**Infection related to Klebsiella pneumoniae producing carbapenemase in renal transplant patients**

**ABSTRACT**

**Objective:** To evaluate the risk factors related to Klebsiella pneumoniae carbapenemase infection after renal transplantation. **Methods:** This was a retrospective epidemiological (case-control) study, conducted from October 2011 to March 2016. Transplanted patients with infection by this bacteria during hospitalization were selected as cases. The controls were paired by age, sex, type of donor and transplant time. The proportion of cases and controls was 1:2. **Results:** Thirty hundred and five patients were included in the study (45 cases and 90 controls). The risk factors found for infection by KPC were: time of hospitalization after the transplant (OR: 4.82; CI95% 2.46-9.44), delayed kidney function (OR: 5.60; CI95% 1.91-11.01) and previous infectious for another microorganism (OR: 34.13 CI95% 3.52-132.00). **Conclusion:** The risk of acquisition of this bacterium was directly related to invasive procedures and exposure to the hospital environment. The findings reinforce the importance of prevention measures and control of infection by this microorganism.

**Descriptors:** Infection; Infection Control; Kidney Transplantation; Bacteria; Nursing Care.

---

Monica Taminato¹
ORCID: 0000-0003-4075-2496

Dayana Fram¹
ORCID: 0000-0001-6366-2325

Rogério Rodrigues Floriano Pereira²
ORCID: 0000-0003-0560-1422

Ricardo Sesso¹
ORCID: 0000-0002-1062-0073

Angélica Gonçalves Silva Belasco¹
ORCID: 0000-0002-1062-0073

Antonio Carlos Pignatari³
ORCID: 0000-0003-0560-1422

Dulce Aparecida Barbosa¹
ORCID: 0000-0002-9912-4446

¹Universidade Federal de São Paulo. São Paulo, Brasil.
²Universidade de São Paulo. São Paulo, Brasil.
³Universidade Federal do ABC. Santo André, São Paulo, Brasil.

How to cite this article:

Corresponding Author:
Monica Taminato
E-mail: taminatomonica@gmail.com

Submission: 01-11-2019 Approval: 03-23-2019

---

How to cite this article:


Corresponding Author:
Monica Taminato
E-mail: taminatomonica@gmail.com

Submission: 01-11-2019 Approval: 03-23-2019

---

RESUMEN

**Objetivo:** Evaluar los factores de riesgo relacionados con la infección por Klebsiella pneumoniae carbapenemasa después del trasplante renal. **Método:** Estudio retrospectivo epidemiológico (caso-control), realizado de octubre de 2011 a marzo de 2016. Pacientes transplantados con infección por esta bacteria durante la internación fueron seleccionados como casos. Los controles fueron pareados por edad, sexo, tipo de donador y tiempo de trasplante. La proporción de casos y controles fue de 1:2. **Resultados:** Treinta y cinco pacientes fueron incluidos en el estudio (45 casos y 90 controles). Los factores de riesgo para la infección por KPC fueron: tiempo de hospitalización después del trasplante (OR: 4,82; IC95% 2,46-9,44), función renal retardada (OR: 5,60; IC95% 1,91-11,01) y anterior infecciosa para otro microorganismo (OR: 34,13 IC95% 3,52-132,00). **Conclusión:** El risco de exposición a esta bacteria fue directamente relacionado a procedimientos invasivos y exposición al ambiente hospitalario. Los hallazgos refuerzan la importancia de medidas de prevención y control de la infección por este microorganismo.

**Descritores:** Infecciones; Control de Infecciones; Transplante Renal; Bacterias; Cuidados de Enfermería.

---

RESUMEN

**Objetivo:** Evaluar los factores de riesgo relacionados con la infección por Klebsiella pneumoniae carbapenemasa después del trasplante renal. **Método:** Estudio retrospectivo epidemiológico (caso-control), realizado de octubre de 2011 a marzo de 2016. Pacientes transplantados con infección por esa bacteria durante la internación fueron seleccionados como casos. Los controles se parearon por edad, sexo, tipo de donante y tiempo de trasplante. La proporción de casos y controles fue de 1:2. **Resultados:** Treinta y cinco pacientes fueron incluidos en el estudio (45 casos y 90 controles). Los factores de riesgo para la infección encontrados por KPC fueron: tiempo de hospitalización después del trasplante (OR: 4,82; IC95% 2,46-9,44), función renal retardada (OR: 5,60; IC95% 1,91-11,01) y anterior infecciosa para otro microorganismo (OR: 34,13 IC95% 3,52-132,00). **Conclusión:** El risco de exposición a esta bacteria fue directamente relacionado a procedimientos invasivos y exposición al ambiente hospitalario. Los hallazgos refuerzan la importancia de medidas de prevención y control de la infección por ese microorganismo.

**Descritores:** Infecciones; Control de Infecciones; Transplante de Riñón; Bacteria; Atención de Enfermería.
BACKGROUND

Management of infections caused by multidrug-resistant bacteria impacts considerably on health costs and becomes major modifier of health expenses in the ongoing antibiotic resistance crisis(10).

*Klebsiella pneumoniae* carbapenemase (KPC), has become one of the most important contemporary pathogens, among specific populations include solid organ transplantation (SOT), immunosuppressant therapy, mechanical ventilation, prolonged use of invasive devices, use of antimicrobial agents, and a high Acute Physiology and Chronic Health Evaluation score (2,7).

Gram negative bacteria are important pathogens related to healthcare associated infections (HAIs) represents a challenge for the control and treatment of infections related to this microorganism, as well as for the global dispersion of this agent (3-4). Patients undergoing renal transplant are at risk for infection with multidrug-resistant organisms related to the surgical procedure, necessity for invasive devices, clinical complications and immunosuppression, specially in renal transplant was identified as a risk factor for the acquisition of KPC (2,7,8).

SOT recipients infected with KPC-producing Enterobacteriaceae, mortality can be as high as 71 % (2,6,11). Among reporting that such infection occurred more frequently in the first 2 months after Kidney Transplantation, with a combined incidence of 13 % in that period, and that the urinary tract was the most common site of infection. In those studies, mortality associated with KPC-HAI ranged from 33 to 50 % (9-11).

The CDC has offered recommendations since 2009 that indicate efforts to contain dissemination, exchange information among the services, and include experiences to improve the approach to the bacteria (12-13).

The first report of KPC in Brazil occurred in 2006, with the identification of *K. pneumoniae* KPC-2 in an intensive care unit (ICU)(14).

Brazil is part of the SENTRY study, a program of antimicrobial surveillance that showed a significant increase in *K. pneumoniae* KPC-2 identification between 2008-10. In 2008, isolates of KPC were not found, 10 samples were isolated in 2009, and 44 samples were positive for KPC in 2010 in the three participating Brazilian centers (15). This study, together with other findings, showed that KPC is endemic in Brazil and that this bacteria was encountered in the waste water of hospitals in the country (16).

Facing this epidemiological panoramic of KPC, the gravity of the situation, and the fact that transplant patients are considered to be at risk for acquisition of multidrug-resistant bacteria.

OBJECTIVE

To evaluate the risk factors, investigated the epidemiology, clinical outcomes and mortality associated to KPC infection in patients undergoing renal transplant.

METHODS

Ethical aspects

The study was preceded by approval of the Ethics Committee of the Universidade Federal de São Paulo (Federal University of São Paulo, UNIFESP).

**Design, period and location of study**

This was a retrospective study (case-control), conducted from October 2011 to march 2016, in the Hospital do Rim e Hipertensão da Fundação Oswaldo Ramos (HRim/FOR), a university hospital and a worldwide reference in renal transplant which performs approximately 900-1000 renal transplants/year. This institution follows rigorous standards for medical record keeping.

**Patients**

Transplant recipients, regardless of the time of renal transplantation, who in the occasion of hospitalization developed infection with positive culture for *Klebsiella pneumoniae* carbapenemase (KPC), identified by active surveillance. Active surveillance of multidrug-resistant organisms is routinely performed by the Commission for the Prevention and Control of Infection of the HRim/FOR with patients hospitalized in this service (17). Throughout the study period, surveillance cultures (perineal– rectal swab samples in Stuart's transport medium) were collected from all patients admitted from other hospitals, as well as on a weekly basis from patients in intensive care units (ICUs).

Were considered cases: renal transplant recipients (TX), aged > 18 years; those who presented with infection at the initiation of the study with positive culture for KPC in one or more of the following samples: blood culture, urine culture; broncho-alveolar lavage; wound secretion with positive which KPC. Only the first infection was included for risk factors analysis, occurrence of death and clinical variables of interest of the case group and control were recorded for a period of one year prior to the infection by KPC.

Were considered controls: kidney transplant patients at the same institution and period of transplant, with age, sex, time of transplant (in days, the closest to the respective case), and donor type (live or deceased) similar to the cases, without positive culture for KPC or other type of infection at the initiation of the study. The ratio between the cases and controls was 1:2.

**Protocol and data collection**

Data collected from patient files considered cases and controls were: sociodemographic characteristics, etiology of end stage renal disease (ESRD), laboratory tests at entry into the study, results of cultures for microorganisms according to the location of standardized collection, type and duration of dialysis treatment prior to transplant, clinical complications and death, if that occurred. For identification of the risk factors, the parameters were monitored for a period of six months preceding the date of infection or colonization by KPC, and for the controls, the six months prior to the index date of enrollment was considered as the date of the pairing with the data of the TX with the respective case. The occurrence of death in both groups was verified until one week after infection by KPC.

Laboratory Methodology: For the detection of carriers of KPC, samples of blood, urine, bronchoalveolar lavage and surgical wound secretions (surgical site infection) were collected. Biological materials were sent to the Clinical Microbiology Special Laboratory (LEMC) of the Discipline of Infectious Diseases of UNIFESP, where microbiological testing was performed in accordance with the recommendations of the Clinical and Laboratory Standards Institute (CLSI) (20), which include
isolation, identification, culture and antimicrobial sensitivity testing. Carbapenem-resistant strains of K. pneumoniae were identified with an automated susceptibility testing system (VITEK; bioMérieux, Marcy l’Étoile, France). Minimum inhibitory concentrations (MICs) were interpreted according to the Clinical and Laboratory Standards Institute breakpoints (18).

**Statistical analysis**

A descriptive analysis of the cases and controls was conducted which examined the variables of interest. Data are presented using absolute frequencies and percentages for categorical variables, and mean ± standard deviation for continuous variables. A descriptive analysis of groups of case versus control patients was conducted, considering the demographic, clinical, and laboratory parameters related to treatment. The primary results were the presence of infection by KPC at some isolated sites. The univariate analyses were performed, comparing the variables between the groups (case and control). The association between infection and categorical variables were tested with the chi-square or Fisher’s exact test, and the association between continuous variables and the presence of infection was done using the Student t-test, as appropriate. In the analysis of the associations, the odds ratios with confidence intervals at the 95% level were calculated. A multivariate analysis using logistic regression to investigate factors associated with infection/colonization was also performed. In this final analysis, the dependent variable was presence of infection by KPC, and the independent variables tested were those who present a value of p<0.20, in the initial univariate analyses. All analyses were performed using the Statistical Package for Social Sciences software (SPSS 17.0, Chicago, IL, USA).

**RESULTS**

In the 45 patients considered cases, 277 samples were identified as positive for KPC. The site of KPC isolation in 62 % of the samples was the urine culture, 25 % blood culture, 9% surgical site infection (wound secretions) and 4% in tracheal aspirate.

The sociodemographic characteristics, etiology of chronic renal disease (ESRD), comorbidities, type and duration of dialysis treatment before transplant, and the laboratory tests at entrance into the study of patients included in the study are presented in Table 1. There was no statistically significant difference between variables studied in the cases compared to controls and both group were considered homogeneous.

Infection by KPC in patient cases occurred in 73% of the occasions during hospitalization for kidney transplant, and, in 27% of episodes that occurred in the post-transplant hospitalizations related to clinical events during the first year after the transplant in the followed up period.

**Table 1 - Sociodemographic characteristics, etiology of chronic renal disease (ESRD) and clinical variables of interest of the case group and control, São Paulo, Brazil, 2017**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Cases (n=45)</th>
<th>Controls (n=90)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>47.20±12.11</td>
<td>46.12±12.31</td>
<td>0.325</td>
</tr>
<tr>
<td>Male %</td>
<td>28 (61%)</td>
<td>54(59%)</td>
<td>0.974</td>
</tr>
<tr>
<td>Etiology</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glomerulonephritis</td>
<td>18 (40%)</td>
<td>41(45%)</td>
<td>0.304</td>
</tr>
<tr>
<td>Polycystic kidneys</td>
<td>10 (22%)</td>
<td>18(20%)</td>
<td>0.962</td>
</tr>
<tr>
<td>Undetermined</td>
<td>4 (10%)</td>
<td>20(22%)</td>
<td>0.038</td>
</tr>
<tr>
<td>Others</td>
<td>3(9%)</td>
<td>11(13%)</td>
<td>0.421</td>
</tr>
<tr>
<td>Comorbidities</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>23(52%)</td>
<td>41(45%)</td>
<td>0.441</td>
</tr>
<tr>
<td>DM</td>
<td>6 (13%)</td>
<td>13(14%)</td>
<td>0.987</td>
</tr>
<tr>
<td>Other*</td>
<td>16(35%)</td>
<td>46(41%)</td>
<td>0.369</td>
</tr>
<tr>
<td>Type of dialysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemodialysis</td>
<td>41 (93%)</td>
<td>85 (94%)</td>
<td>0.953</td>
</tr>
<tr>
<td>Peritoneal dialysis</td>
<td>3(6%)</td>
<td>3(3%)</td>
<td>0.235</td>
</tr>
<tr>
<td>Conservative</td>
<td>1(1%)</td>
<td>2(3%)</td>
<td>0.378</td>
</tr>
<tr>
<td>Time of dialysis, preceded months</td>
<td>51.32±43.23</td>
<td>49.92±43.11</td>
<td>0.241</td>
</tr>
<tr>
<td>Time of transplantation, months</td>
<td>23.90±19.28</td>
<td>19.91±18.32</td>
<td>0.117</td>
</tr>
<tr>
<td>Urea, mg/dl</td>
<td>133 ± 46</td>
<td>116±43</td>
<td>0.221</td>
</tr>
<tr>
<td>Creatinine, mg/dl</td>
<td>7.34±3.27</td>
<td>6.61±3.21</td>
<td>0.067</td>
</tr>
<tr>
<td>Leukocytes /mm3</td>
<td>740±7348</td>
<td>8831±7776</td>
<td>0.085</td>
</tr>
<tr>
<td>Hemoglobin g/ℓ</td>
<td>11.05±1.75</td>
<td>11.5±1.51</td>
<td>1</td>
</tr>
<tr>
<td>Hematocrit %</td>
<td>34.4±5.86</td>
<td>36.96±5.62</td>
<td>0.083</td>
</tr>
</tbody>
</table>

*Variables: neoplasm and systemic lupus erythematosus.

**Table 2 - Characteristics of renal transplant and previous clinical complications of infection by KPC between cases and controls, São Paulo, Brazil, 2017**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Simple logistic regression</th>
<th>Multiple logistic regression</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Type of donor</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deceased donor</td>
<td>40 (88%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Time of hospitalization after TX*, days</td>
<td>51.31±42.26</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Delayed kidney function</td>
<td>23 (49.3%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Thymoglobulin induction</td>
<td>8.2 (18%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Basiliximab induction</td>
<td>10 (22%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Mycophenolic acid</td>
<td>9 (18.8%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Calcineurinhibitors</td>
<td>7.65 (17%)</td>
<td>30 (65%)</td>
</tr>
<tr>
<td>Previous infection</td>
<td>30 (65%)</td>
<td>22.5 (25%)</td>
</tr>
<tr>
<td>Acute rejection</td>
<td>26 (57%)</td>
<td>17 (19%)</td>
</tr>
<tr>
<td>CMV infection</td>
<td>14 (29%)</td>
<td>13 (19%)</td>
</tr>
</tbody>
</table>

*TX: transplante.
In univariate logistic regression analysis the most significant events were: time of hospitalization after transplant (OR: 5.61, 51.3±42.2 days versus 2.46±9.44 days, p<0.001), delayed kidney function (OR: 7.85, 49.3% versus 11%, p<0.001) and use of mycophenolic acid (OR: 3.20, 18.8% versus 6.5%, p<0.006). Clinical events were: occurrence of previous infection by other microorganisms (OR: 53.68, 65% versus 25%, p<0.001), acute rejection (OR: 4.96, 57% versus 19%, p<0.004) and cytomegalovirus infection (OR: 7.60, 29% versus 19%, p<0.003).

Twelve (27%) deaths occurred due to KPC infection in patient cases. The control group there was no death related to KPC or other infectious causes during the study period (p<0.001).

In the multivariate analysis using logistic regression, risk factors associated with infection by KPC were identified in Table 2. The parameters that were significantly associated with the outcome of KPC infection were: time of hospitalization (OR 4.82, CI 2.46-9.44, p<0.001), delayed kidney function (OR 5.60, CI 1.92-11.01, p<0.001), previous infections by another microorganism (OR 34.13, CI 3.52-132.00, p<0.001) and CMV infection (OR 9.20 – 95% CI 2.17 – 24.29, p< 0.005).

Discussion

The analysis of risk factors for infection by KPC among renal transplant patients was the first proposition of the study. The characteristics of the groups of cases and controls were homogeneous in regard to age, sex, type of dialysis treatment, donor type and clinical conditions to analyze the exams of biochemical input.

The risk factors found for infection by KPC among patients was the length of hospital stay after transplant, delayed kidney function and requiring dialysis after transplantation and clinical complications such as previous infectious episodes and invasive cytomegalovirus that lead to invasive diagnostic procedures and greater use of antimicrobials during six months after the transplant.

In renal transplant patients, infections consist of one of the principal causes of hospital readmissions, and represent 51% of admissions that occur within six months after transplantation, preceded only by surgical complications (19). Despite the development of surgical techniques and immunosuppressive schemes, infections remain the second leading cause of mortality among renal transplant patients (19-20).

Previous studies with other patient populations demonstrated that hospitalization in ICU, prolonged length of hospital stay, invasive procedures, solid organ transplant and recent exposure to broad-spectrum antibiotics were directly related to infection/colonization by KPC (6-7,9, 21-25).

In the present study the mortality associated with infection KPC was 27%, a lower rate than other studies as high as 71% in vulnerable population, solid organ transplantation (7,9,11).

The lower mortality rate (27%) found in this study suggested that in this service, early detection and intervention occurred. Other data that reinforced the measures of early intervention were a form of identification of the bacteria in the study population, in which 41% were obtained through surveillance of cultures.

A recent literature review about global epidemiology of KPC showed that in several studies the mortality rate was found to be similar to that found in the present study, and ranged from 13 to 34% for the groups who had early identification of bacteria and combined antimicrobial therapy (21). McNeil et al. have evaluated the implementation of surveillance cultures for patients who were scheduled to undergo renal transplantation, and demonstrated that patients colonized before transplant have a higher morbidity when compared with those non-colonized patients (26).

Routine rectal swab surveillance of KPC contacts is an important measure to enhance identification and isolation of carriers as well as with other interventions are more likely to be successful included minimizing use of invasive devices, promotion of antimicrobial stewardship, a standardized approach for active surveillance of at risk populations, improve the hand hygiene adherence and protocols for discontinuation of carrier status (2,27-30).

The high risk of acquiring and disseminating KPC in renal transplant patients is recommended as a component of infection control programs and the development of safe techniques for the rapid identification of this microorganism and the qPCR technique demonstrated in provides better results compared to the culture method and allows the rapid implementation of control measures and interventions to reduce the spread of health services (33).

Confronted by infectious complications in patients with ESRD, Barbosa et al. (34) evaluated the prevalence of colonization with vancomycin-resistant Enterococcus (VRE) in 300 patients on dialysis and 280 transplant recipients treated at Hospital do Rim e Hipertensão da UNIFESP. A prevalence rate of 14.5% was verified in dialysis, and of 14% in renal transplant recipients. With molecular typing of these samples, the important detection of cross-transmission of VRE among patients treated at the Serviço de Dialise e Transplante da UNIFESP (Dialysis and Transplant Service of UNIFESP) was possible (35).

Cytomegalovirus (CMV) infection is the important infectious complication in kidney transplantation, specially in the first year and it is one reason for high morbidity and mortality rates. Infection for CMV increasing the risk factors for infection by KPC related to hospitalization and invasive procedures (31).

This becomes even more important when recent studies suggest there are no differences in the survival of the patient and of the graft in renal transplant recipients receiving azathioprine or mycophenolate in combination with tacrolimus and steroids, indicating that azathioprine should be considered a therapeutic option for a selected group of patients (32).

It was observed that most donors were deceased, reflecting the national and international policies for organ donation stimulus. In recent systematic review and meta-analysis published has revealed that deceased kidney donor recipients have 20% higher risk for developing infections (36).

The risk factors found in this study should lead to greater vigilance in actions with this population, from hospital admission to discharge, especially the establishment of measures of prevention and control of infection, in particular the implementation of bundles that include washing of hands, surface cleaning, education of health staff, surveillance cultures, bathing with chlorhexidine, and contact precautions are among the key measures described to contain the spread of this bacteria (2,27,30).

Limitations of the Study

We highlight as a study limitation the fact that the majority of the patients had renal replacement therapy in other services, and because of this further analysis was not possible to deepen the analysis. Another aspect that must be emphasized is that, because it was a retrospective...
study that made use of information obtained in patients’ records, the collection and insertion of data in the database analysis were confirmed by two researchers, aiming at reliability of the results.

Contributions to the Area

As implications for practice, the need for adherence to preventive measures against infections related to health care, in particular hand hygiene, are emphasized. Early identification and infection control measures for patients at higher risk of infection as delayed kidney function and risk for infection for CMV. Identification of patients colonized by KPC in this study was probably an important measure for the reduction of mortality rates, as compared to the literature.

Conclusions

The main risk factors for infection by KPC were: time of hospitalization after the transplant, delayed kidney function and clinical events that lead to the need for invasive procedures, and consequently greater exposure to the hospital environment.

Additional measures include minimizing use of invasive devices, promotion of antimicrobial stewardship, a standardized approach for active surveillance of at risk populations, and protocols for discontinuation of carrier status.

Competing interests

The authors declare that they have no competing interests.

Acknowledgements

We wish to acknowledge the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (Capes) and Fundação de Amparo à Pesquisa do Estado de São Paulo for the financial support for this study.

REFERENCES

Infection related to Klebsiella pneumoniae producing carbapenemase in renal transplant patients
Taminato M, Fram D, Pereira RRF, Sesso R, Belasco AGS, Pignatari AC, Barbosa DA.

from: https://doi.org/10.1053/j.ajkd.2004.04.038
