Review Article

Dissemination of carbapenemases in Brazil: an integrative review

Comment [ДС1]: Where? In hospitals? Outpatients? Animals?

ABSTRACT

Antimicrobial resistance is a natural biological phenomenon. As antimicrobial agents become incorporated into clinical practices, strains of resistant microorganisms are detected in laboratories. The most frequently identified groups of carbapenemases include KPC, NDM, and OXA-48, which limit the available treatment options. The rapid spread of acquired resistance to carbapenems is increasingly facilitated by mobile genetic elements, such as plasmids, which transfer carbapenemase genes both intra- and interspecies among gramnegative bacilli (GNB). The β-lactamase genes present in mobile genetic elements play a crucial role in the rapid global spread of antibiotic-resistant GNB. This study presents an integrative review of the literature from the scientific databases of the National Library of Medicine (PubMed) and the regional portal of the Virtual Health Library (BVS), comprising scientific articles published between 2020 and 2024 to assess the epidemiological situation of carbapenemases in Brazil. After applying the inclusion and exclusion criteria, 26 articles were selected, covering different Brazilian states. The studies analyzed revealed that the state of Pernambuco in northeastern Brazil is noteworthy for reporting the first identification of the blavim-2 gene in Acinetobacter baumannii isolates, as well as the identification of the blaoxa-23like and blaoxa-143 genes. It is also noteworthy that the first description of the association of blaNDM1 and blaKPC-2 in Proteus mirabilis and Serratia marcescens, as well as the presence of the blandm gene in Klebsiella aerogenes, was recorded in this state. The cohabitation of multiple genes resistance in microorganisms isolated from several Brazilian states is of particular importance, with the KPC carbapenemase producer gene being the most prominent. Understanding this genetic diversity is crucial for the development of effective control and prevention strategies aimed at mitigating the impacts of antimicrobial resistance in the country.

Keywords: Carbapenemases, Carbapenem-Resistant Enterobacteriaceae, Carbapenem Resistance, Gram-Negative Bacilli, and Brazil.

1. INTRODUCTION:

Since the late 20th century, infections caused by multidrug-resistant bacteria have become a significant global concern, recognized as a serious public health problem. These infections are often associated with treatment failures and the increased costs related to morbidity and mortality in patients, particularly those hospitalized. In addition to impacting the morbidity and mortality of diseases, bacterial resistance to antibiotics results in high hospital costs, as this resistance is frequently linked to the indiscriminate use of antibiotics [1].

Antimicrobial resistance is a natural biological phenomenon. As antimicrobial agents are incorporated into clinical practices, resistant strains of microorganisms are detected in laboratories[2].

Carbapenems are antimicrobial drugs used in hospitals and are often the last resort for the treatment of infections caused by multidrug-resistant microorganisms (MRM). These drugs are typically the preferred option for managing severe infections caused by enterobacteria that produce extended-spectrum β-lactamases (ESBLs) [3].

The most frequently identified groups of carbapenemases include KPC, NDM, and OXA-48, which limit the available treatment options. The rapid spread of acquired carbapenem resistance is increasingly facilitated by mobile genetic elements, such as plasmids, which transfer carbapenemase genes both within and between Gram-negative bacilli (GNB). The β-lactamase genes present in mobile genetic elements play a crucial role in the rapid global spread of antibiotic-resistant GNB [4].

Carbapenem resistance can develop due to the poor penetration of active agents, which results from genetic alterations leading to different types of porin membrane proteins and/or efflux mechanisms, in addition to the production of enzymes called carbapenemases. This latter aspect is particularly important due to the great diversity of these enzymes, especially the metallo- β -lactamases (MBLs) [5].

Carbapenemases have the ability to hydrolyze a variety of β -lactam antimicrobial agents, including carbapenems, cephalosporins, penicillins, and monobactams (such as aztreonam). These enzymes can be inhibited by clavulanic acid and tazobactam. They are classified into Ambler classes A, B, and D, with serine carbapenemases (classes A and D) and metallo- β -lactamases (class B) [6].

Metallo- β -lactamases are frequently identified in Enterobacteriaceae and Pseudomonas aeruginosa. Among MBLs, New Delhi metallo- β -lactamases (NDM), Verona integron-encoded metallo- β -lactamases (VIM), and imipenem-hydrolyzing metallo- β -lactamases (IMP) are among the most common worldwide [7].

Comment [ДС2]: A multidrug resistant organism (MDRO)?

Thus, considering the importance of multidrug-resistant bacteria in the healthcare field, this article aims to provide an integrative review of the literature addressing the frequency of carbapenemases in Brazil, with the objective of offering an updated perspective on the serious clinical and epidemiological issue of the spread of microorganisms producing these enzymes, highlighting the healthcare institutions where they have been isolated in Brazil.

2. METHODOLOGY

This is an integrative literature review based on scientific articles to analyze the resistance profile of carbapenemase-producing microorganisms in Brazil, during the period from 2020 to 2024.

The methodology applied in this study is based on synthesizing results from various sources regarding the proposed topic, aiming to gather, update, and synthesize the available scientific evidence on the subject, thereby contributing to the deepening of knowledge on the investigated theme.

The guiding question of this work was formulated by incorporating the identification of keywords to facilitate the search and guidance of the research. Thus, it was defined as: "What is the resistance profile of carbapenemase-producing Gram-negative bacilli in Brazil?"

The search was conducted in the MEDLINE (PubMed) and regional portal of the Virtual Health Library (BVS) databases, using scientific articles published between 2020 and 2024, and applying the following keywords: Carbapenemases, Carbapenem-Resistant Enterobacteriaceae, Resistant to Carbapenems, Gram-Negative Bacilli, and Brazil. Boolean operators AND and OR were used to refine the keyword search.

Those descriptors and keywords were entered in both Portuguese and English, and the selected scientific articles were in the respective languages. The established filter period was set to include studies published between 2020 and 2024 for the evaluation of the epidemiological situation of carbapenemases in Brazil. Literature reviews and other studies that did not meet the specified criteria were excluded, as well as works that did not describe the analysis of carbapenem genes resistance (Figure 1).

Data collected from the selected articles were summarized in Table 1, which was organized with the following items: article title, publication year, biological examined material, isolated microorganisms, resistance profile, and genes resistance. An Excel spreadsheet was developed for the storage and systematization of the data.

3. RESULTS

Comment [AC3]: I would add information about where this bactéria circulate – in hospitals or other settings

3.1. SELECTED ARTICLES

A total of 776 (PubMed) and 272 (BVS) articles were found as a result of the search, based on the descriptors used in the searched databases. After applying the inclusion and exclusion criteria, 75 articles remained. After duplicates were removed and the articles that did not align with the proposed topic were excluded, 26 articles were selected for qualitative analysis (Fig. 1).

The selected studies were published between 2020 and 2024 and analyzed hospital infections caused by carbapenemase-producing microorganisms in Brazil.

Database and number of articles identified between 2020 and 2024:

- Pub med 776
 - BVS 272



Articles included after reading the title:

- Pub med 48
- BVS 27



Duplicate articles removed - 49



Articles included in this literature review - 26

Image 1 - Flowchart of the methodological steps for article selection for inclusion in the review.

Source: Authors (2024).

Comment [AC4]: 75 articles were retained for the study after applying the inclusion and exclusion criteria. And then out of these 75, 26 articles remained based on the removal of duplicates and exclusion of articles not corresponding to the proposed topic. 49 articles were removed just because they were duplicates? Or were other inclusion/exclusion criteria applied? Fig 1 only mentions duplicates in this regard.

Comment [ДС5]: Presumably the fact that you studied hospital strains should be displayed in the title of the article?

Comment [ДС6]: Fig?

Table 1: Highlighted results by study for investigation of carbapenemases dissemination in Brazil (2020-2024)

Table 1: Highlighted results by study for investigation of carbapenemases dissemination in Brazil (2020-2024)					Comment [ДС7]: Many articles provide
AUTHOR, YEAR OF PUBLICATION AND STATE	EXAMINED MATERIAL	MICROORGANISMS	RESISTANCE PROFILE	GENES RESISTANCE	data not only on isolated genes, but also on sequence type bacteria (ST). Including these data in the table, figures, and the article as a whole would greatly enhance your publication.
Flores <i>et al.</i> , 2020; Rio de Janeiro[9]	Rectal swabs	Klebsiella pneumoniae	All 11 isolates were resistant to β-lactams, 90% to ciprofloxacin, 82% to tigecycline, 73% to gentamicin and 18% to amikacin.	All presented <i>bla</i> _{NDM} , 27% showed cooccurrence with the <i>bla</i> _{KPC,OXA-48} and _{VIN} genes, 46% with <i>bla</i> _{KPC} and <i>bla</i> _{VIM} and 18 with <i>bla</i> _{VIM} .	very informative as it would show not only the presence of carbapenemases but also phenotypic resistance/sensitivity. Unfortunately, some studies do not
Urzedo <i>et</i> al.,2020;Uberlândia[10]	Unspecified	Pseudomonas aeruginosa	ciprofloxacin (100%), gentamicin (100%) and cefepime (87.5%).	52.6% isolates presented <i>bla_{VIM}</i> and <i>bla_{IM}</i>	provide these data or do not give a M complete picture. I would recommend to clarify these data again to improve the quality of the article.
Cury et al., 2020; São Paulo	Rectal swabs	Unspecified	Unspecified	8.4% presented the <i>bla</i> - _{KPC} gene, 0.5%	Comment [ДС9]: Check it please
[11]				bla _{VIM} , and 0.2% bla _{NDM} .	Comment [ДС10]: Check it please
Firmo <i>et al.</i> ,2020; Pernambuco [17]	Urine (34.3%), tracheal aspirate (20.0%), blood (1%), cerebrospinal fluid (8.6%), surgical wound (5.7%), rectal swab (5.7%), bone (2.9%), lesion swab (2.9%), abscess (2.9%), sacral ulcer (2.9%) and	Klebsiella pneumoniae (45,7%), Proteus mirabilis (28,6%) and Serratia marcescens(25,7%)	All isolates were resistant to one or more carbapenems. 17.1% of isolates were resistant to amikacin, gentamicin and tobramycin.	25.7% presented <i>bla</i> _{NDM-1} , 88.6% <i>bla</i> - _{KPC-} 14.3% presented these two genes concomitantly.	2,

AUTHOR, YEAR OF PUBLICATION AND STATE	EXAMINED MATERIAL	MICROORGANISMS	RESISTANCE PROFILE	GENES RESISTANCE	_
Vivas <i>et al</i> .,2020; Sergipe [20]	Unspecified	Klebsiella pneumoniae	1.3% of the <i>bla</i> _{NDM} -positive isolates were susceptible to imipenem and meropenem. 14.3% of the <i>bla</i> _{KPC} -positive isolates were susceptible to imipenem and meropenem. 97.4% of the <i>bla</i> _{NDM} -positive isolates were considered susceptible to polymyxin.	One or more of the genes analyzed were detected in 56.5% of the isolates, with 50.3% harboring <i>bla_{NDM}</i> , 5.4% <i>bla_{KPC}</i> , and 1.2% <i>bla_{NDM}</i> and <i>bla_{KPC}</i> .	
Santos et al.,2020; Pará [21]	Blood (17,2%) and urine	Klebsiella pneumoniae	12.5% were resistant to	10.9% of the isolates tested positive for	Comment [ДС11]:
	(15,6%)		carbapenems and 81.2% were multidrug-resistant (MDR).	bla _{KPC-1} .	
Soares et al.,	Unspecified		Sensitive to amikacin,	This was the first report of the bla _{NDM} gene	
2020;Pernambuco[16]		Klebsiella aerogenes	gentamicin, tobramycin, tigecycline, colistin and polymyxin B, and susceptible, with increasing exposure (I), to ciprofloxacin, according to BrCAST.	in clinical isolates of K. aerogenes in Braz	Comment [ДС12]: K. aerogenes
Freitas et al., 2020. Rio	Unspecified	Acinetobacter baumannii	95.4% and 77.3% of the isolates		Comment [ДС13]: Check it please
Grande do Sul [24]			were resistant to meropenem and imipenem, respectively	isolates.	
Junior <i>et al.</i> , 2021.Pernambuco [18]	Unspecified	Pseudomonas aeruginosa	Unspecified	32.4% presented bla _{KPC} , 38.2% bla- _{VIM} , a	
				one isolate presented both genes concomitantly.	

AUTHOR, YEAR OF PUBLICATION AND STATE	EXAMINED MATERIAL	MICROORGANISMS	RESISTANCE PROFILE	GENES RESISTANCE	_
Wink <i>et al.</i> , 2021. Porto Alegre[26]	Unspecified	Unspecified	Unspecified	85.8% presented the <i>bla</i> _{KPC-like} gene and o	Comment [ДС15]: Check it please
Stallbaum <i>et al.</i> , 2021. Rio Grande do Sul[23]	Unspecified	Klebsiella pneumoniae	83.3% of isolates were resistant to imipenem and meropenem.	All isolates evaluated in this study presented the <i>bla</i> _{KPC} gene.	_
Higashino <i>et al.</i> , 2021. São Paulo [12]	Rectal swabs	Klebsiella pneumoniae	Unspecified	90% presented the <i>bla</i> _{KPC} gene, of which 3 presented two genes concomitantly <i>bla</i> _{KPC} and <i>bla</i> - _{NDM} .	_
Souza et al., 2021. Mato Grosso do Sul[28]	Unspecified	Pseudomonas aeruginosa	100% of <i>P. aeruginosa</i> resistant to carbapenems	11% presented the <i>bla</i> _{KPC-2} and <i>bla</i> _{TEM} genes, and 7% the <i>bla</i> _{SHV} gene.	_
Gollino <i>et al.</i> , 2021. Rio Grande do Sul[25]	Tracheal aspirate	Acinetobacter baumannii	100% susceptibility to polymyxin B	94.4% of the isolates presented the <i>bla</i> _{OXA-2} gene.	3
Ribeiro <i>et al.</i> , 2021.Pará [22]	Tracheal aspirate (50%), blood (16.7%), postoperative wound swabs (16.7%), catheter tips (11.1%) and inguinal swabs (5.6%).	Acinetobacter baumannii	Isolates were resistant to almost all antimicrobials tested and remained susceptible to tigecycline and polymyxin B.	All isolate characterized as carbapenemase producers harbored the <i>bla</i> _{OXA-23} and <i>bla</i> _{OXA-51} genes.	
Neto <i>et al.</i> , 2022. (MA),(PI), (SE); (GO),(RJ), (MG), (ES), (RS) [8]	Unspecified	Klebsiella pneumoniae	29.5% resistant to colistin	77% presented the <i>bla</i> _{KPC} gene. (Comment [ДС16]: Check it please
Freire <i>et al.</i> , 2022. São Paulo [13]	Swabs retais	Klebsiella pneumoniae	Não Especificado		Comment [ДС17]: ? Comment [ДС18]: Check it please

AUTHOR, YEAR OF PUBLICATION AND STATE	EXAMINED MATERIAL	MICROORGANISMS	RESISTANCE PROFILE	GENES RESISTANCE	_
Rodrigues <i>et al.</i> , 2022; Mato Grosso do Sul [27]	Unspecified	Klebsiella pneumoniae	Unspecified	88.9% presented the $bla_{\rm KPC}$ gene, 73.5% $bla_{\rm SHV}$, 72.2% $bla_{\rm TEM}$ and 43.9% $bla_{\rm CTX-M}$.	Comment [ДС19]: Check it please
Altafini <i>et al.</i> , 2022; Paraná [31]	Blood, urine, and tracheal aspirate	Klebsiella pneumoniae	Unspecified	75% of the isolates presented the <i>bla</i> _{KPC} gene, 19.2% <i>bla</i> _{NDM} and 5.7% presented two genes concomitantly <i>bla</i> _{KPC} and <i>bla</i> _{NDM}	
Rocha <i>et al.</i> , 2022. Bahia [32]	Rectal swabs	Klebsiella pneumoniae	100% de resistência à colistina	94.7% presented the <i>bla</i> _{KPC} gene, 16.0% <i>bla</i> _{NDM} and 1.7% <i>bla</i> _{GES} .	
Arend <i>et al.</i> , 2023. Paraná[29]	Urine (48%), blood (14%), tracheal aspirate (16%), body fluids (4%), and other clinical specimens (18%).	Klebsiella pneumoniae (68,5%), Escherichia coli (10,1%), Enterobacter cloacae complexo(7,6%), Acinetobacter baumannii (2,2%) and Pseudomonas aeruginosa (0,6%).	All isolates were resistant to imipenem, 92% to amikacin, 72% were sensitive to colistin and 40% to gentamicin.	4,55% presented the <i>bla_{NDM}</i> gene.	
Barroso <i>et al.</i> , 2023. São Paulo[14]	Swab (65.4%) blood (11.5%), urine (11.6%), tracheal aspirate (3.8%), pleural fluid (3.8%), and surgical wound (3.8%)	Klebsiella pneumoniae	Unspecified	72.2% carried <i>bla</i> _{KPC} , the only carbapenemase-coding gene found.	
Rocha <i>et al.</i> , 2024. Pernambuco [15]	Unspecified	Acinetobacter baumannii	Widespread resistance to multiple antimicrobials was evident	genes were detected. Furthermore, this is the first report of blavim-2 genes in A. baumannii isolates carrying the bla _{OXA-23-1}	Comment [JC21]: It is better to clarify specific data Comment [JC22]: Check it, please. "A total of 163 different bacterial
Santos, <i>et al.</i> , 2024. Rondônia [33]	Unspecified	Acinetobacter baumannii, Klebsiella pneumoniae, Pseudomonas aeruginosa Enterobacter cloacae	Unspecified	A. baumannii isolates bla _{OXA-23-like} 75.2% bla _{OXA-58-like} 19% and bla _{OXA-143-like} 4.1%. I K. pneumoniae isolates, the bla _{KPC} gene encoded the majority of carbapenemase 95.7%.	species were identified over 4 years of research, among which 110/163 (67.48%) had AST profiles for ertapenem, meropenem and imipenem (Table 1). Among them, 72/110 (65.45%) species showed resistance to carbapenems (based on phenotypic tests)."

AUTHOR, YEAR OF PUBLICATION AND STATE	EXAMINED MATERIAL	MICROORGANISMS	RESISTANCE PROFILE	GENES RESISTANCE	
Krul <i>et al.</i> , 2024. Paraná [30]	Unspecified	Klebsiella pneumoniae	Unspecified	The most frequent resistance gene	was Comment [ДС23]:
				bla _{KPC}	Comment [ДС24]: Check it, please. "Twenty-five CRKP isolates were identified, and
Martins <i>et al.</i> , 2024. Pernambuco[19]	Unspecified	Klebsiella pneumoniae	Unspecified	46.66% had <i>bla_{NDM},</i> 35.55% <i>bla_{KPC}</i> 17.79% had both genes concomita	antimicrobial susceptibility was assessed using broth microdilution. Carbapenemase production and β -lactamase genes were detected by phenotypic and genotypic tests"
		POLICE			

4. Qualitative analysis of studies

A study conducted by the Hospital Infection Research Laboratory (LAPIH - FIOCRUZ) covered four regions of Brazil, including eight states: Northeast; Central-West; Southeast and South. A total of 502 clinical isolates of *Klebsiella pneumoniae* were analyzed, collected from samples of blood, urine, tracheal aspirate, rectal swab, catheter tips, sputum, tissue fragments and wounds. The investigation included resistance to polymyxins, genetic factors related to clonality, in addition to the search for genes associated with resistance to carbapenems and an assessment of antimicrobial resistance. The results of the study showed a colistin resistance rate of 29.5% in the *K. pneumoniae* samples. The *bla*_{KPC} gene was found in 77% of the samples, indicating high resistance to polymyxins due to their use as a treatment. Recently, the WHO classified polymyxins as "critical" antimicrobials, since they are among the few options available for the treatment of severe bacterial infections. On the other hand, the increase in resistance to carbapenems in the last decade has favored the spread of KPC, also resulting in an increase in resistance to polymyxins [8].

Rio de Janeiro: In a study on the surveillance culture of patients hospitalized in an intensive care unit of a hospital in Rio de Janeiro, the genetic relationship of carbapenem-resistant *Klebsiella pneumoniae* carrying the bla_{NDM} gene was analyzed. The study identified that 27% of the strains showed correlation with KPC, OXA-48, and VIM, 46% with KPC and VIM, and 18% with the VIM type. None of the strains analyzed presented the resistance genes bla_{KPC} , bla_{SPM} , or bla_{IMP} . All 11 isolates demonstrated resistance to β -lactams and were classified as multidrug-resistant. The coexistence of *K. pneumoniae* producing NDM with other carbapenemases has been frequently reported in various regions worldwide. Furthermore, for the first time, it was observed that *K. pneumoniae* can harbor up to four types of carbapenemases in active surveillance cultures[9].

Minas Gerais: A study conducted in a tertiary university hospital in the city of Uberlândia, MG, in 2020, analyzed the detection of resistance genes in 138 carbapenem-resistant *Pseudomonas aeruginosa* isolates, identified as bla_{SPM} , bla_{VIM} , and bla_{IMP} , with high resistance frequencies to ciprofloxacin (100%), gentamicin (100%), and cefepime (87.5%). These results highlight a significant shift in the epidemiology of carbapenem-resistant *P. aeruginosa* isolates in a Brazilian hospital, with low detection of SPM and VIM, suggesting the involvement and coexistence of other resistance mechanisms in these strains and encouraging a reevaluation of empirical antibiotic use guidelines in institutions[10].

São Paulo: The detection of the *bla_{KPC}* gene in *Klebsiella pneumoniae* in three surveillance studies using rectal swab samples from various hospitals in the city of São Paulo indicates the need for the adoption of infection control strategies to minimize costs and harm to patients, particularly those undergoing transplantation or in critical health conditions[11],[12],[13]. In another study conducted in a pediatric hospital in São Paulo, where samples from different

Comment [AC25]: In the table when referring to this article, the EXAMINED MATERIAL section indicates "Unspecified".

Comment [AC26]: The first time a microorganism is mentioned, the species and genus name should be written complitley (ex. Klebsiella pneumoniae). Then we can write the abbreviated name throughout the text (ex. K. pneumoniae). In this manuscript there is an alternation of full names and abbreviations. I recommend a uniform format.

Comment [ДC27]: Other carbapenemases are described in this publication, which could be included.

biological sites were analyzed, it was found that 72.2% of carbapenem-resistant K. pneumoniae isolates also carried the bla_{KPC} gene. These results highlight that the strains found in the pediatric population are similar to those detected in adults, underscoring the importance of epidemiological surveillance for the effective implementation of preventive and control measures[14].

Pernambuco: In a reference hospital located in Recife, the first case of bla_{VIM-2} in *Acinetobacter baumannii* isolates carrying the $bla_{OXA-23-like}$ gene and the $bla_{OXA-143}$ gene was recorded in Brazil. Among the 78 *A. baumannii* isolates analyzed, widespread resistance to various antimicrobials was observed[15]. Additionally, the first case of the bla_{NDM} gene in clinical isolates of *Klebsiella aerogenes* in Brazil was documented in 2020 in the state of Pernambuco[16]. A study conducted in three hospitals in Recife also reported the first case of the association between bla_{NDM-1} and bla_{KPC-2} in *Proteus mirabilis* and *Serratia marcescens* in Brazil[17].

Another study conducted in Pernambuco highlighted the analysis of clinical P. aeruginosa isolates, where the bla_{KPC} and bla_{VIM-2} genes were detected in 32.4% and 38.2% of the samples, respectively, with one of the isolates carrying both the bla_{KPC} and bla_{VIM-2} genes[18]. The reports of antimicrobial resistance, with the detection of the bla_{VIM-2} , bla_{NDM} , and bla_{KPC} genes in bacterial isolates in Brazil, particularly in Recife, indicate a concerning scenario of bacterial resistance. These findings underscore the urgent need for monitoring and controlling hospital infections, as well as the importance of effective antimicrobial stewardship strategies to contain the spread of these resistant strains.

A study conducted in Pernambuco, involving patients with and without COVID-19, showed that one of the main species of microorganisms identified was *Klebsiella pneumoniae*, with the detection of the resistance genes bla_{NDM} (46.66%), bla_{KPC} (35.55%), and both concurrently (17.79%). The presence of *Enterobacterales* carrying bla_{KPC} and bla_{NDM} , particularly *K. pneumoniae*, in infections and colonizations of patients with and without COVID-19, highlights the genetic diversity and carbapenem resistance observed in various species of this order[19].

Sergipe: A study conducted in a public hospital in Aracaju revealed that the majority of *Klebsiella pneumoniae* isolates (50.3%) were positive for bla_{NDM} , while 1.2% were positive for both bla_{NDM} and bla_{KPC} . Among the isolates that tested positive for bla_{NDM} , 97.4% were considered sensitive to polymyxin, and only 1.3% showed sensitivity to imipenem and meropenem. The presence of NDM has been associated with multidrug resistance and has been reported in various Brazilian states, affecting different species of Gram-negative bacteria. These findings highlight the rapid spread of NDM-positive isolates and underscore the urgent need for alternative therapies to treat infections caused by these multidrug-resistant isolates. The KPC and NDM enzymes are rarely found in the same strain, and the concurrent production of these carbapenemases in a single strain can lead to significant resistance to carbapenems[20].

Belém: A study conducted to investigate the clinical, epidemiological, and resistance-related aspects of infections caused by Klebsiella pneumoniae in cancer patients treated at a reference oncology center in the state of Pará identified isolates that tested positive for the blaker gene, with 12.5% being resistant to carbapenems and 81.2% being multidrug-resistant (MDR). A high prevalence of MDR K. pneumoniae was observed in this study population, along with a significant occurrence of carbapenem resistance, emergence of colistin resistance, and the detection of the blakec gene. The data obtained emphasize the need for strategies addressing epidemiological surveillance, combining continuous work between hospital infection control committees and healthcare professionals[21]. Another study conducted in a public institution in the southeastern region of the state of Pará, with kidney transplant services and renal replacement therapy, analyzed Acinetobacter baumannii isolates resistant to imipenem and meropenem found in different biological sites (tracheal aspirates, blood, catheter bridge), with bla_{OXA-23} and bla_{OXA-51} genes detected in all these microorganisms. Bacterial resistance is an emerging problem that demands the utmost attention and effort for mitigation. The importance of screening colonized or infected patients and providing frequent training to healthcare professionals in ICUs and clinics is emphasized. Additionally, surveillance for A. baumannii resistant to imipenem and meropenem, along with rational antimicrobial administration, must be reinforced[22].

Rio Grande do Sul: In an investigation conducted at a hospital in Pelotas, *Klebsiella pneumoniae* isolates associated with hospital infections were analyzed. The strains carried the bla_{KPC} gene, and 83.3% of them were resistant to carbapenems[23]. In another study, also conducted in Pelotas by Freitas et al.[24], which analyzed *Acinetobacter baumannii* strains, the researchers revealed that 95.4% of these isolates carried the bla_{VIM} gene, and 77.3% of them were resistant to meropenem and imipenem. The combination of data regarding the resistance of *K. pneumoniae* and *A. baumannii* strains in this city in Rio Grande do Sul, associated with the presence of bla_{VIM} and bla_{KPC} genes, suggests a similarity in the resistance trajectory to carbapenems that impacts these microorganisms. These findings should serve as a call to action for joint efforts among hospitals, public health authorities, and researchers to combat the growing threat of infections caused by multidrug-resistant bacteria in hospital environments.

Additionally, the first data on the *Acinetobacter baumannii* profile in the western border of Southern Brazil raise concerns about the presence of endemic clones producing OXA-23 in this region. This study demonstrated high resistance to aminoglycosides and fluoroquinolones, contrasting with 100% sensitivity to polymyxin B. The *bla*_{OXA-23} gene was identified in 34 strains[25].

Furthermore, a study conducted at a tertiary hospital in Southern Brazil evaluated the growth in the frequency of bla_{NDM} , highlighting a constant increase of $bla_{NDM-like}$ genes, which rose from 4.2% in 2015 to 24% in 2020. This increase in the detection frequency of bla_{NDM} raises an important issue, as the therapeutic options available for the treatment of patients infected by bla_{NDM} carrying bacteria are currently very limited[26].

Mato Grosso do Sul: A study conducted in Mato Grosso do Sul aimed to determine the molecular epidemiology of *Klebsiella pneumoniae* carrying bla_{KPC} , recovered from three public hospitals in Brazil. The *K. pneumoniae* isolates presented in the study carried the following resistance genes: bla_{KPC} (88.9%), bla_{SHV} (73.5%), bla_{TEM} (72.2%), and bla_{CTX-M} (43.9%). Additionally, new sequences of types that had not been previously identified in the country were detected. It was observed that *K. pneumoniae* belonging to the same clone was present in different hospitals within the same region, highlighting the spread of multidrug-resistant *K. pneumoniae* [27].

At a public tertiary hospital located in the municipality of Dourados, Mato Grosso do Sul, an evaluation of *Pseudomonas aeruginosa* isolates resistant to carbapenems was conducted, of which only 10.7% were *KPC*-producing strains. The production of carbapenemase has emerged in Brazil as the main mechanism of resistance to carbapenems among clinical isolates of *P. aeruginosa*. However, despite the identification of some *KPC*-producing strains, the low rates suggest that other mechanisms of carbapenem resistance may also be at play [28].

Paraná: A study conducted across multiple hospitals in eight cities in the Southern region of Brazil investigated the spread of bacteria producing NDM in clinical samples from urine, blood, tracheal aspirates, and body fluids. The number of isolates with bla_{NDM} in the southern cities of Brazil has been growing significantly. Moreover, the frequency varies considerably between municipalities, indicating that some cases correspond to small, localized outbreaks, while in others there is a more widespread dissemination. This situation poses challenges for treatment, as carbapenem antibiotics are the last β-lactams with proven efficacy against nearly all Gram-negative microorganisms study conducted in Paraná examined the molecular epidemiology and mechanisms of carbapenem resistance in K. pneumoniae isolates from pediatric patients, identifying that the most common resistance gene found was blaker. Both stight the occurrence of two different resistance genes, but in distinct populations: adults and children. In an additional study in the state of Paraná, in the city of Maringá, among COVID-19-infected patients who were carriers of K. pneumoniae producing carbapenemase, it was found that 75% of isolates carried the bla_{KPC} gene, 19.2% harbored bla_{NDM} , and 5.7% contained both genes. Given the results presented in this research, it is essential to continuously invest in training, both in the management of antimicrobials and in infection control within healthcare settings. This will ensure that, in the post-pandemic period, antimicrobial stewardship and surveillance programs remain active, thereby reducing the emergence and spread of multidrug-resistant (MDR) microorganisms.

Bahia: A study conducted at a reference hospital in Salvador aimed to investigate the genetic association, antimicrobial resistance profile, and resistance mechanisms in K. pneumoniae isolates resistant to carbapenems and colistin (CoIR-CRKP). The material analyzed consisted of rectal swab samples. The bla_{KPC} gene was identified in 94.7% of the isolates, bla_{NDM} in 16.0%, and bla_{GES} in 1.7%. The bla_{OXA-48} , bla_{VIM} , and bla_{IMP} genes were not

Comment [ДС28]: Link to the article (s)?

found. The research indicates a prolonged outbreak of CoIR-CRKP, suggesting the possibility of cross-transmission of *K. pneumoniae* isolates resistant to both carbapenems and colistin [32].

Rondônia: A study conducted in the southern Amazon region aimed to map the diversity of carbapenem-resistant bacteria, with an emphasis on molecular epidemiology and the analysis of genes responsible for carbapenemase production. Seventy-two species were detected showing resistance profiles to these antibiotics, of which 25 contained at least one gene encoding carbapenemases from classes A ($bla_{KPC-like}$), B ($bla_{NDM-like}$, $bla_{SPM-like}$, or $bla_{VIM-like}$), and D ($bla_{OXA-23-like}$, $bla_{OXA-24-like}$, $bla_{OXA-48-like}$, $bla_{OXA-58-like}$, or $bla_{OXA-143-like}$), including species such as K. pneumoniae, P. aeruginosa, A. baumannii, S. marcescens, and Providencia spp. These findings have the potential to contribute to scientific knowledge, providing molecular and epidemiological data that can be used in state-level bacterial resistance surveillance, as well as supporting the formulation of public health policies [33].

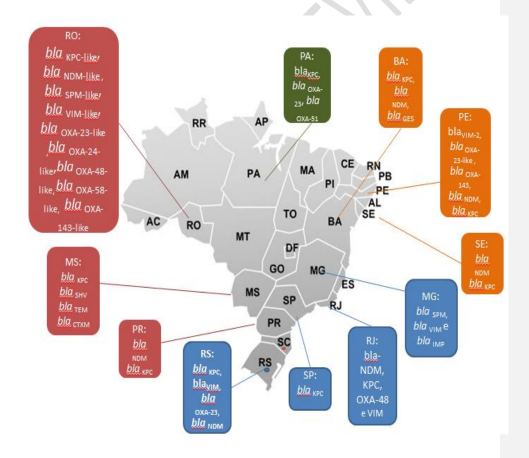


Image 2: National patterns of gene resistance dissemination.

Comment [ДС29]: Optional - Consider to add type of bactéria and ST.

5. Final considerations

The results presented in this study highlight that the state of Pernambuco in northeastern Brazil has been significant in reporting the first case of the bla_{VIM-2} gene in A. baumannii isolates, along with the identification of the $bla_{OXA-23-like}$ and $bla_{OXA-143}$ genes, marking an important advancement in microbiological surveillance. Also notable is the first report of the association between bla_{NDM-1} and bla_{KPC-2} in P. mirabilis and S. marcescens, as well as the presence of the bla_{NDM} gene in K. aerogenes in this state.

Moreover, the cohabitation of multiple resistance genes in microorganisms isolated from several Brazilian states underscores the need for effective control and monitoring strategies, as this situation can further complicate clinical management and the efficacy of antimicrobial treatments. These findings reinforce the importance of ongoing research and the development of public health policies to address the growing threat of infections caused by multidrug-resistant bacteria in the country.

In Brazil, the complexity and diversity of bacterial resistance genes pose a significant challenge to public health. Among these genes, those that express carbapenemase and KPC stand out due to their high prevalence, contributing to resistance to last-line antibiotics. This situation is concerning, as the spread of bla_{KPC} -bearing strains can hinder the treatment of severe infections, increasing morbidity and mortality. Understanding this genetic diversity is crucial for the development of effective control and prevention strategies aimed at mitigating the impact of antimicrobial resistance in the country.

6. REFERENCES:

- Silva AF, Junior OMR. Bacterial resistance dueto in discriminate use of the carbapenems meropenem and imipenem: anintegrative review.Research, Society and evelopment, 2022; 11(7): e44711730195M.
- Melo DB. Clonal pattern and antimicrobial susceptibility profile of Escherichia coli strains isolated from food and clinical specimens .Dissertation (Masters in Food Science) -Faculty of Pharmacy, Federal University of Bahia, Salvador, 2010.
- 3. Oliveira RS, Silva SS. Klebsiella pneumonia and carbapenemase (KPC) as a public health threat in Brazil: a literature review. University academic center uniftc undergraduate course in biomedicine Jequié teaching unit. Salvador, 2022.
- 4. Soares CRP, Silva RFR, Junior JBO, Araujo PSR, Firmo EF. Molecular epidemiology of multidrug-resistant gram-negative bacilli producing carbapenemase isolated from different sites of infection. Research, Society and Development. 2021; 32 (9): p. e30210918070.

Comment [ДС30]: Different spacing in the text

- Figueredo ACF, de Freitas NL, Dalmolin TV, & Brandão,: Pseudomonas aeruginosa: overview of the profile of resistance to carbapenems in Brazil. Brazilian Journal of Development, 2021; 7(1): 9661–9672. https://doi.org/10.34117/bjdv7n1-655.
- 6. Rusth MC, Meurer IR, Vicenti CM, de Lourdes Junqueira M, da Silva Lima J, Fochat, R C, Garcia PG. Prevalence and carbapenem resistance profile of Enterobacterales isolated from tracheal aspirate of patients admitted to a university hospital. Acervo Saúde Electronic Magazine, 2024; 24(1): 14830-14830.
- Bonomo RA, Burd EM, Conly J, Limbago BM, Poirel L, Segre JA, Westblade LF. Carbapenemase-producing organisms: a global scourge. Clinical infectious diseases, 2020; 66(8): 1290-1297.
- 8. Conceição-Neto OC, da Costa BS, Pontes LDS, Silveira MC, Justo-da-Silva LH, de Oliveira Santos IC, Carvalho-Assef AP. Polymyxin resistance in clinical isolates of K. pneumoniae in Brazil: Update on molecular mechanisms, clonal dissemination and relationship with KPC-producing strains. Frontiers in Cellular and Infection Microbiology, 2022; 12: 98125. https://doi.org/10.3389/fcimb.2022.898125.
- 9. Flores C, Bianco K. de Filippis I, Clementino MM, Romão CMC. Genetic relatedness of NDM-producing Klebsiella pneumoniae co-occurring VIM, KPC, and OXA-48 enzymes from surveillance cultures from an intensive care unit. Microbial drug resistance, 2020; 26 (10): 1219-1226.https://doi.org/10.1089/mdr.2019.0483
- 10. Urzedo JE, de Paula Menezes R, Porto JP, Ferreira ML, Gonçalves IR, de Brito CS, Ribas RM. High mortality by nosocomial infections caused by carbapenem-resistant P. aeruginosa in a referral hospital in Brazil: facing the perfect storm. Journal of Medical Microbiology; 2020; 69(12): 1388-1397. https://doi.org/10.1099/jmm.0.001273.
- 11. Cury AP, Junior JA., Costa SF, Salomão MC, Boszczowski Í, Duarte AJ, Rossi F. Diagnostic performance of the XpertCarba-R™ assay directly from rectal swabs for active surveillance of carbapenemase-producing organisms in the largest Brazilian University Hospital. Journal of microbiological methods, 2020; 171:105884.
- 12. Higashino H R, Marchi AP, Ruedas Martins RC, Bubach Carvalho L, Vieira Perdigão Neto L, Farrel Côrtes M, Costa SF. Carbapenem-resistant Klebsiella pneumoniae colonization and infection is associated with lower overall survival in a cohort of haematopoietic stem-cell transplantation patients: mechanism of resistance and virulence by whole-genome sequencing. Journal of Medical Microbiology,2020; 70(10): 001422. https://doi.org/10.1099/jmm.0.001422.

- 13. Freire MP, de Oliveira Garcia D, Lima SG, Pea CRD, Reusing Junior JO, Spadão F, Pierrotti LC. Performance of two methods of carbapenem-resistant Enterobacterales surveillance on a kidney transplant ward: selective culture of and real-time PCR directly from rectal swabs. *Infection*, 2022; 50(6): 1525-1533.
- 14. do Valle Barroso M, da Silva CR, Benfatti LR, Gozi KS, de Andrade LK, Andrade LN, Casella T. Characterization of KPC-2-producing Klebsiella pneumonia e and affected patients of a pediatric hospital in Brazil. Diagnostic Microbiology and Infectious Disease, 2023; 106(2): 115932. https://doi.org/10.1016/j.diagmicrobio.2023.115932.
- 15. Rocha IV, Martins LR, Pimentel MIS, Mendes RPG, Lopes, ACDS. Genetic profile and characterization of antimicrobial resistance in Acinetobacter baumannii post-COVID-19 pandemic: a study in a tertiary hospital in Recife, Brazil. Journal of Applied Microbiology, 2024;135(6). https://doi.org/10.1093/jambio/lxae148
- 16. Soares CRP, Oliveira-Júnior JB, Firmo EF. Firstre port of a Bla NDM-resistant gene in a Klebsiella aerogenes clinical isolate from Brazil. Revista da Sociedade Brasileira de Medicina Tropical, 2020; 54: e02622020. https://doi.org/10.1590/0037-8682-0262-2020
- 17. Firme EF, Beltrão EMB, da Silva FRF, Alves LC, Brayner FA, Veras D ,Lopes ACS. Association of blaNDM-1 with blaKPC-2 and aminoglycoside modify ingenzyme genes among Klebsiella pneumoniae, Proteus mirabilis and Serratia marcescens clinical isolates in Brazil. Journal of Global Antimicrobial Resistance, 2020; 21: 255-261.
- 18. Costa-Júnior SD, da Silva AMCM, Niedja da Paz Pereira J, da Costa Lima JL, Cavalcanti IMF, Maciel MAV. Emergence of rmtD1 gene in clinical isolates of Pseudomonas aeruginosa carrying bla KPC and/orbla VIM-2 genes in Brazil. Brazilian Journal of Microbiology, 2021: 52(4): 1959-1965.
- 19. Martins LR, Pimentel MIS, de Oliveira ÉM, Jucá MB, Beltrão EMB, Lopes ACDS. Occurrence of bla NDM-1, bla NDM-5, bla NDM-7, and bla KPC-2 genes in clinical isolates of enterobacterales with high genetic variability, from colonization and infection in patients with or without COVID-19, from a hospital in Brazil. Journal of Applied Microbiology,2024; 135(8): Ixae212. https://doi.org/10.1093/jambio/lxae212.
- 20. Vivas R, Dolabella SS, Barbosa AAT, S. Prevalence of Klebsiella pneumonia carbapenemase-and New Delhimetallo-beta-lactamase-positive K. pneumoniae in Sergipe, Brazil, and combination therapy as a potential treatment option. Revista da

- Sociedade Brasileira de Medicina Tropical, 2020; 53: e20200064.https://doi.org/10.1590/0037-8682-0064-2020.
- 21. Santos ALSD, Rodrigues YC, Melo MVHD, Silva dos Santos PA, Oliveira TNDC, Sardinha DM,Lima KVB. First Insights into Clinical and Resistance Features of Infections by Klebsiella pneumoniae among Oncological Patients from a Referral Center in Amazon Region, Brazil. Infectious Disease Reports, 2020; 12(3): 110-120.
- 22. Ribeiro EA, Gales AC, Oliveira APSD, Coelho DD, Oliveira RAD, Pfrimer IAH, Carmo Filho JRD. Molecular epidemiology and drugresistance of Acinetobacter baumanni iisolated from a regional hospital in the Brazilian Amazon region. Revista da Sociedade Brasileira de Medicina Tropical, 2020; 54: e20200087.
- 23. Stallbaum LR, Pruski BB, Amaral SC, de Freitas SB, Wozeak DR, Hartwig DD. Phenotypic and molecular evaluation of biofilm formation in Klebsiella pneumonia carbapenemase (KPC) isolates obtained from a hospital of Pelotas, RS, Brazil. Journal of Medical Microbiology,2021: 70(11): 001451.
- **24.** Freitas SB. Molecular Characterization of Carbapenem-Resistant Acinetobacter baumannii Associated with Nosocomial Infection in the Pelotas, RS, Brazil. CurrMicrobiol., 2020; 77: 2724-2734.
- **25.** Paula Gollino G, Escobar BM, da Silveira ID, Siqueira RHR, Ferreira JC, da Costa Darini AL, Ribeiro VB. Molecular epidemiology of carbapenem-resistant Acinetobacter baumannii from Southern Brazil. Revista de Epidemiologia e Controle de Infecção, 2021; 11(1): 26-31.
- **26.** Wink PL, Martins AS, Volpato F, Zavascki AP, Barth AL. Increased frequency of bla NDM in a tertiary care hospital in southern Brazil. Brazilian Journal of Microbiology, 2021; *52*: 299-301.
- 27. Rodrigues ACS, Chang MR, Santos ICDO. Carvalho-Assef APD. A. Molecular Epidemiology of bla KPC-Encoding Klebsiella pneumoniae Isolated from Public Hospitals in Midwest of Brazil. Microbial Drug Resistance, 2022; 28(1): 1-6.
- **28.** Souza GHDAD, Rossato L, Brito GT, Bet GMDS, Simionatto S. Carbapenem-resistant Pseudomonas aeruginosa strains: a worrying health problem in intensive care units. Revista do Instituto de Medicina Tropical de São Paulo, 2021; *63*: e71.

- **29.** Arend LN, Bergamo R, Rocha FB, Bail L., Ito C, Baura VA, Tuon FF. Dissemination of NDM-producing bacteria in Southern Brazil. Diagnostic Microbiology and Infectious Disease,2023; *106*(2): 115930.
- **30.** Krul D, Rodrigues LS, Siqueira AC, Mesa D, Dos Santos ÉM, Vasconcelos TM, Dalla-Costa L. M. High-risk clones of carbapenem resistant Klebsiella pneumoniae recovered from pediatric patients in Southern Brazil. Brazilian Journal of Microbiology,2024; 1-7.
- 31. Dambroso-Altafini D, dos Santos Saalfeld SM, de Mattos MDSF, Martinez HV, Shinohara DR, Zarpellon MN, Tognim MCB. Overuse of empirical antibiotics in a COVID-19 intensive care unit led to the spread of carbapenem-resistant Gram-negative bacteria in a teaching hospital. Journal of Global Antimicrobial Resistance, 2022; 30; 100.
- **32.** Rocha VFD, Barbosa MS, Leal HF, Silva GEO, Sales NMMD, Monteiro ADSS, Reis JN. Prolonged outbreak of carbapenem and colistin-resistant Klebsiella pneumoniae at a large tertiary hospital in Brazil. Frontiers in microbiology, 2022;13: 831770.
- 33. Dos Santos LA, Cayô R, Valiatti TB, Gales AC, de Araújo LFB, Rodrigues FM, Campanharo M. Biodiversity of carbapenem-resistant bacteria in clinical samples from the Southwest Amazonregion (Rondônia/Brazil). Scientific Reports, 2024; 14(1): 9383.