Detection of Amp-C Beta Lactamase Enzyme Production among Enterobacteriaceae and Comparison of Different Inducer Substrate Combinations for Detection of Inducible Amp-C

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A B S T R A C T

AmpC beta-lactamases hydrolyse penicillins, monobactams, cephalosporins and cephamycins. AmpC producers are resistant to beta-lactam/beta-lactamase inhibitor combinations therapeutically. AmpC is generally underreported which leads to therapeutic failures and uncontrolled spread of these resistant strains. Hence, there is an increased need to detect AmpC routinely in the laboratory. To detect AmpC beta-lactamase production among Enterobacteriaceae isolated from clinical samples and to compare different inducer substrate combinations for the detection of inducible Amp-C (iAmpC). 100 clinical isolates of Enterobacteriaceae were tested. Constitutive AmpC (cAmpC) detected using inhibitor based method using Cefoxitin (CN) and CN with Phenylboronic acid (PBA). Inducible AmpC detected using disk approximation test using inducers Imipenen (I) and Cefoxitin (CN), and substrates Cefotaxime (CTX), Ceftazidime (CAZ) and Piperacillin-Tazobactum (PT). Various combinations tested were I/PT, I/CTX, I/CAZ, CN/CTX, CN/CAZ. AmpC production was detected in 30% of isolates, 23% were constitutive and 7% were inducible. Commonest AmpC producer was Enterobacter sp with 7(36.84%) and 4(21.05%), followed by E. coli 14(26.92%) and 3(5.76%) constitutive and iAmpC respectively. 2(8.69%) Klebsiella demonstrated only cAmpC. I/PT combination detected all the 7 iAmpC, others I/CTX and I/CAZ detected only 3isolates. Simple disk method of cefoxitin with boronic acid and I/PT combination can be used to detect constitutive and inducible AmpC respectively.

Keywords: beta-lactamases, AmpC, Cephalosporinases, Boronic acid, Inducers

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Introduction

Beta-lactamases are clinically important cephalosporinases encoded on the chromosomes of many Enterobacteriaceae and a few other organisms, where they mediate resistance to cephalothin, cefazolin, cefoxitin, most penicillins, and beta-lactamase inhibitor/beta-lactam combinations (Gunjan Gupta et al., 2014). Isolates that coproduce both an Extended spectrum beta-lactamase (ESBL) and a high level of AmpC are becoming more common (Naveen Grover., 2013). Mechanism of drug resistance in
AmpC β lactamase can be chromosomal or plasmid mediated. Chromosomal mediated resistance is due to mutation in the nucleotide sequence at some point of the DNA of the bacteria and such genes are not easily transferable to other bacterial species. Plasmid mediated AmpC β lactamases have arisen by the transfer of chromosomal genes for AmpC β-lactamase onto plasmids. These genetic determinants can spread laterally and to other bacteria through lateral transfer of plasmids (Ascelijn Reuland, 2015).

Majority of AmpC β lactamases are chromosomally mediated (Unlike ESBLs which are plasmid mediated) and are found in SPACE bugs (Serratia, Pseudomonas, Acinetobacter, Citrobacter and Enterobacter spp.). Plasmid mediated AmpC β lactamases are seen in isolates of E.coli, K. pneumoniae, Salmonella spp, Citrobacter freundii, Enterobacter aerogenes, and Proteus mirabilis (George, 2009). Failure of empirical therapy is a frequent and common problem in infections caused by Amp-C producing isolates leading to significant morbidity and mortality (Harris PN., 2012, Conan MacDougall., 2011). There are no standard Clinical Laboratory Standards Institute (CLSI) guidelines to detect the presence of Amp-Cβ-lactamase. Amp-C is generally underreported which leads to therapeutic failures and spread of these resistant strains (George A 2009). Hence, there is an increased need to detect AmpC routinely in the laboratory.

Objectives

1) To detect AmpCβ-lactamase enzyme production among Enterobacteriaceae isolated from various clinical samples

2) To compare different inducer substrate combinations for the detection of inducible Amp-C(iAmpC)

Materials and Methods

A total of 100 clinical isolates of Enterobacteriaceae from various samples like pus, urine, blood and sputum were included in the study. The isolates were identified by standard biochemical techniques.

Detection of AmpC

Constitutive AmpC (cAmpC) detected by inhibitor based method using Cefoxitin (CN) (30μg) and CN with Phenylboronic acid (PBA) (30/400 μg) disks (Philip E. Coudron, 2005; Shoorashetty, 2011)

120 mg of phenylboronic acid was dissolved in 3 ml of dimethyl sulfoxide. Three milliliters of sterile distilled water was added to this solution. Twenty microliters of the stock solution was dispensed onto disks containing 30 μg of Cefoxitin. The boronic acid disk test was performed by inoculating Mueller-Hinton agar by the standard disk diffusion method and placing a disk containing 30 μg of cefoxitin and a disk containing 30 μg of Cefoxitin and 400 μg of boronic acid onto the agar. Inoculated plates were incubated overnight at 35°C. Bacteria that demonstrated a zone diameter around the disk containing cefoxitin and boronic acid that was 5 mm or greater than the zone diameter around the disk containing cefoxitin was considered an AmpC producer (Figure 1).

Double disk approximation method for the detection of iAmpC (Michael Dunne., 2005)

Disk diffusion susceptibility testing was performed as per standard guidelines. Inducers Imipenem (I) (10μg) and Cefoxitin (CN) (30μg), and substrates Cefotaxime (CTX) (30μg), Ceftazidine (CAZ) (30μg) and Piperacillin-Tazobactum (PT)(100/10 μg) were used. Various combinations tested were I/PT, I/CTX, I/CAZ, CN/CTX, CN/CAZ.
Inducer/substrate disks were placed on the surface of Mueller-Hinton agar plates at a distance of 25 mm apart. After incubation, zones of inhibition were measured on both the induced (adjacent to the inducer disk) and the uninduced side of the substrate disk from disk edge to zone edge. A test was considered positive if the zone of inhibition was reduced by ≥2 mm on the induced side of the substrate disk (Figure 2).

**Results and Discussion**

Out of 100 Enterobacteriaceae isolates, 52 were *E. coli*, 23 were *Klebsiella* spp, 19 were *Enterobacter* spp and 6 were *Citrobacter* spp. Amp C production was detected in 30% of isolates, 23% were constitutive and 7% were inducible (Figure 3). Commonest AmpC producer was *Enterobacter* sp with 7 (36.84%) and 4 (21.05%), followed by *E. coli* 14 (26.92%) and 3 (5.76%) constitutive and iAmpC respectively. 2 (8.69%) *Klebsiella* demonstrated only cAmpC (Figure 4). Sample wise distribution of cAmpC and iAmpC is depicted in Figure 5. I/PT combination detected all the 7 (100%) iAmpC, others I/CTX and I/CAZ detected only 3 (42.85%) isolates (Figure 6). Distribution of cAmpC and iAmpC among different strains in various clinical specimens is shown in Table 1.

Resistance to broad-spectrum beta lactams mediated by extended spectrum beta lactamases (ESBLs) and AmpC beta lactamases (AmpC βLs) enzymes is an increasing problem worldwide. In our study 23% of isolates were cAmpC and 7% of isolates were iAmpC. Shoorashetty and Nagarathnamma (2011) reported 33.5% cAmpC and 7% iAmpC. (Shoorashetty, 2011) Tan TY, Nq SY et al., (2009) reported 26% cAmpC (Thean Yen Tan, 2009)

In our study, I/PT combination detected 100% iAmpC. W. Michael Dunne, Jr. et al., in 2005, also found that I/PT combination provided the greatest sensitivity (97.1%) to detect iAmpC (W. Michael Dunne., 2005). Bacteria expressing AmpC β lactamases are of major clinical concern because these are usually resistant to all beta lactam antimicrobials, except for cefepime, cefpirome and carbapenems. The accurate detection of plasmid mediated AmpC is important to improve the clinical management of infection and to provide epidemiological data.

**Table 1 Distribution of cAmpC and iAmpC among different strains in various clinical specimens**

<table>
<thead>
<tr>
<th>Samples</th>
<th><em>E. coli</em>, n=52</th>
<th><em>Klebsiella</em> spp, n=23</th>
<th><em>Enterobacter</em> spp, n=19</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>cAmpC</td>
<td>iAmpC</td>
<td>cAmpC</td>
</tr>
<tr>
<td>Urine</td>
<td>9 (17.3%)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Exudates</td>
<td>4 (7.69%)</td>
<td>1 (1.92%)</td>
<td>-</td>
</tr>
<tr>
<td>Blood</td>
<td>-</td>
<td>2 (3.84%)</td>
<td>2 (8.69%)</td>
</tr>
<tr>
<td>Sputum</td>
<td>1 (1.92%)</td>
<td>-</td>
<td>1 (4.34%)</td>
</tr>
<tr>
<td>Stool</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
**Fig. 1** Constitutive AmpC producer

**Fig. 2** Inducible AmpC producer
**Fig. 3** Percentage of AmpC producers

**Fig. 4** Distribution of cAmpC and iAmpC among various isolates

**Fig. 5** Sample wise distribution of AmpC isolates
In conclusion, performing simple tests using boronic acid compounds as specific class C β-lactamase inhibitors for cAmpC and disk approximation test using I/PT combination for iAmpC enables clinical microbiology laboratories to report those strains producing class C β-lactamases thus helping physicians to select appropriate antimicrobial therapy. Determination of the prevalence of resistant strains is essential to formulate an effective antibiotic policy and hospital infection control measures.

References


RM Shoorashetty, T Nagarathnamma, J Prathibha. Comparison of the boronic
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