**Antimicrobial Resistance in *Klebsiella pneumoniae* Strains: Mechanisms and Outbreaks †**

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**Abstract:** The enterobacteria that produce β-lactamases are the main focus of infections in the healthcare environment. This is due to the difficulty they present in terms of treatment, their ease of transmission, and the impact they represent at the economic and personal level. The bacteria of greatest clinical relevance are those with resistance to third and fourth generation cephalosporins, extended spectrum β-lactamase and AmpC. Currently, carbapenemics are one of the few antimicrobials effective against multi-drug resistant organisms. However, the emergence of carbapenem-resistant enterobacteria has increased health concerns. These microorganisms include *K. pneumoniae*, a pan-resistant bacteria with high morbidity and mortality rates in public health facilities. In this work, we have carried out a review on the antimicrobial resistance genes found in its genome, as well as the resistance mechanisms involved. Finally, we will focus on the main outbreaks causing nosocomial infections during the last few years.

**Keywords:** multi-drug resistant organisms; extended spectrum β-lactamase; cephalosporins; carbapenem-resistant enterobacteria

1. Introduction

*Klebsiella pneumoniae* is an encapsulated Gram-negative bacillus belonging to the Enterobacteriaceae family. It was first described in 1882 by Carl Friedlander, from lung tissues of dead pneumonia patients. It presents a mucoid phenotype in media such as MacConkey or eosin methylene blue agars. Moreover, it is capable of fermenting lactose and mannitol, and reducing nitrate to nitrite. *K. pneumoniae* is a bacterium able to infect humans, causing different types of infections, including respiratory and urinary tract infections, soft tissue infections, surgical wounds and sepsis. After *E. coli*, *K. pneumoniae* is the second most important opportunistic enterobacterium causing nosocomial and community infections. However, it is a ubiquitous species in nature, although environmental strains are more susceptible to antibiotics than clinical ones. It is a consequence of the selection pressure exerted by antibiotics in health care settings [1].

The pangenome of *K. pneumoniae* (total set of genes) is approximately 5.5 Mbp in size and codes for an average of 5500 genes. More than 30,000 unique sequences have been identified that code for different proteins. However, it is still an open pangenome, in which there are accessory genes that have not yet been characterized [1,2]. Its core genome (set of genes shared by all members of the species) is composed of approximately 2000 genes. These genes have been determined by examining their presence in more than 95% of the isolates from that species [1,2]. The accessory genome contains genes that vary between bacteria of the same species. This includes both chromosomally encoded genes and those located in the mobile genetic elements. In *K. pneumoniae*, most of the genome is made up of accessory genes. This allows their classification into different strains, and different pathotypes.
can be catalogued in the clinical setting [opportunistic, Carbapenem-Resistant Enterobacteria (CRE) or hypervirulent (hv) bacteria] (Figure 1) [1].

**Pool of Accessory Genes**

| Aerobactin siderophore (Aer) | Nitrogen fixation operon (nif) | Regulator of mucoid phenotype A (rmpA) | Salmochelin siderophore (Sal) | Versiniabactin siderophore (Ybt) |
|-----------------------------|-----------------------------|-------------------------------------|-------------------------------|----------------------------------|
| Carbapenemase (bla_KPC)     | ESBL                        | Regulator of mucoid phenotype A2 (rmpA2) |                               |                                  |

![Figure 1](image)

**Figure 1.** *K. pneumoniae*, *K. variicola* and *K. quasipneumoniae* are three species that share a set of accessory genes. Ent (Enterobactin, blue) and Fimbriae (dark green) represent conserved genes. The accessory genes are shown as examples, they are not a definitive list. The evolutionary tree is not drawn to scale [1].

These accessory genes can also be combined to form new pathotypes (hvCRE) as they can be shared between species by THG [1,2]. Therefore, the accessory genome reveals the virulence of a bacterium and its study is essential to understand the clinical features of an infectious disease. Moreover, this type of genome can evolve due to the changes in the genome of the pathogen, or by the acquisition of new genes for resistance or variation in its expression. Table 1 shows *K. pneumoniae* genes that determine the development of antimicrobial resistance with their respective enzymes or target molecules [3]. The high number of genes encoding for β-lactamases supports that they are crucial to the antibiotic resistance.

In recent decades, *K. pneumoniae* has acquired a wide variety of antibiotic resistance mechanisms and, thus, the nosocomial infections it can cause are particularly problematic. This increases the complexity of detecting resistant strains and assigning the appropriate treatment for each bacterium. In a healthcare center, there are several potential foci that facilitate the transmission of *K. pneumoniae* such as: person-to-person contact between health workers and patients, contaminated surfaces, instrumentation, and some medical techniques (endoscopy). Diseases that weaken the patient, such as cancer, diabetes mellitus, some neurological disorders, or fluid and electrolyte disorders may be classified as risk factors for infection. Despite *K. pneumoniae* being part of the normal microbiota of the intestine, it is able to colonize other mucosal surfaces, including the nasopharyngeal, urinary, and gastrointestinal tracts, leading to serious infectious diseases in immunocompromised patients [1].
Table 1. Antimicrobial resistance genes in the genome of *K. pneumoniae* [1].

| **β-Lactamases** | **bla genes conferring resistance (* intrinsic*) | **Mutations** |
|------------------|-----------------------------------------------|---------------|
| Class A          | CARB-3, PSE-1, SCO-1, SHV-1 *, TEM-1         |               |
| - ESBL          | CTX-M, SHV-5, TEM-10, VEB                |               |
| - Carbapenemase  | KPC, GES-5 Class B Metallo              |               |
| - β-lactamase    | CphA, IMP, NDM, SIM, VIM Class C        |               |
| (Cephalosporinase) | AmpC, CMY, DHA, FOX, MIR                  |               |
| Class D          | OXA-1, OXA-2, OXA-7, OXA-9, OXA-10, OXA-12 |               |
| - ESBL          | OXA-11, OXA-15                              |               |
| Carbapenemase    | OXA-48, OXA-51, OXA-181, OXA-237          |               |

| **Other AMR** | **Genes conferring resistance (* intrinsic*) | **Mutations** |
|---------------|---------------------------------------------|---------------|
| Aminoglycosides | aac, aadA, aadB, aph, armA, rmt, strAB (see carbapenemase bla genes, class A and D above) | Mutations in ompK35, ompK36 |
| Colistin      | mcr-1, mcr1.2                                | Inactivation of pmrB, mcrB; mutations in crrB Fluoroquinolones |
| OqxAB efflux Macrolides | ereA, ereB, ermB, mef, mph, msrE | - |
| Phenicols     | catA, catB, cml, floR                        | Rifampin |
| Sulfonamides  | folP, sul1, sul2, sul3                        | - |
| Tetracycline  | tet genes                                    | Uregulation of acrAB or OqxAB efflux; mutation in rpsJ |
| Tigecycline   | -                                           | - |
| Trimethoprim  | dfr genes                                    | - |

2. Main Resistance Mechanisms in *K. pneumoniae*

*K. pneumoniae* has experienced an increase in resistance to different families of antibiotics. Most of these antibiotic resistance mechanisms (AMR) have been acquired through horizontal gene transfer (HGT) conferring high level resistance to antibiotics of the type β-lactams and quinolones.

(a) Mechanisms of antibiotic resistance to β-lactam antibiotics

One of the resistance mechanisms in *K. pneumoniae* consists in the alteration of penicillin binding proteins (PBPs), enzymes that catalyze peptidoglycan synthesis and specific target of the β-lactam antibiotics. Modifications in their structure decrease the affinity of PBPs for the β-lactam antibiotics, increasing resistance to them [4,5]. Moreover, outer membrane (OM) permeability modifications are critical in *K. pneumoniae* resistance (hydrophilic drugs). OM is a barrier that must be crossed before it can bind to PBPs [5]. The antibiotic molecules must use porins (OmpK35 and OmpK36), which are reduced in number or modified, decreasing the permeability of the membrane [4,6]. Another common mechanism in *K. pneumoniae* strains are active ejection pumps like AcrAB and OqxAB pumps [4].

(b) Quinolone resistance

Mechanisms of fluoroquinolone resistance are due to point mutations in specific areas of DNA gyrase (genes gyrA and gyrB) and topoisomerase IV (genes parC and parE), called quinolone resistance determining region. The first target for quinolone resistance in Gram-negative microorganisms is gyrA and then parC. Several simultaneous mutations in these genes increase high level resistance [7]. In addition, resistance mechanisms acquired by mobile genetic elements, such as the acquisition of Qnr (quinolone resistance) proteins that protect DNA gyrase and topoisomerase IV. Altered permeability and the presence of efflux pumps are also mechanisms involved in quinolone resistance [8,9].

3. Outbreaks Caused by Resistant Strains of *K. pneumoniae*

More than 100 different AMR acquired genes have been identified in *K. pneumoniae*. The presence of SHV-1 penicillinase on the chromosome confers intrinsic resistance to ampicillin, amoxicillin and ticarcillin [2]. Resistance of enterobacteria to the β-lactam antibiotics is associated to the presence of genes encoding for β-lactamases [2,3]. These β-lactamases are frequently encoded by the *bla* genes and are part of the CTX-M, TEM, SHV and OXA families, sharing different hydrolytic profiles [2]. In the TEM family, mutations of individual nucleotides in genes encoding β-lactamases
of the TEM-1 type change the phenotype of BLEE. A wide diversity of members of the TEM family is known, which facilitates the search for transmission of individual resistance genes between strains [2,10]. The amino acid sequence of the SHV-1 family has a high rate of coincidence with TEM-1, sharing its molecular structure. The CTX-M family shows greater activity against cefotaximes than cephtazidime (third-generation cephalosporins), because of the geometry of the antibiotic binding site, which allows efficient recognition of cefotaxime, but not of cephtazidime (a larger molecule). The CTX-M producing strains have the same clinical implications as the production of other BLEE. Nowadays, the number of variants of this family is increasing [2,10]. The main members of the OXA family show low level hydrolytic activity towards clavulanic acid and high level towards carbapenemes. They are the most complex strains to detect and, due to their extensive resistance to \(\beta\)-lactams, therapy has been limited, resulting in high mortality (conditioned by host, type of infection and treatment administered). OXA-48 producing \(K.\ pneumoniae\) is dangerously common in healthcare centers. These are the strains that cause most of the bacteremia outbreaks in hospitals throughout Europe and the Middle East.

The European Commission and the European Centre for Disease Prevention and Control (ECDC) have proposed to increase surveillance and control of outbreaks, in order to reduce the impact of infectious diseases caused by multi-drug resistant organisms on the population [2,10,11]. \(K.\ pneumoniae\) resistant strains to multiple drugs are being more frequently isolated, and this leads to the phenotyping techniques becoming little sensitive or unspecific, making it difficult to identify the strain causing the nosocomial disease. Table 2 shows the main families of plasmids that confer resistance to clinically relevant antibiotics, such as extended spectrum cephalosporins, fluoroquinolones and aminoglycosides. The number of references obtained for TEM, KPC and CTX-M type enzymes is very high compared to the rest of the enzymes, supporting their importance for the health field and the scientific community. In the 1970s, most outbreaks of healthcare associated infections (HAI) caused by \(K. pneumoniae\) were due to aminoglycoside resistant strains. Between 1990 and 2000, outbreaks caused by BLEE producing strains were described, while those caused by carbapenemic resistant strains were described from 2000 onwards. Today, strains producing carbapenemes such as KPC and NDM-1, are significant in clinical practice [11].

On the one hand, \(K. pneumoniae\) NDM-1 is a zinc dependent metalo-\(\beta\)-lactamase that was first identified in 2009 from a New Delhi patient. This enzyme, anchored to the external membrane, can capture the zinc present in the host. In addition, vesicles can carry the active enzyme NDM-1 by exocytosis. Therefore, it can reach other bacteria and bind to the membrane acting as if it were also expressed by other bacteria, making them temporarily resistant [12]. Furthermore, NDM-1 is encoded by the plasmid gene blaNMD-1 and can reach other strains through HGT. Strains producing this enzyme hydrolyze all \(\beta\)-lactam antibiotics, except aztreonam. Among the possible inhibitors studied, hesperidins show high activity against NDM-1. Due to the low toxicity hesperidins contain, they are on the list of possible NDM-1 inhibitor drugs [12]. Since 2009, nearly 650 articles have been reported in PubMed on the presence of NDM-1 in \(Klebsiella\).

On the other hand, KPC is an enzyme capable of conferring resistance to carbapenemes, as well as inactivating penicillins, cephalosporins or monobacterics. It was first identified in 2001 in the United States from a superbug that had suffered a mutation that conferred resistance to multiple drugs [13]. Bacteria with KPC enzymes are more virulent because they have acquired resistance to multiple antibiotics, leading to a high mortality rate in immunocompromised patients. Research confirms that previous use of carabapenemes increases the likelihood of a patient developing a HAI caused by KPC-producing bacteria [13]. However, there is some controversy in the scientific community about this claim, as a study on the clinical impact of carabpenemically resistant \(K. pneumoniae\) concluded that previous use of fluoroquinolone and broad-spectrum cephalosporin was independent of infections with KPC producing organisms [14].
Table 2. Review of resistant plasmid families associated with AmpC, BLEE, 16S rRNA methylases, Qnr proteins and MBLs in *K. pneumoniae*.

| Enzyme                      | Replicons | No. of Plasmids | Genbank References |
|-----------------------------|-----------|-----------------|--------------------|
| CMY-8 (CTX-M-3)             | HI2       | 1               | 3                  |
| CMY-36                      | ColE      | 1               | 7                  |
| CTX-M-3 (ArmA)              | L/M       | 73              | 11                 |
|                             | HI2       | 68              |                    |
| CTX-M-9                     | P         | 10              | 636                |
|                             | I1        | 8               |                    |
| CTX-M-14                    | FII, FIB  | 13              | 11,375             |
| CTX-M-15 [TEM-1, AAC(6′)-IB-CR] | FII, FIA, FIB | 152          | 17                 |
|                             | NT        | 22              |                    |
| CTX-M-17                    | ColE      | 1               | 5                  |
| CTX-M-62                    | NT        | 1               | 5                  |
| SHV-5                       | L/M       | 118             | 397                |
|                             | FII, FIB  | 2               |                    |
|                             | I1        | 12              |                    |
|                             | K         | 5               | 1189               |
| SHV-12                      | FII, FIB  | 9               |                    |
|                             | FI1K      | 1               |                    |
| TEM-1                       | ColE      | 2               | 67,442             |
| KPC-2                       | N         | 1               | 29,077             |
| IMP-4 (QnrB2, QnrB-4, ArmA) | L/M       | 22              | 1888               |
|                             | A/C       | 15              |                    |
|                             | I1        | 17              |                    |
| VIM-1                       | N         | 15              | 2379               |
|                             | NT        | 1               |                    |
| VIM-4 (CMY-4) ArmA          | A/C       | 2               | 5                  |

4. Conclusions

The massive and uncontrolled use of antimicrobials in clinical practice (mainly the β-lactam type) has contributed to the accelerated development of antimicrobial resistance. The increase in nosocomial infections caused by BLEE-producing enterobacteria such as *K. pneumoniae* is generating a global economic stress due to the numerous treatments that have been compromised. Therefore, the main objective is not to expand the therapeutic arsenal, but to develop other techniques far from the direct use of antimicrobials, reducing the effects and impact of multi-drug resistant organisms on the population.

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Abbreviations

ESBL  Extended Spectrum β-Lactamase
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