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Jornal Brasileiro de Patologia e Medicina Laboratorial, vol. 50, núm. 6, noviembre-diciembre, 2014, pp. 421-427

Sociedade Brasileira de Patologia Clínica/Medicina Laboratorial
Rio de Janeiro, Brasil

Available in: http://www.redalyc.org/articulo.oa?id=393541985006
Microbial resistance and frequency of extended-spectrum beta-lactamase (ESBL) in isolated from blood cultures

Resistência microbiana e frequência de betalactamase de espectro estendido (ESBL) em isolados de hemoculturas

Ruan Carlos Gomes da Silva¹; Amanda Cristina de Oliveira Silva¹; Sibele Ribeiro de Oliveira²

ABSTRACT

Introduction: The emergence and spread of isolated carriers of extended-spectrum beta-lactamase (ESBL) have complicated the treatment of nosocomial infections, since its production is not easily identified by the sensitivity tests, routinely performed in clinical laboratories, leading to difficulties in the hospital control of resistant microorganisms and antibiotics misuse. Objective: The objective of this study was to analyze the resistance profile and the frequency of ESBL in Gram-negative bacteria isolated from blood cultures. A hundred bacterial samples from blood cultures of adult patients were analyzed, which were phenotypically identified by biochemical tests of carbohydrates fermentation and submitted to determination of the resistance profile by disc diffusion test and ESBL screening by disc approximation and disc replacement methods. Results: Among the bacterial samples tested, 30 were identified as Gram-negative bacteria, predominantly by Proteus mirabilis, Pantoea agglomerans, and Escherichia coli. Of these, 73.33% were positive for the detection of ESBL by phenotypic tests, and was found mainly in Pantoea agglomerans, Proteus mirabilis, and Enterobacter cloacae. Conclusion: The increase in the occurrence of ESBL in different Enterobacteriaceae shows the importance of the amplification of detection in other species than Escherichia coli or Klebsiella sp., so that the assistance to the patient is not restrained, since these resistant bacteria cannot be detected by the laboratories. Considering the frequency of ESBL in this study, we highlight the importance of its detection, aiming to its contribution to the development of improvements in the health care policies of hospitals.

Key words: bacterial resistance; extended-spectrum beta-lactamase; ESBL; hospital infection.

INTRODUCTION

Nosocomial infections are a significant public health problem, because the effect of an unsuccessful treatment create situations that increase the time of hospitalization and the expenditure on medicines, generating recurrence, which are caused, among others, by the production of enzymes responsible for degrading the main antimicrobial used in clinical practice(10).

One of the main mechanisms of bacterial resistance is the production of extended-spectrum beta-lactamases (ESBL), enzyme mediated by non-inducible plasmid gene, capable of hydrolyzing the oxyimino-beta-lactams chain present in the chemical structure of beta-lactams, inactivating them and providing greater bacterial resistance, which extends its spectrum of activity to the broad-spectrum antimicrobials(11, 21).

ESBLs provide resistance to penicillins, all cephalosporins, and monobactams (including, aztreonam), but do not provide resistance to cephamycins (cefoxitin and cefotetan) and carbapenems (imipenem, meropenem, and ertapenem)(18), however, they are inhibited by beta-lactamases inhibitors, such as clavulanic acid, sulbactam and tazobactam(18).

ESBLs are derived from genetic mutations (temoneira [TEM] and sulfidril variable [SHV]), and are found mainly in Escherichia coli and Klebsiella sp., but can also be detected in Proteus mirabilis, Enterobacter sp., Acinetobacter sp.,...
*Pseudomonas* sp., and other Enterobacteriaceae species\(^{(8)}\). Currently, a major expansion of CTX-M type ESBL, whose preferred substrate is cefotaxime\(^{(16)}\), made this type in the most widely distributed, exceeding the incidence of SHV and TEM\(^{(10)}\).

In samples of Gram-negative bacilli isolated from blood cultures, there is an association between the emergence and spread of patients isolated with ESBL and multiple antibiotic resistance, particularly the 3rd and 4th-generation cephalosporins\(^{(9)}\), resulting in fewer treatment options, increased risk of failures in the treatment of patients infected with these strains, as well as morbiditymortality\(^{(9)}\).

ESBL production is not easily recognized by *in vitro* antimicrobial susceptibility testing, routinely performed in a clinical laboratory, confirmatory tests are needed to detect the phenotype and, in some cases, the resistance genotype\(^{(13)}\). These limitations in ESBL detection have hampered the hospital control of resistant microorganisms and favored the increase in prescription and overuse of antibiotics\(^{(21)}\). Furthermore, the involvement of ESBL-producing bacteria in the blood of hospitalized patients is directly related to complications in pathological conditions, worsens septicemia\(^{(17)}\).

Clinical and Laboratory Standards Institute (CLSI) recommends that screening and confirmatory assays are routinely performed to *Klebsiella pneumoniae*, *Klebsiella oxytoca*, and *Escherichia coli*, are considered the classic production of the enzyme, besides *Proteus mirabilis*\(^{(3)}\). However, other standardization can be used in phenotypic research on ESBL in isolates that are not covered by CLSI, including the British Society for Antimicrobial Chemotherapy Methodology (BSCA)\(^{(8)}\).

In view of the possibility of ESBL positive bacteremia and the importance of this hospital notification, the present study aimed to analyze the antimicrobial susceptibility profile of Gram-negative bacteria from positive blood cultures, as well as the frequency of ESBL-producing strains of patients of a hospital at Agreste Pernambucano, located in the city of Caruaru-PE.

**MATERIAL AND METHODS**

**Location of study and population**

In the period between February and June 2014, 100 samples from blood cultures of hospitalized patients were analyzed in a hospital at Agreste Pernambucano, located in the city of Caruaru-PE.

This study was approved by the Research Ethics Committee of the Caruaruense Association of Higher Education (Associação Caruaruense de Ensino Superior [ASCES]) under the number 421338.

**Samples collection and processing**

From the blood culture bottles provided by the Hospital Regional do Agreste (HRA), aliquots were aspirated for preparing a slide, which was stained by the Gram method, and for sowing in the medium Sheep Blood agar and MacConkey agar. Based on the colonies obtained, the phenotypic identification of the genus was carried out and, where possible, also the species, according to the macroscopic and microscopic characteristics of the colonies and with the results of biochemical tests of carbohydrates fermentation in Triple Sugar Iron agar (TSI), Sulphide Indole Motility (SIM), Simmons Citrate agar, Christensen Urea agar, and Instituto Adolfo Lutz medium (IAL).

**Determination of sensitivity pattern**

The identified strains were submitted to disc diffusion test, using broad-spectrum beta-lactam proposed by CLSI 2014. To test performing, the suspension of the bacteria in test, adjusted to McFarland 0.5 scale standard, was inoculated on the surface of a Müeller-Hinton agar plate. Subsequently, the ceftriaxone, cefotaxime, ceftazidime, cefepime and aztreonam discs were placed. The strains were evaluated for resistance according to CLSI 2014 guidelines, regarding their inhibition zones sizes\(^{(2)}\).

**Phenotypic detection of ESBL-producing strains**

ESBL enzyme research was performed by two phenotypic tests:

1) disc approximation – a plate of Müeller-Hinton agar was used, previously inoculated with the strain to be studied and adjusted to McFarland 0.5 scale. One amoxicillin/clavulanic acid disc (20 µg/10 µg) was placed in the center of the plate and around it the antimicrobial markers: cefotaxime/ceftriaxone (30 µg), ceftazidime (30 µg), cefepime (30 µg), and aztreonam (30 µg), at a distance of 20 mm, center to center in relation to the central disc. After incubation for 18-24 hours at 35-37ºC, the test interpretation was performed. The increase of the inhibition zone diameter or appearance of an empty area (distortion of the zone around the beta-lactam disc) indicated the presence of a ESBL-producing sample\(^{(8)}\);

2) disc replacement – consisted of placing two amoxicillin/clavulanic acid discs (20 µg/10 µg) on the surface of Müeller-
Hinton agar plate, and incubated for 1 hour at 35-37ºC in a bacteriological incubator, for diffusion of clavulanic acid in agar medium. Subsequently, the discs were removed with sterile forceps and, in the same location, were placed Ceftazidime (30 µg) and Cefotaxime (30 µg) discs. After appropriate incubation, the zones containing clavulanic acid were measured and compared with those of the discs containing only cephalosporin. In the presence of an ≥ 5 mm increase in the disc combined with the inhibitor, in relation to the disc without clavulanate, the strain was considered positive for ESBL(12).

RESULTS

Among the 100 samples obtained from blood cultures, in 30 Gram negative bacteria were identified and therefore, selected for the study. According to the interpretation of biochemical tests, bacterial samples were classified into five distinct species, with the predominance of *Proteus mirabilis* (23.33%), *Pantoea agglomerans* (20%), *Escherichia coli* (16.66%), and *Enterobacter cloacae* (10%), and a variety of other bacterial species, such as *Proteus penneri* (6.66%), *Acinetobacter* sp. (6.66%), *Enterobacter gergoviae* (6.66%), *Enterobacter sakazakii* (6.66%), and *Pseudomonas* sp. (3.33%) (Figure 1).

From the identification of the strains, the disc diffusion test was performed, and we observed that cephalosporins and monobactam showed the following resistance rates: ceftriaxone (80%), ceftazidime (56.66%), cefotaxime (86.66%), cefepime (60%), and aztreonam (60%) (Figure 2). Figure 2 shows the high resistance rates and low rates sensitivity of the isolates for the broad spectrum beta-lactam, widely used in clinical practice.

Regarding ESBL research, among the 30 bacterial strains, 73.33% (22) were positive by phenotypic tests, among them, 27.27% (six) *Pantoea agglomerans*, 18.18% (four) *Proteus mirabilis*, 13.63% (three) *Enterobacter cloacae*, 9.09% (two) *Proteus penneri*, *Enterobacter gergoviae* and *Acinetobacter* sp. and 4.54% (one) *Escherichia coli*, *Pseudomonas* sp. and *Enterobacter sakazakii* (Figure 3).

Whereas a strain is ESBL-producing when the test is positive for one or more of the employed substrates, it was found that the 22 samples tested were positive for the enzyme by phenotypic tests. The disc approximation method confirmed the presence of ESBL in 22 strains, all detected by cefotaxime/ceftriaxone, and 20 (66.66%) also positive for ceftazidime, aztreonam, and...
cefepime. The disc replacement test confirmed the presence of ESBL in 18 strains (60%), all positive for cefotaxime, and 16 (53.33%) were also positive with the ceftazidime substrate. Four samples corresponding to Pantoea agglomerans (one), Acinetobacter sp. (two), and Pseudomonas sp. (one) were positive to ESBL only for the disc approximation test, with all employed substrates (Table).

Evaluating the positive blood cultures by the presence of ESBL, we find that most of these cases occurred in female patients, with a prevalence of 59.10% ($n = 13$), while in males the prevalence was 40.90% ($n = 9$). Correlating the presence of ESBL with the place inside the hospital where the presence of bacteria was detected, from the blood cultures collected, it was noted that this enzyme was present most often in the Intensive Care Unit (ICU), showing prevalence of 73% ($n = 16$), when compared with its frequency in wards 27% ($n = 6$) (Figure 4).

**DISCUSSION**

Infections caused by ESBL-producing microorganisms have become a concern for the development of antimicrobial therapy, since these infections have major impact on the morbidity and mortality rates, and in hospital costs. This fact is due to the fact that the clinical implications of ESBL are serious, and it is essential to understand this mechanism resistance using sensitive and specific diagnostic techniques, in such a way as to properly conduct the treatment, monitor antimicrobial resistance, and implement intervention strategies(20).

In the present study, we observed a diversification of the isolated bacterial genus, with predominance of Proteus mirabilis (23.33%) and Pantoea agglomerans (20%). Gram-negative bacilli are among the main bacterial strains responsible for nosocomial infections, thus, it is important to study them in this environment, especially as regards the development of resistance mechanisms(19). In contrast to these results, Alves et al. (2012) analyzed 170 blood cultures of patients admitted to the ICU, from a hospital in São José dos Campos-SP, from January to July 2011, and found the following bacterial genus: Pseudomonas aeruginosa (12.2%), Klebsiella pneumoniae (8.1%), Acinetobacter baumannii (3.3%), Enterobacter cloacae (0.8%), Escherichia coli (0.8%), and Proteus mirabilis (0.8%) (1).

Dantas (2011) also found very low prevalence of Proteus mirabilis (1.85%) in nosocomial bacteremia, from May 2009 to February 2010 (4). This fact emphasizes the importance of this study results with respect to the current prevalence in the studied region, since this pathogen has significant potential for involvement in severe infections.

The susceptibility pattern of isolated strains for the beta-lactams showed high percentage of resistance, which can be linked

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>$n$ (%)</th>
<th>DA CTX/CRO (%)</th>
<th>DA CAZ/CPM/ATM (%)</th>
<th>DR CTX</th>
<th>DR CAZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pantoea agglomerans</td>
<td>6 (27.27%)</td>
<td>6 (27.27%)</td>
<td>6 (27.27%)</td>
<td>5 (27.77%)</td>
<td>5 (27.77%)</td>
</tr>
<tr>
<td>Proteus mirabilis</td>
<td>4 (18.18%)</td>
<td>4 (18.18%)</td>
<td>3 (13.63%)</td>
<td>4 (18.18%)</td>
<td>3 (13.63%)</td>
</tr>
<tr>
<td>Enterobacter cloacae</td>
<td>3 (13.63%)</td>
<td>3 (13.63%)</td>
<td>2 (9.09%)</td>
<td>3 (13.63%)</td>
<td>2 (9.09%)</td>
</tr>
<tr>
<td>Proteus penneri</td>
<td>2 (9.09%)</td>
<td>2 (9.09%)</td>
<td>2 (9.09%)</td>
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<tr>
<td>Enterobacter gergoviae</td>
<td>2 (9.09%)</td>
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<tr>
<td>Acinetobacter sp.</td>
<td>2 (9.09%)</td>
<td>2 (9.09%)</td>
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<tr>
<td>Escherichia coli</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
</tr>
<tr>
<td>Pseudomonas sp.</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Enterobacter sakazakii</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
<td>1 (4.54%)</td>
</tr>
</tbody>
</table>


![FIGURE 4 – Relationship between positive blood culture due to the presence of ESBL and the location of patient inside the hospital.](image)
to the overuse of third and fourth generation cephalosporins, as well as difficulty in hospital controlling resistant microorganisms, bringing consequences such as unsuccessful treatments and unfavorable clinical outcomes.

Cabrál (2011), analyzing the microorganisms resistance to beta-lactam antibiotics, reveal the true problems for the treatment of multiresistant isolates and established an association between the resistance profile and genes for beta-lactamases, since isolates of the same resistance pattern had the same genetic content, and thus, they were able to hydrolyze fourth generation cephalosporins and aztreonam(2).

The Clinical Microbiology Laboratory plays a key role in the detection of ESBL-producing Enterobacteriaceae, contributing to the orientation of a therapeutic approach and the application of preventive measures to control these nosocomial agents and isolation of patients, in order to minimize the spread of these pathogens(7, 12, 13).

In this study, the production of ESBL showed a frequency of 73.33%, and was mainly detected in Pantoaea agglomerans (27.27%), Proteus mirabilis (18.18%), and Enterobacter cloacae (13.63%), showing the dissemination of this resistance mechanism in the Enterobacteriaceae family. Phenotypic confirmatory methods are routinely performed for ESBL detection in Escherichia coli and Klebsiella sp., according to the CLSI, which recommends its screening only in these isolated, since they are considered the main producers. However, it is noteworthy that the increase on ESBL occurrence in other Enterobacteriaceae reveals the importance of extending this detection in other species than Escherichia coli and Klebsiella sp.(10).

Lago et al. (2010) reported that 49.6% of their isolated ESBL-producing were Enterobacter sp., showing that, since the pathogen is identified as Enterobacter sp., it is 8.43 times more likely to be ESBL-producing(10). In Japan, a study carried out by Datta et al. (2014) showed that in a number of 60 Enterobacteriaceae strains, 60% (n = 36) were identified as Proteus mirabilis and considered ESBL-producing, drawing attention to the fact these bacteria may not be detected by most clinical laboratories, suppressing some managements of assistance to the patient(10).

The sensitivity of phenotypic methods was satisfactory, of low cost, and thus feasible for use in the ESBL screening. However, we found that the disc replacement test was unable to detect the ESBL in four isolated, perhaps this was due to the fact that a smaller range of substrates was employed in the test, when compared to the disk approximation method. According to Karoline et al. (2008) and Wieand et al. (2007), the disc approximation method uses many antibiotics, so that it presents a sensitivity of 94.1% and specificity of 81.4%(8, 12). Martins et al. (2011) found in their study 91.6% of sensitivity for the disc replacement test(12).

Regarding the distribution of hospital infections by the ESBL-producing bacteria, it was observed that 73% of patients were admitted to the ICU, and, thus, showing higher risks of infections by these microorganisms, since, in this environment are performed: invasive procedures, prolonged hospitalization, and use of broad-spectrum antibiotics(15).

One limitation of this study was not using of the combined disc method recommended by the CLSI for phenotypic research on ESBL, due to the difficulty in obtaining the discs in the studied location. However, it is demonstrated the possibility of using alternative techniques in the study of this bacterial resistance mechanism.

CONCLUSION

ESBL frequency was high, prevailing Enterobacter genus as potential producer, demonstrating that the production of this enzyme in Enterobacteriaceae not belonging to the Klebsiella genus is no longer an exception. This reinforces the need for knowledge and use of techniques in its research. This high positivity rate may occur due to the selective pressure resulting in the indiscriminate use of broad-spectrum cephalosporins.

The techniques proposed in this paper proved to be useful for ESBL screening, with satisfactory sensitivity and low cost, and able to confirm this important resistance mechanism that hinders the beta-lactam antibiotics therapy used in clinical. We also observed the importance of testing a wider variety of substrates, thus increasing test sensitivity.

With the spread of bacterial strains in hospitals worldwide, it is necessary to know the prevalence of ESBL production, in an attempt to contribute to the formulation of a policy of therapy in high-risk units, where infection rates by multidrug resistant microorganisms are high. In addition, the knowledge of resistance standard in a geographical area, guide the proper and discerning use of antimicrobials.

ACKNOWLEDGEMENTS,

To the Microbiology Laboratory of the Hospital Regional do Agreste, for the supply of blood cultures that were analyzed, and to the Asces School, for the financial support.
RESUMO

Introdução: A emergência e a disseminação das isolados portadores de betalactamase de espectro estendido (ESBL) têm complicado o tratamento das infecções nosocomiais, uma vez que sua produção não é facilmente identificada nos testes de sensibilidade realizados rotineiramente em laboratórios clínicos, levando a dificuldades no controle hospitalar de microrganismos resistentes, bem como ao uso inadecuado de antibióticos. Objetivo: O objetivo deste trabalho foi analisar o perfil de resistência a e frequência de ESBL em bactérias Gram negativas isoladas de hemoculturas. Foram analisadas 100 amostras bacterianas, provenientes de hemoculturas de pacientes adultos, as quais foram identificadas fenotípicamente por provas bioquímicas de fermentação dos carbohidratos e submetidas à determinação do perfil de resistência por meio do teste disco difusão e da pesquisa de ESBL, pelos métodos de aproximação de diso e substituição de disco. Resultados: Entre as amostras bacterianas analisadas, 30 foram identificadas como bactérias Gram negativas, com predomínio de Proteus mirabilis, Pantoea agglomerans e Escherichia coli. Destas, 73,3% foram positivas para a pesquisa de ESBL pelos testes fenotípicos aplicados, sendo detectada, principalmente, em Pantoaea agglomerans, Proteus mirabilis e Enterobacter cloacae.

Conclusão: O aumento da ocorrência de ESBL em diversas enterobactérias revela a importância da ampliação desta detecção em espécies não Escherichia coli e não Klebsiella sp., de modo que as condutas de assistência ao paciente não sejam ameaçadas, já que essas bactérias resistentes podem não ser detectadas pelos laboratórios. Considerando a frequência de ESBL deste estudo, ressalta-se a importância de sua detecção, tendo em vista sua contribuição na possibilidade de melhorias das políticas de terapia em hospitais.

Unitermos: resistência bacteriana; betalactamase de espectro estendido; ESBL; infecção hospitalar.

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