Detection and characterization of multidrug-resistant enterobacteria bearing aminoglycoside-modifying gene in a university hospital at Rio de Janeiro, Brazil, along three decades

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Introduction: Multidrug-resistant Enterobacteriaceae, particularly those resistant to gentamicin, have become one of the most important causes of nosocomial infections.

Objective: We sought to investigate the presence of genes conferring resistance to aminoglycosides, specially to gentamicin, in Klebsiella pneumoniae and Escherichia coli multidrug-resistant strains isolated from different clinical materials among patients hospitalized in a university hospital in Rio de Janeiro, Brazil.

Materials and methods: Ten colonization strains and 20 infection strains were evaluated during three decades (1980 to 2010) using selective media containing 8 µg/ml of gentamicin. Thirty strains were tested for antimicrobial susceptibility. Twenty two strains were subjected to plasmid DNA extraction and 12 to hybridization assays using as probe a 1.9 kb plasmid DNA fragment from one of the K. pneumoniae strains isolated from faecal samples. This fragment was sequenced and assigned to the GQ422439 GenBank record. PCR was also performed using oligonucleotides designed for aminoglycoside-modifying enzymes.

Results: An accC2 acetylase, besides transposons and insertion sequences, were evidenced. Twenty-four (80%) of the isolates were positive for the accC2 gene in agreement with antibiotic susceptibility testing profiles, indicating the persistent presence of this gene throughout the three decades. We detected high molecular weight plasmids in 54.5% of the strains. Of the tested strains, 91% showed positive signal in the hybridization assays.

Conclusion: A gene codifying for one specific aminoglycoside-modifying enzyme was detected all throughout the three decades. Our data back the adoption of preventive measures, such as a more conscious use of antimicrobial agents in hospital environments, which can contribute to control the dissemination of microorganisms harboring resistance gene plasmids.

Key words: Enterobacteriaceae, infection; drug resistance, multiple, bacterial; enzymes, aminoglycoside, plasmids, acetyesterase.

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Verônica Dias Gonçalves and Françoise Bohrer-Lengruber contributed equally to the manuscript through their dissertations, which had the guidance and cooperation of the other six authors.
30 cepas en las que se determinó la resistencia a los antibióticos por medios fenotípicos. Veintidós muestras se sometieron a extracción de ADN plasmídico y se hicieron ensayos de hibridización con fragmentos de ADN plasmídico de 1,9 kb obtenido de una cepa de *K. pneumoniae* aislada de muestras fecales. Este fragmento fue secuenciado y correspondió al registro GQ422439 del GenBank. Se verificó la presencia de genes de enzimas modificadoras de aminoglucósidos mediante reacción en cadena de la polimerasa.

**Resultados.** En las cepas analizadas se evidenció la presencia de la acetilasa accC2, además de transposones y secuencias de inserción. Veinticuatro aislamientos (80 %) fueron positivos para el gen *aacC2* en concordancia con los perfiles de sensibilidad a los antibióticos, lo que indicó su persistencia a lo largo de las tres décadas. Se detectaron plásmidos de alto peso molecular en 54,5 % de las cepas. El 91 % de las cepas analizadas mostró signos positivos en las pruebas de hibridación.

**Conclusión.** Se detectó la persistencia de un gen codificador de una enzima modificadora de aminoglucósidos a lo largo de las tres décadas. Los resultados indican que las medidas de prevención, tales como un uso más responsable de los agentes antimicrobianos en el ambiente hospitalario, pueden contribuir al control de la diseminación de microorganismos que albergan plásmidos de genes de resistencia.

**Palabras clave:** Enterobacteriaceae, infección, farmacorresistencia bacteriana múltiple, enzimas, aminoglucósidos, plásmidos, acetilesterasa.

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In the last decades, multidrug-resistance of nosocomial bacteria has been a concern for healthcare professionals around the world (1,2). Through gene acquisition or mutation, bacteria can become resistant to one or several antimicrobial drugs. Gene acquisition involving horizontal transfer is a more frequent resistance mechanism than mutations, especially among enterobacteria, and it involves plasmid, gene cassette and transposon participation (3).

Besides resistance genes, resistance plasmids can carry other genes that also contribute to the pathogenicity of microorganisms (4). The integrons can house several resistance genes in Gram-negative bacteria. These genes may be found within resistance cassettes, and their different combinations can be generated by site-specific recombination (5). Transposons are also important in establishing bacterial resistance genes. The capacity of these mobile genetic elements to ‘circulate’ between plasmids and chromosomes is decisive for the dissemination of resistance-encoding genes to antimicrobial agents as beta-lactams and aminoglycosides (6,7). Aminoglycosides are highly potent, broad-spectrum antibiotics, widely used for the treatment of life-threatening infections. Resistance to these drugs occurs through several mechanisms that can coexist in the same cell; nevertheless, these resistance is often due to enzymatic inactivation by acetyltransferases, nucleotidytransferases (adenyllyltransferases), and phosphotransferases. The *accC2* gene is among the most frequently detected in strains of Enterobacteriaceae isolated from clinical samples (8). Many of these genes are associated with transposons, which help to the rapid dissemination of drug resistance across species boundaries (9).

In a previous study with Enterobacteriaceae strains of hospital origin we searched for genetic elements which express multidrug-resistance including resistance to aminoglycosides (10). Probes were built to detect plasmids encoding these multidrug-resistance traits. In the present work, we sought to investigate the presence of multidrug-resistant bacteria harboring aminoglycoside-resistance-encoding genes, particularly to gentamicin, in clinical specimens from a university hospital, and the persistence of these genetic elements during three decades. In another study, Gonçalves (11) detected the *aacC2* gene in strains of hospital origin, showing how the gene sequence was flanked by transposons and insertion sequences.

**Materials and methods**

*Isolated strains under study*

We analyzed 30 Enterobacteriaceae strains (18 *K. pneumoniae* and 12 *E. coli* strains) from 28 patients interned in a 600-bed tertiary university hospital in...
Rio de Janeiro, Brazil. Strains isolated from stool and urine samples of two of the patients were used. The strains were randomly selected from the multidrug-resistant Enterobacteriaceae (gentamicin-resistant) collected in the university hospital. Both infection and colonization strains were isolated from samples recovered in different years: Five colonization and three infection strains recovered in 1980 and 1981; four colonization strains in 1990; seven infection and one colonization strains in 2000, and 10 infection strains in 2010. The intestinal colonization strains, isolated in 1990 and 2000, were obtained from a primary streak in Eosin Methylene Blue Agar medium (EMB agar - Difco Laboratories, Detroit, MI) containing 8 µg/ml of gentamicin. The intestinal colonization strains and the infection strains recovered in 1980 and 1981 were isolated and identified by the hospital bacteriology laboratory.

**Antimicrobial susceptibility testing**

Antimicrobial susceptibility testing (AST) was performed using the agar diffusion method as set by the Clinical Laboratories Standards Institute (CLSI) (12). *Escherichia coli* strain ATCC 25922 was used as control, and the following antimicrobial agents were tested (suppliers’ individual concentrations appear in brackets): sulfamethaxazole-trimethoprim (St-23.75-1.25 µg), cephalotin (Cp- 30 µg), ceftazidime (Cz- 30 µg), cefoxitin (Fx- 30 µg), ciprofloxacin (Ci- 30 µg), chloramphenicol (Ch- 30 µg), ciprofloxacin (Cf- 30 µg), tetracycline (Tt- 30 µg) and ampicillin (Ap- 10 µg). Aminoglycosides tested were: gentamicin (Gn- 10 µg), amikacin (Ak- 30 µg), kanamycin (Kn- 30 µg), tobramycin (Tb- 10 µg), neomycin (Ne- 30 µg) and netilmicin (Nt- 30 µg).

Strains resistant to second and third generation cephalosporins in the AST were subjected to confirmatory tests for extended-spectrum beta-lactamases (ESBL) production using the double-disc synergy and the approximation tests as established by the CLSI (12).

Bacterial strains were grouped based on the results obtained in the susceptibility tests.

**Plasmid DNA extraction and agarose gel electrophoresis in wild bacterial strains**

Twenty two of 30 strains were randomly selected and subjected to DNA extraction and electrophoresis (in 0.8% agarose gel) for plasmid detection following the Kado and Liu protocol (13). We used the *E. coli* R861 strain plasmid DNA as molecular weight marker. Electrophoresis gels were dyed with ethidium bromide solution (0.5 µg/ml), analyzed in ultraviolet transiluminator and photographed with a Kodak EDAS 120 system.

**B2d DNA fragment cloning**

The 1.9 Kb B2d DNA fragment (GenBank accession number GQ422249) was obtained from Kp 401F10 plasmid digestion with restriction enzymes. The plamid was obtained from the *K. pneumoniae* 20 Kp strain isolated from the faeces of a surgical patient in the 80’s. B2d fragment sequencing was also performed to use it as a positive control for PCR and a probe for DNA hybridization.

**Automated DNA sequencing**

B2d DNA fragment gene sequencing was done according to Otto, *et al.* (14). Sequences were analyzed and compared using the BioEdit Sequence Alignment Editor software (15).

**PCR conditions**

The thermal cycling conditions were performed in a Cetus model 480 thermal cycler (Perkin-Elmer, Norwalk, CT). The primers used were: *aacC2* gene yielding a 237 bp product (5’-ACT GTG ATG GGA TAC GCG TC-3’ and 5’-CTC CGT CAG CGT TTC AGC TA-3’); *aadB* gene yielding a 320 bp product (5’-GAG CGA AAT CTG CCG CTC TGG-3’ and 5’-CTG TTA CAA CGT CTC TGG-3’ and 5’-CTG TTA CAA CGT CTC TGG-3’ and 5’-CTG TTA CAA CGT CTC TGG-3’), and *aacC3* gene yielding a 815 bp product (5’-AAA CTG GTG GCA ATA GAA GGA T-3’ and 5’- CTA TCC GTA TGA CGC TGA GTC3’), according to van de Klundert and Vliengenthart’s protocol (16). PCR assays were performed in a 50 µl total volume adding the following components to the reaction tubes: 1 µL of target DNA (obtained from dilutions of colonies in 50 µL of 10 mM Tris, 1 mM EDTA -pH 8.0), 1.5 mM MgCl$_2$, 0.2 mM of dNTP mixture (dATP, dTTP, dCTP and dGTP), 20 pmol of each primer, 1x PCR buffer, and 1.25U of Taq DNA Polymerase. The amplified products were subjected to electrophoresis in a 2% agarose gel.

**DNA hybridization assays**

For the DNA hybridization assays we used Thomas protocol (17): After electrophoresis in agarose gel, DNA extracted from *K. pneumoniae* and *E. coli* wild strains was transferred to a nylon membrane by capillarity system. The 1.9 Kb B2d DNA fragment was labeled with α[dATP] P$^{32}$ through random primed labeling (Gibco®, Life Technologies, US) and used as probe according to Feinberg and Vogelstein’s method (18).
Results

Identifying the antimicrobial susceptibility profiles

Antimicrobial resistance profiles were determined based on the results of the AST performed with aminoglycosides and other drugs. *K. pneumoniae* and *E. coli* strains AST for aminoglycosides indicated a high frequency of resistance to Gn, Kn and Tb (21 strains: 70%). Regarding the other antimicrobial agents, AST indicated a high frequency of resistance to Cx, Ch, Tt and Ap (17 strains: 57%) (table 1).

Fifteen strains were subjected to ESBL-production confirmatory tests and seven (46.7%) presented the ESBL phenotype. ESBL positive strains distribution was as follows: 1 (25%) from strains isolated in 1990; 1 (12.5%) from strains isolated in 2000, and 5 (50%) from strains isolated in 2010 (table 1). None of the strains isolated in 1980 or 1981 were ESBL positive. However, one strain (12.5%) isolated in this latter period, as well as one strain (12.5%) isolated in 2000, showed resistance to the third generation cephalosporin Cx (table 1).

Table 1. Characterization of resistance profiles to antimicrobials, and enzymatic and hybridization signals of *E. coli* and *K. pneumoniae* strains isolated from different material collected from hospitalized patients in a university hospital in Rio de Janeiro

<table>
<thead>
<tr>
<th>Strains</th>
<th>Mat</th>
<th>Origin</th>
<th>Antimicrobial profile</th>
<th>CBL</th>
<th>aacC2</th>
<th>HS</th>
</tr>
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<tr>
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<tr>
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<td>-</td>
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</tbody>
</table>

Kp: Klebsiella pneumoniae; Ec: Escherichia coli; Mat: Material; Fc: Faeces; Ur: Urine; Bl: Blood; DS: Different secretions; PF: Peritoneal fluid; TCT: Tenckhoff catheter tip; GS: General surgery; IE: Infectious diseases unit; GICU: General intensive care unit; NCU: Neonatal intensive care unit; VS: Vascular surgery; Nph: Nephrology; AU: Adolescent unit; A: Ambulatory; TS: Thoracic surgery; DIP: Infectious diseases unit; Gn: Gentamicin; Ak: Amikacin; Kn: Kanamycin; Tb: Tobramycin; Ne: Neomycin; Nt: Netilmicin; St: Sulfamethaxazole-trimethoprim; Cp: Cephalotin; Cf: Cephalozine; Fx: Cefoxitin; Ct: Cefotaxime; Cz: Ceftazidime; Ci: Ciprofloxacin; Nr: Norfloxacin; Ch: Chloramphenicol; Tt: Tetracycline; Ap: Ampicillin; CBL: Confirmatory test to ESBL; Neg: Negative result; Pos: Positive result; HS: Hybridization signal; -: Data not observed for this strain.
**Plasmid profile of strains**

Out of the 22 strains randomly selected, 14 corresponded to *K. pneumoniae* and eight to *E. coli*. They were subjected to plasmid DNA extractions with the following results: Nine strains (41%) had plasmids with molecular weights of 147 kb or more, and plasmids of around 7 kb (figure 1). Strains 9 Ec, 10 Ec and 11 Ec isolated in 1990 had similar electrophoretic patterns, as well as strains 18 Kp, 20 Kp and 22 Kp isolated in 1980 and 1981, while only strain 38 Kp from 1990 and strains 2 Kp, 5 Kp and 13 Kp from 2000 showed such similarity.

**Analysis of DNA sequences**

DNA sequencing was performed for the B2d DNA fragment originated from plasmid 20 Kp. The 1.9 kb repeat sequence obtained indicated the presence of DNA sequences homologous to the *aacC2* gene or to the aminoglycoside-(3)-N-acetyltransferase (AAC (3) II). The BLAST™ analysis of the B2d fragment allowed us to identify a subset of repeats that showed over 90% of overall identity and E values <0.05 when compared to other GenBank records, mostly related to acetyltransferases, insertion sequences and transposons found in Gram-negative bacteria (figure 2). The most significant sequences aligned to the B2d fragment were: *Salmonella enterica* subsp. *enterica* serovar Choleraesuis TnpA_rve (GI: 161867979); *Escherichia coli* IAI39 transposase, IS26 (GI: 218700153); *Acinetobacter baumannii* AB0057 transposase IS26 (GI: 213155651); *Escherichia coli* AAC(3)-II (GI: 41056930), and *Citrobacter freundii* aminoglycoside acetyltransferase (GI: 27383509).

**Detection of sequences coding for aminoglycoside modifying enzymes**

Amplification products indicated the presence of the *aacC2* gene (GenBank access number X51534) in 13 (72.2%) *K. pneumoniae* strains and nine (75%) *E. coli* strains (table 1). We did not detect amplification for the other genes under study.

**Hybridization profiles**

Plasmid DNA was extracted from 12 strains which were subsequently subjected to DNA hybridization assays using the B2d fragment as probe (data not shown). Positive signals were obtained for 11 (91.7%) of the 12 strains tested (table 1). Regarding the extracts of Kp 18 and Kp 38 strains isolated in 1981 and 1990, respectively, hybridization signals were detected in the chromosome. Strain Kp 5, isolated in 2000, and strain Ec 11, isolated in 1990, showed more than one band, indicating that the gene sequences identified in the B2d fragment were present in several regions of the bacterial genome. In the other strains with more than one plasmid, the hybridization signal was detected in just one of the plasmid bands.

**Discussion**

To characterize the resistance profile of the isolated strains, we selected different classes of antimicrobial agents: those which have been used for a long time, as well as some classes introduced more recently in therapeutics, such as beta-lactam antibiotics and aminoglycosides, important antimicrobial drugs used in hospital environments to which microorganisms have increasingly become resistant (19,20). ESBL-producing Gram-negative bacteria have been detected since the 1980s as important causes of nosocomial infections (21). After the introduction of expanded-spectrum cephalosporins, *K. pneumoniae* and *E. coli* resistant strains emerged in certain areas of the world (22). In fact, this study detected multidrug-resistant, aminoglycoside resistant and ESBL-producing strains in the different decades. Among those isolated in the 80’s, despite the absence of ESBL-producing strains, we detected resistance to third generation cephalosporins.

Aminoglycosides susceptibility profiles in strains isolated between September 1980 and January 1981 were similar to those in strains isolated in 1990, 2000 and 2010. TSA results agreed with
those described in the literature for the acetylase identified as AAC(3)-II (Gen/Tob/Net) in most of the strains (8). Small changes in DNA sequences in genes throughout the replication processes, and/or acquired genes, may have resulted in these phenotypical differences.

Several environmental and nosocomial bacteria can contain transferable plasmids that carry genes expressing several traits, one of which is multidrug-resistance to antimicrobial drugs (23-25). In fact, we found plasmids with molecular weights varying between 147 kb or more and 7 kb or less in multidrug-resistant strains isolated from specimens of patients hospitalized in different units and in different decades. These genetic elements can be transferred from a microorganism to another in vivo, vertically or horizontally, contributing to worsen hospital infections (3).

The similarity of plasmid profiles in strains isolated in different decades indicates the persistence of bacterial clones and/or of specific plasmids. Well-established populations of bacterial clones can extend their resistance phenotypes by acquiring genetic elements (plasmids, integrons and transposons), facilitating co-selection processes under different levels of pressure and favoring the permanence of these microorganisms and their genetic constituents in the environment (26). The possibility of co-transmission may not only contribute to increase resistance markers, but also confer an evolutionary benefit to these strains, leading to selection in an environment with persistent antibiotic pressure by beta-lactams, aminoglycosides and quinolones (3,24,27).

The B2d clone sequencing showed that, besides sequences compatible with the aacC2 gene (ACC (3)–II), there are other sequences compatible with transposons, insertion sequences and plasmid-mediated resistance to macrolides and other antimicrobial drugs. With respect to the presence of more than one hybridization signal observed in two strains (Kp 5 and Ec 11), we considered the possibility of different insertion events occurring throughout several generations of bacteria, which may have involved different plasmids in the same bacterial cell. We observed how these DNA sequences can determine DNA insertions in one or more genomic sites, originating resistant strains, irrespective of the presence of original mobile genetic elements. When we compared sequence frames with those already described in the GenBank, we were able to detect identity regions in microorganisms from different species, indicating the possibility of horizontal resistance gene transfer involving plasmids, transposons and insertion sequences. When analyzing 160 strains of gentamicin-resistant E. coli isolated from human and animal samples, Ho, et al. (28), identified the aacC2 gene in 81.3% of them, thus confirming this possibility.

Based on the results obtained with the hybridization and PCR assays, it can be said that genetic elements housing multidrug-resistance encoding sequences presumably remain in circulation among microorganisms in hospital environments for long periods of time, and that these microorganisms may colonize and cause infections in hospitalized patients. Isolated strains were tested not only for the same period as the Kp 20 strain from which the fragment used as probe was extracted (1980/81), but in those isolated 10, 20 and even 30 years later (1990, 2000 and 2010, respectively). Selective pressure can be acting in favor of the persistence of several genes grouped in one operon.

Figure 2. Linear map of the partial sequence of the B2d DNA fragment originated from the plasmid 20 Kp examined in this study showing sequences compatible with the aacC2 gene (ACC (3)–II), as well as transposons, insertion sequences and plasmids of resistance to macrolides and other antimicrobial drugs.
that, otherwise, could be eliminated. Besides, it is important to note that gene cassettes can contain many of the resistance genes expressed in Gram-negative bacteria (6).

The Kp 20 strain was originally isolated from faeces of a patient admitted in a general surgery ward. It is known that intestinal microbiota components are affected during antimicrobial therapeutics decreasing their number, and favoring proliferation of opportunistic microorganisms which can disseminate and cause infections (29). The human intestine provides an important reservoir for multidrug-resistant Gram-negative bacteria, including Enterobacteriaceae species involved in infectious processes both in community and nosocomial environments, and the use of antimicrobial agents is one of the important factors for the selection of multidrug-resistant microorganisms (20,30).

We want to emphasize the importance of reducing hospital stays, as well as of a more discerning and conscious use of antimicrobial agents in inpatient and outpatient hospital environments, and of educational programs aimed at updating healthcare professionals, especially on the importance of adequate hand washing before and after patient care, to name only some of the measures to control the persistent dissemination of microorganisms with resistance genes and plasmids.

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Conflict of interest

The authors declare no conflicts of interest.

References


