecycline naïve. One would have to be very guarded to attempt any extrapolations. Nevertheless, even with not very many isolates, one could infer that tigecycline would be very unlikely to be the exclusive recipe in the future. That would be important for any empirical therapy before report on in-vitro antibiotic susceptibility was known. The above in-vitro data on tigecycline efficacy supports the University of Philadelphia Hospital clinical and microbiologic data. In all probability, chances of tigecycline emerging as the first favorite for clinicians’ in severe gram negative infections would be remote.

To conclude, those responsible for treatment of serious infections would be obliged to offer the most efficient recipe. They should also bear in mind risks for emergence of antimicrobial resistance during clinical use of antibiotics. Their hard work to manage patients with use of the most potent agents around could affect the prevalence of antimicrobial resistance. The antibiotic prescription practices would vary dramatically in different countries. In several countries, a prior in-vitro susceptibility result on patient's isolates would be necessary before prescribing any newer antimicrobial. On the contrary, in countries with over-the-counter availability of antimicrobials and no antibiotics policy around, clinicians would be tempted to prescribe recently marketed antibiotics. To prescribe newer antibiotics expecting a swift and optimistic response might not be rational all the times. A better strategy towards patient cure would be the retrospective antibiotic susceptibility profiles, shown in Table 1 in the above study, to select the ideal empirical therapy in serious gram negative infections.

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**References**