Characteristics of Klebsiella Pneumoniae ST4 Coharboring qnrB1, aac-Ib-cr, CTX-M-15 and SHV 11 in a Tunisian Hospital

MARIEM Nasri Yaiche (mariem_nasri@hotmail.com)  
The University of Manchester Faculty of Biology Medicine and Health  https://orcid.org/0000-0002-8751-3647

Ikbel Denden Rafraf  
Faculty of pharmacy

qinglan guo  
Fudan University Huashan Hospital Institute of Antibiotics

Maha Mastouri  
Faculty of Pharmacy

Mahjoub Aouni  
Faculty of pharmacy

Minggui Wang  
Fudan University Huashan Hospital Institute of Antibiotics

Research article

Keywords: PMQR, ESBL, MLST, K.pneumonia, conjugation, transformation

DOI: https://doi.org/10.21203/rs.3.rs-42514/v1

License: ☑️ This work is licensed under a Creative Commons Attribution 4.0 International License. Read Full License
Abstract

Background: Multiresistant Klebsiella pneumoniae are predominant cause of hospital acquired infection. This work describes the molecular epidemiology of these isolates in Tunisian Hospital.

Methods: Between October 2010 and June 2013, 50 non-duplicated clinical K. pneumoniae were selected on the basis of nalidixic acid (NA) resistance and were characterized. Isolates were identified using APi 20E system. Susceptibility testing was determined using the disc diffusion method and the microdilution technique to determine the MIC of ciprofloxacin.

PMQR and ESBL genes were detected by PCR and positive results were confirmed by direct sequencing of PCR products. Multilocus sequence typing (MLST) was performed to determine the genetic relationship among isolates. Conjugation and transformation were done to know if PMQR and ESBL were carried with one or two plasmids.

Results: A total of 20 PMQR harboring K. pneumoniae representing 40% of all NA resistant isolates were characterized. Among PMQR positive K. pneumoniae 13 were resistant to amoxicilline, amoxicilline/clavulanic acid, ticarcilline, piperacilline, cefaloridine, cefotaxime (CTX) and ceftazidime. The rate of resistance to gentamicine, tobramicine and amikacine were 85%, 95% and 25% respectively. Out of 20 K. pneumoniae (60%) were qnr positive (1 qnrA6 and 11 qnrB1) and (60%) were aac-lb-cr positive. 33.3% coharbored the aac-lb-cr and qnrB1 determinants. Out of all PMQR positive strains, 65% harbored ctx-M-15 gene. It was associated to shv11 in three cases and tem1 in two cases. The predominant types were ST4 (35%) and ST15 (20%). ST 101(15%) and ST 147 (10%) come in second order. one case of each ST14,ST86,ST336 and ST307 were also observed. qnrB1, aac-lb-cr, Ctx-m-15 can be carried with more than one plasmid in the same bacteria.

Conclusion: The co-existing of different genes conferring resistance among the same and different family of antibiotics is a big threat to patient because it limits the therapeutic process. This phenomonen is a problem of concern that needs to improve the resistance surveillance of multi gene carrying K pneumoniae.

1. Background

Pathogenic Klebsiella pneumoniae resistant to quinolones and β-lactam drugs is widely recognized as important bacteria causing array of diseases [1]. Antibiotic sensitivity assays have revealed high frequencies of quinolone drug resistance in extended spectrum beta lactamases (ESBL) producing clinical isolates [1]. The distribution of qnr and ESBL alleles varies among countries. In 1998, the first plasmid-mediated mechanism of resistance to quinolones was described in multiresistant K. pneumoniae harbouring the qnrA gene on pMG252 plasmid and associated to the cephalosporinase Fox-5 [2,3]. Several studies have found the qnr determinants mainly qnrA and qnrB that were more frequent in ESBL
producing Enterobacteriacae; the genes were carried by large plasmids (40-320Kb) in Class1 integron type sul1,qnr S was more associated to tem-1 and lap-1[4,5,6,7,8]

The aac-lb-cr gene was found in various integrons, especially on IncF11 plasmid expressing ctx-m-15 that have spread rapidly so that ctx-m-15 has become predominant ESBL in many countries around the world [9,10,11,12,13,14,15,16,17,18].

Aac-lb-cr has been associated with other PMQR genes including qnrA1[19,20,21], qnrB1 [6], qnrB2 [17,22], qnrB4 [19,23], qnrB6 [19,23], qnrB10 [24], qnrS1 [19,23], qnrS2[25] and qepA [23] and with other β-lactamases including, ctx-m-1 [26], ctx-m15 [6], ctx-M14 [19], CTX-M-24[19], DHA-1 [23], SHV-12[23].

Very few studies have reported on the problem of the coexisting of PMQR and ESBL in K. pneumonia in Africa. [4,6]

In Tunisia, qnrA6 and CTX-m-15 were the predominant allele among K.pneumoniae isolates [4, 27,28]. qnrB1 coexisted with ctx-m-15, SHv-28 and Tem-1 in 125 Kb plasmid through the study of Dahmen et al in a Tunisian hospital [4].

Despite the high prevalence of these isolates in nosocomial infections, large studies to investigate the molecular epidemiology of these isolates in Africa are still lacking and the sequence type of cohaboring PMQR and ESBL K.pneumoniae are not yet known.

We therefore conducted the first epidemiological study in Africa to determine the molecular epidemiologie of PMQR and ESBL coproducing k.pneumoniae. This is also the first report of aac-lb-cr gene among K.pneumoniae despite its huge distribution in several countries.

2. Methods

2.1 Sample selection

In total, 50 non-duplicate clinical K.pneumoniae isolates recovered between October 2010 and June 2013 at Fattouba Bourguiba hospital (Tunisia) were selected on the basis of nalidixic acid resistance.

The K.pneumoniae isolates were identified by biotyping by using API20E system ((bioMérieux SA, Marcy-l’Etoile, France).

2.2 Antimicrobial susceptibility testing

The antibiotics susceptibilities of the K.pneumoniae isolates were determined on Mueller-Hinton agar by the standard disk diffusion procedure as described by the Antibiogram Committee of the French Society for Microbiology (http://www.sfm.asso.fr).

The microbroth dilution technique was used to determine the MIC for ciprofloxacin.
2.3 PMQR and ESBL screening and DNA sequencing.

The *qnrA*, *qnrB*, and *qnrS*, *qnrD*, *aac-Ib*, *qepA*, *gyrA*, Ctx-M1G, *SHV* and *Tem* genes were detected by PCR with the primers described previously [29,30,31,32,33,34,35,36,37].

Positive results were confirmed by direct sequencing of PCR products.

Analysis and comparison of nucleotide and amino acid sequence data were carried out using Lasergene software ((version 7.1; DNASTAR, Wisconsin, USA) and programs available from the national Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov).

2.4 Multi locus sequence typing (MLST)

The genetic relationship among *K. pneumoniae* isolates was assessed based on the Multilocus Sequence Typing Method as described previously on the website (http://www.pasteur.fr/recherche/genopole/PF8/mlst/Kpneumoniae.html). PCR for seven housekeeping genes; *rpoB*, *gapA*, *mdh*, *pgi*, *phoE*, *infB* and *tonB* was conducted and products were directly sequenced. Analysis was carried out as described on the website.

2.5 Location of PMQR and ESBL genes

Conjugation

The transferability of *qnr*, aac-Ib-cr and ctx-m-15 genes between the *K. pneumoniae* isolates and the *E. coli* J53Az· recipient were carried out in LB broth. Cultures of donor and recipient cells in logarithmic phase (0.5 ml of each) were added to 4 ml of fresh LB broth and incubated overnight without shaking. Transconjugants were selected on Trypticase soy agar (TSA) plates containing sodium azide (100 µg/ml; Sigma Chemical Co., St. Louis, Mo.) for counterselection and cefotaxima (10 µg/ml) to select for plasmid-encoded resistance.

To determine if quinolone and ESBL resistance were cotransferred, colonies were replicated onto TSA with and without ciprofloxacin (0.06 µg/ml). MICs of ciprofloxacin for the donor, recipient, and transconjugant strains were measured by E-test.

Transformation

For transformation reactions, Top 10 competent cells and plasmid extracted from transconjugants were taken to ice for 2 min. 5 to 10µl of plasmid were taken and added to 100µl of competent cells. The mixture was taken to ice for 30 min, then to 42°C for 1 min. After this it was taken to ice again for 3 min without mixing. After this step 900µl of LB broth were added to each mixture then taken to 37°C for one hour with checking.

100µl of transformants were plated on 3 LB agar containing separately Amp (1µg/ml), Cip (0.06µg/ml) and Ctx (1µg/ml).
3. Results

3-1 Distribution of PMQR positive isolates

Out of 50 *K pneumonia* resistant to quinolones, 20 (40%) harbored PMQR over a period of three years. Table I characterize these isolates.

Table I: Features of PMQR positive *Klebsiella pneumoniae* isolates.
| Strain code | Date of isolation | Sample | Unit       | PMQR | ST | MIC of CIP µg/ml | ESBL genes |
|-------------|------------------|--------|------------|------|----|----------------|------------|
| 154         | 3/12/10          | Blood  | Pediatric  | qnr B1 | 4  | 128             | blaCtx-m-15, blushv11 |
|             |                  |        |            | aac-lb-cr |    |                 |            |
| 2116        | 18/03/11         | Blood  | Pediatric  | qnr B1 | 4  | 512             | blaCtx-m-15 |
|             |                  |        |            | aac-lb-cr |    |                 |            |
| 3142        | 31/05/11         | Urine  | Nephrology | aac-lb-cr | 101 | 256            | blaCtx-m-15 |
| 4144        | 27/05/11         | Pus    | ICU        | qnr A6 | 147 | 512            | -          |
| 5189        | 28/12/11         | Blood  | Pediatric  | qnr B1 | 4  | 32              | blaCtx-m-15 |
|             |                  |        |            | aac-lb-cr |    |                 |            |
| 6208        | 7/04/12          | Blood  | Surgery    | qnr B1 | 14 | 16              | blaCtx-m-15, blushv11 |
|             |                  |        |            | aac-lb-cr |    |                 |            |
| 7228        | 29/01/13         | Blood  | ICU        | aac-lb-cr | 147 | >1024         | blaCtx-m-15, blashv11 |
| 880         | 25/01/11         | Urine  | nephrology | aac-lb-cr | 15 | 512            | blaCtx-m-15, blatem1 |
| 99          | 30/10/10         | Pus    | Pediatric  | qnr B1 | 86 | 32             | -          |
| 10127       | 09/05/11         | Urine  | nephrology | aac-lb-cr | 15 | 128            | blaCtx-m-15 |
| 11113       | 15/03/11         | Urine  | nephrology | aac-lb-cr | 15 | 256            | blaCtx-m-15 |
| 12231       | 30/01/13         | Urine  | nephrology | aac-lb-cr | 101 | 256         | blaCtx-m-15 |
| 1387        | 9/02/11          | T.S    | ICU        | aac-lb-cr | 101 | >1024        | blaCtx-m-15 |
| 1442        | 22/11/10         | Urine  | Nephrology | aac-lb-cr | 15 | 256            | blaCtx-m-15 |
All of PMQR harboring *K. pneumoniae* were from inpatients. Samples were gathered from different organs and systems: urine (45%), blood (25%), pus (20%) and respiratory samples (10 %). They were collected from patients admitted in Pediatrics (40%), nephrology (30 %), intensive care units (20 %), cardiology (5 %), and surgery (5 %).

During the survey period of quinolone resistance in *K. pneumoniae*, all of the strains had high MIC of ciprofloxacin (16-1024µg/mL). A higher rate of resistance to commonly used non-quinolones was observed. Among PMQR harbouring *K.pneumoniae* isolates 13 were resistant to Amoxicillin, Amoxicillin/clavulanic Acid, ticarcillin, piperacillin, cefaloridine, cefotaxim and ceftazidim. As well as, they possessed an ESBL by phenotypic testing. The rate of resistance to gentamicin, tobramicin and amikacin were 85%, 95% and 25% respectively. Resistance to trimethoprime/sulfamides was 60%. All isolates were sensitive to imipenem using the disc diffusion method.

### 3-2 PMQR and ESBL alleles

Out of 20 PMQR harboring *K.pneumoniae*, 60 % were found to contain *qnr* type determinants. The breakdown of the determinants detected was as follows: 1 *qnrA6* and 11 *qnrB1*. The rate of *aac(6')-Ib-cr* gene was 60 % also.

Among the *aac(6')-Ib-cr* positive strains, there were 4 (33.3%) which were also *qnr* positive.

Among the twenty PMQR positive *K.pneumoniae*, 13 (65%) harbored the *blaCTX-M-15* determinant. The previous gene was associated to the *blaSHV11* gene in three cases and coexisted with the *blaTEM1* gene in two cases.

### 3-3 Genetic relatedness
Seven ST4 *K. pneumoniae* variant were isolated in the Pediatric Unit. Four isolates belonging to ST15 were found in the nephrology unit. Three strains assigned to ST101 were recovered from the nephrology unit (n=2) and the ICU (n=1). Tow ST147 *K. pneumoniae* variant were also isolated from the ICU. One case of each ST14, ST86, ST336 and ST307 were isolated from Surgery, Pediatric, ICU and Cardiology respectively.

**3-4 PMQR and Ctx-M-15 genes transferability**

**Conjugation**

A: transconjugants 154\(^{c+I}\) of donor 154 inoculated in CTX+azide plate. B: transconjugants 154\(^{c+I}\) of donor 154 inoculated in CIP+azide plate. C: transconjugants 5189\(^{c}\) of donor 5189 inoculated in CTX+azide. D: transconjugant 5189\(^{c+I}\) of donor 5189 inoculated in ctx+azide plate. E: transconjugants 5189\(^{c+I}\) inoculated in Cip+azide plate. F: transconjugants 2116\(^{c+I}\) inoculated in ctx+azide plate, G: transconjugants 2116\(^{c+I}\) inoculated in ctx+azide plate.

Notes:

c+I: the same transconjugants give colonies on CTX+azide plate and on CIP +azide plate.

c: the transconjugant give colonies on CTX+azide plate only.

All the conjugation reactions between *Ecoli* J53 and the donor 154 or the donor 2116 has given transconjugants able to give colonies on CTX+azide plates and Cip+azide plates.

However, conjugation reactions between *E.coli* J53 and the donor 5189 have given two types of transconjugants. First type: when inoculated from CTX+azide plate to Cip+azide plate they give colonies. Second type gives colonies only on CTX+ azide plates.

**Table II: Genes detected in transconjugants and the corresponding ciprofloxacin MIC**

| Donor  | 154 | 5189 | 2116 |
|--------|-----|------|------|
| Transconjugants | A | B | C | D | E | F | G |
| Qnr B1 | + | + | - | + | + | + | + |
| Aac-Ib-cr | + | + | + | + | + | + | + |
| Ctx-m-15 | + | + | + | + | + | + | + |
| MIC Cip µg/ml | 0,5 | 0,5 | 0,064 | 0,5 | 0,5 | 0,5 | 0,5 |

**Transformation**
A': transformant with plasmid A, B': transformant with plasmid B, C': transformant with plasmid C, D': transformant with plasmid D, E': transformant with plasmid E, F': transformant with plasmid F, G': transformant with plasmid F, H': transformant with plasmid G.

Table III: Genes detected on transformants and the corresponding ciprofloxacin MIC

| Plasmid donor | 154 | 5189 | 2116 |
|---------------|-----|------|------|
| Transformants | A   | B   | C   | D   | E   | F   | G   | H   |
| Qnr B1        | +   | +   | -   | -   | +   | -   | +   | +   |
| Aac-Ib-cr     | +   | +   | +   | +   | +   | +   | +   | +   |
| Ctx-m-15      | +   | +   | +   | +   | +   | +   | +   | +   |
| MIC cip µg/ml  | 0,19 | 0,19 | 0,008 | 0,006 | 0,19 | 0,008 | 0,19 | 0,19 |

Plasmid extraction from transformants has shown the harbor of one plasmid in each one. Consequently, all genes detected in one transformant were carried in the same plasmid. Therefore, the donors 154 and 2116 are carrying qnrB1, aac-Ib-cr and CTX-m-15 in the same plasmid. However, the donor number 5189 contains two types of plasmids, the first one is carrying the three cited genes (transformant E) and the second one is carrying aac-Ib-cr and CTX-m-15.

3-Discussion

The results of this study provide insights into the molecular epidemiology of PMQR and ESBL producing *K. pneumoniae* isolates in the university hospital Fattouma Bourguiba in Monastir, Tunisia, thereby representing the first study to characterize PMQR and ESBL producing *K. pneumoniae* in Tunisia.

During our three years study, most of the *K. pneumoniae* was isolated from patients with urinary tract infection at inpatient department. Blood was the second site colonized by these bacteria. It was shown that multidrug resistant *K. pneumoniae* is one of the important nosocomial and community acquired opportunistic bacteria causing urinary tract infection, pneumonia, septicemia etc [1].

The majority of the isolates characterized in this study were from pediatric units. Infections caused by PMQR and ESBL harboring organisms in neonates and pediatric wards are usually reported to be hospital acquired and associated with invasive procedures [38,39,40]. The predominance of these isolates in these units could be explained by the high use of the third generation of cephalosporins, as cefotaxim was the most prescribed drug in pediatric unit during the study period. Comparing to other units, the MIC of ciprofloxacine (16-64µg/ml) was lower in pediatric wards as ciprofloxacine is usually contraindicated in children and neonates. High rates of ciprofloxacine resistance were detected in the ICU, the second ward
contaminated with multidrug resistant *K. pneumonia*. These units have high rates of empirical treatment using fluoroquinolones, 3rd generation cephalosporins and also carbapenems.

Among quinolone resistant *K. pneumoniae*, 40% only harbored the PMQR genes. It has been demonstrated that quinolone resistance in Enterobacteriaceae family usually results from mutations in genes carried by chromosomally encoded type II topoisomerases, efflux pump or porin related protein [41].

PMQR such as *qnr* and *aac(6')-Ib-cr* may facilitate the spread and increase the prevalence of quinolone resistant strains. To date, *qnr* genes have been widely detected in Africa, Asia, America and Europe. The co-harbour of *aac(6')-Ib-cr* and *qnr* genes have been reported in several studies. In *K. pneumoniae* the *aac(6')-Ib-cr* gene was related to *qnr B1* [6], to *qnr B32* [5], to *qnrS1*, *qnrB6* and *qnrB4* [42]. It was also shown that PMQR and ESBL could be cotransferred in several studies and this is amplify the infection risk due to these strains of *K. pneumoniae* [1,4,5,42]

This is the first report in Tunisia of coexisting *aac(6')-Ib-cr* and *qnrB1* in *K. pneumoniae*. Moreover, these two PMQR coexisted with bla CTX-m15 and shv11 in three cases and with ctx-m15 and tem1 in tow cases. To the best of our knowledge, this is the first report of coexisting *qnrB1*, *aac-lb-cr*, *ctx-m-15* and *shv11/tem1* in *K. pneumoniae*. Referring to conjugative reactions, *qnrB1*, *aac-lb-cr* and *ctx-m-15* were cotransferred from tow donors.

Plasmids extracted from the corresponding transformants were the same in size and there were only one plasmid for each transformant. This is confirming that these three genes were carried by the same plasmid. The third donor gave transconjugants with *qnrB1*, *aan-LB-cr* and *ctx-m-15* or transconjugants with *aac-lb-cr* and *ctx-m-15* only. This is had already influence on MIC of ciprofloxacin on transconjugants. Effectively, the MIC of ciprofloxacin on transconjugants with *qnrB1* and *aac-lb-cr* genes was higher (0.5µg/ml) then those with *aac-lb-cr* only (0.064µg/ml). This is mean the expression of *qnrB1* in resistance to ciprofloxacin is match more important than the expression of *aac-lb-cr*. The transformation of top 10 with plasmid exracted from transconjugants coming from the third donor has shown tow types of transformants; the fist type is carrying one plasmid harboring the three cited genes (transformant E), the second type is carrying one smaller plasmid harboring *aac-lb-cr* and *ctx-m-15* only. Referring to the previous data the donor 1 and 3 are harboring one plasmid that carry 3 genes and the third donor harbor tow type of plasmids the first carry the 3 genes and the second carry *aac-lb-cr* and *ctx-m-15*. Bought type of donor with one or tow plasmid are prohibiting the patient health and this is need important survey of the epidemiology of these strains.

This is the first epidemiological report in Tunisia using MLST to type strains of *K. pneumoniae* harboring PMQR. The results revealed mainly sequence type 4, 15,101 and 147 compared to previous reported ST 101, ST 107 and ST147 [27]. There is a significant distribution of the ST and the corresponding unit. The ST4 *K. pneumoniae* were found in pediatric wards, the ST 15 in nephrology unit, ST147 in ICU and the ST101 between the nephrology and the ICU. This phenomenon can be explained by
the nosocomial spread of strains with the same sequence type among patients of the same unit. The study of Filippa et al, have demonstrated the spread of an unusual \textit{K.\textit{pneumoniae}} isolated from Nigerian patient among 10 French patients in the ICU of French hospital [6]. The ST4 and ST 15 \textit{K.\textit{pneumoniae}} were first described in France, but until 2013, the \textit{ctx-m-15} doesn't exist in France and it was introduced among this country because of a strain coming from Nigerian patient [6]. The nosocomial contamination between patients could explain the harbor of new genes in ST4 and ST15 \textit{K.\textit{pneumoniae}}. The ST101 and ST147 \textit{K.\textit{pneumoniae}} were first reported respectively in Poland and Hungary. Since that these sequence type have been reported to occur worldwide and are associated with multidrug resistant \textit{K.\textit{pneumoniae}} [43,44]. The ST14 found in the surgery unit was described previously in Africa [28]. The ST86, ST336 and ST307 are strange to Africa and this is the first report of these types in Tunisia since they were described respectively in France, America and Netherlands. Tunisia has important relationship with the cited countries and this can explain the first appearance of these ST types \textit{K.\textit{pneumoniae}}.

4. Conclusion

This study highlights the need to improve the antimicrobial resistance surveillance in Tunisia for \textit{K.\textit{pneumoniae}} to monitor the new types of resistance mechanisms emerging in different ward in the same or different hospitals. As well as, the factors responsible for the selection and dissemination of the high conjugative plasmids encoding the \textit{qnr} genes, \textit{aac-ib-cr} determinant and different ESBL must be considered for the antibiotic policies and controlled to prevent other outbreaks in future.

**Abbreviations**

- CIP: ciprofloxacin
- CTX: cefotaxime
- ESBL: Extended spectrum beta lactamases
- ICU: Intensive care unit
- MIC: Minimal inhibitory concentration
- MLST: Multi locus sequence typing
- NA: nalidixic acid
- PMQR: Plasmid mediated quinolone resistance
- ST: Sequence type

**Declarations**
Ethics approval and consent to participate: Not applicable because strains were already collected thinks to the clinical microbiological laboratory of Fattouma Bourguiba Hospital.

Consent for publication: Not applicable

Availability of data and material: Not applicable

Competing interests: The authors declare that they have no competing interests

Funding:

Experiments of MLST, conjugation, transformation and plasmid extraction were performed in the Institut of Antibiotics, Huashan hospital in Shanghai during the training period wish was financed by the Chinese government scholarship.

Authors’ contributions: MNY carried the bacteria collection and typing antibiotic susceptibility, PCR screening, conjugation and transformation reactions, sequence blasting and writing the manuscript. IDR is project designer and co-supervisor of research work. QG Laboratory supervisor, she controlled the analysis, interpretation of data and controlled the manuscript. MM provided the necessary materials for collection, typing and doing the susceptibility tests. MA project designer and general supervisor of research group. MW General supervisor of research experiments and controlled the manuscript. All authors read and approved the final manuscript.

Acknowledgements: not applicable

References

[1] Tripathi A, Dutta SK, Majumdar M, Dhara L, Banerjee D, Roy K (2012) High Prevalence and Significant Association of ESBL and QNR Genes in Pathogenic Klebsiella pneumoniae Isolates of Patients from Kolkata, India. Indian J Microbiol 52(4):557–564.

[2] Martinez-Martinez LA, Pascual A and Jacobey GA (1998) quinolone resistance from transferable plasmid. Lancet 351:797-799.

[3] Strahilevitz J, Jacobey GA, Hooper DC and Robicsek A (2009) Plasmid-Mediated Quinolone Resistance: a Multifaceted Threat. Clin Microbiol Rev 664-689.

[4] Dahmen S, Poirel L ,Mansour W, Boualle`gue O and Nordmann P (2010) Prevalence of plasmid-mediated quinolone resistance determinants in Enterobacteriaceae from Tunisia. Clin Microbiol Infect 7:1019-23.

[5] Wang D,Wang H, Qi Y, Liang Y,Zhang J, Yu L (2012) Characteristics of Klebsiella pneumoniae harbouring QnrB32, Aac(6’)-Ib-cr, GyrA and CTX-M-22 genes. Folia Histochem Cyto 50 : 68-74.
[6] Filippa N, Carricajo A, Grattard F, Fascia P, El Sayed F, Defilippis JP, Berthelot P and Aubert G (2013) Outbreak of multidrug-resistant Klebsiella pneumoniae carrying qnrB1 and blaCTX-M15 in a French intensive care unit. Annals of Intensive Care 3:18

[7] A. Merens, Servonnet A (2010) Mécanismes et épidemiologie de la résistance aux fluoroquinolones en 2010. RFL N°422 pp-31-41

[8] Bourouis A, Dubois V, Chihi H, Beladj C, Ben Moussa M, Quentin C et al (2010) Detection of plasmid-mediated quinolone resistance determinant QnrS1 in an extended-spectrum β-lactamase-producing Enterobacter cloacae in Tunisia. Ann Microbiol 60:423–427.

[9] Jouini A, Ben Slama K, Vinué L, Ruiz E, Saenz Y, Somalo S, Klibi N, Zarazaga M, Ben Moussa M, Boudabous A, Torres C (2010) Detection of unrelated Escherichia coli strains harboring genes of CTX-M-15, OXA-1, and AAC(6')-Ib-cr enzymes in a Tunisian hospital and characterization of their integrons and virulence factors. J Chemother 5:318-23.

[10] Boyd, D. A., S. Tyler, S. Christianson, A. McGeer, M. P. Muller, B. M. Willey, E. Bryce, M. Gardam, P. Nordmann, and M. R. Mulvey (2004) Complete nucleotide sequence of a 92-kilobase plasmid harboring the CTX-M-15 extended-spectrum beta-lactamase involved in an outbreak in long-term-care facilities in Toronto, Canada. Antimicrob. Agents Chemother. 48:3758–3764.

[11] Cordeiro, N. F., L. Robino, J. Medina, V. Seija, I. Bado, V. Garcia, M. Berro, J. Pontet, L. Lopez, C. Bazet, G. Rieppi, G. Gutkind, J. A. Ayala, and R. Vignoli (2008) Ciprofloxacin-resistant enterobacteria harboring the aac(6)-Ib-cr variant isolated from feces of inpatients in an intensive care unit in Uruguay. Antimicrob. Agents Chemother. 52:806–807.

[12] Fihman, V., M. F. Lartigue, H. Jacquier, F. Meunier, N. Schnepf, L. Raskine, J. Riahi, M. J. Sanson-le Pors, and B. Bercot (2008) Appearance of aac(6)-Ib-cr gene among extended-spectrum β-lactamase-producing Enterobacteriaceae in a French hospital. J. Infect. 56:454–459.

[13] Karisik, E., M. J. Ellington, R. Pike, R. E. Warren, D. M. Livermore, and N. Woodford. (2006) Molecular characterization of plasmids encoding CTXM-15 β-lactamas from Escherichia coli strains in the United Kingdom. J.Antimicrob. Chemother. 58:665–668.

[14] Machado, E., T. M. Coque, R. Canto´n, F. Baquero, J. C. Sousa, and L Peixe (2006) Dissemination in Portugal of CTX-M-15-, OXA-1-, and TEM-1-producing Enterobacteriaceae strains containing the aac(6)-Ib-cr gene, which encodes an aminoglycoside- and fluoroquinolone-modifying enzyme. Antimicrob. Agents Chemother. 50:3220–3221.

[15] Pallecchi, L., A. Bartoloni, C. Fiorelli, A. Mantella, T. Di Maggio, H.Gamboa, E. Gotuzzo, G. Kronvall, F. Paradisi, and G. M. Rossolini (2007) Rapid dissemination and diversity of CTX-M extended-spectrum β-lactamase genes in commensal Escherichia coli isolates from healthy children from low-resource settings in Latin America. Antimicrob. Agents Chemother. 51:2720–2725.
[16] Pitout, J. D. D., Y. Wei, D. L. Church, and D. B. Gregson (2008) Surveillance for plasmid-mediated quinolone resistance determinants in Enterobacteriaceae within the Calgary Health Region, Canada: the emergence of aac(6\_)-Ib-cr. J. Antimicrob. Chemother. 61:999–1002.

[17] Pomba, C., J. D. da Fonseca, B. C. Baptista, J. D. Correia, and L. Martínez-Martínez. (2009) Detection of the pandemic O25-ST131 human virulent Escherichia coli CTX-M-15-producing clone harboring the qnrB2 and aac(6\_)-Ib-cr genes in a dog. Antimicrob. Agents Chemother. 53:327–328.

[18] Sabtcheva, S., M. Kaku, T. Saga, Y. Ishii, and T. Kantardjiev (2009) High prevalence of the aac(6\_)-Ib-cr gene and its dissemination among Enterobacteriaceae isolates by CTX-M-15 plasmids in Bulgaria. Antimicrob. Agents Chemother. 53:335–336.

[19] Jiang, Y., Z. Zhou, Y. Qian, Z. Wei, Y. Yu, S. Hu, and L. Li (2008) Plasmid mediated quinolone resistance determinants qnr and aac(6\_)-Ib-cr in extended-spectrum \_\_lactamase-producing Escherichia coli and Klebsiella pneumoniae in China. J. Antimicrob. Chemother. 61:1003–1006.

[20] Robicsek, A., J. Strahilevitz, G. A. Jacoby, M. Macielag, D. Abbanat, C. H. Park, K. Bush, and D. C. Hooper (2006) Fluoroquinolone-modifying enzyme: a new adaptation of a common aminoglycoside acetyltransferase. Nat. Med. 12:83–88.

[21] Xu, X., S. Wu, X. Ye, Y. Liu, W. Shi, Y. Zhang, and M. Wang (2007) Prevalence and expression of the plasmid-mediated quinolone resistance determinant qnrA1. Antimicrob. Agents Chemother. 51:4105–4110.

[22] Yang, H., H. Chen, Q. Yang, M. Chen, and H. Wang (2008) High prevalence of plasmid-mediated quinolone resistance genes qnr and aac(6\_)-Ib-cr in clinical isolates of Enterobacteriaceae from nine teaching hospitals in China. Antimicrob. Agents Chemother. 52:4268–4273.

[23] Wu, J. J., W. C. Ko, C. S. Chiou, H. M. Chen, L. R. Wang, and J. J. Yan (2008) Emergence of Qnr determinants in human Salmonella isolates in Taiwan. J. Antimicrob. Chemother. 62:1269–1272.

[24] Quiroga, M. P., P. Andres, A. Petroni, A. J. Soler Bistue´, L. Guerriero, L. J. Vargas, A. Zorreguieta, M. Tokumoto, C. Quiroga, M. E. Tolmasky, M. Galas, and D. Centro´n. (2007) Complex class 1 integrons with diverse variable regions, including aac(6\_)-Ib-cr, and a novel allele, qnrB10, associated with ISCR1 in clinical enterobacterial isolates from Argentina. Antimicrob. Agents Chemother. 51:4466–4470.

[25] Pica˜o, R. C., L. Poirel, A. Demarta, C. S. Silva, A. R. Corvaglia, O. Petrini, and P. Nordmann (2008) Plasmid-mediated quinolone resistance in Aeromonas allosaccharophila recovered from a Swiss lake. J. Antimicrob. Chemother. 62:948–950.

[26] Soge, O. O., B. A. Adeniyi, and M. C. Roberts (2006) New antibiotic resistance genes associated with CTX-M plasmids from uropathogenic Nigerian Klebsiella pneumoniae. J. Antimicrob. Chemother. 58:1048–1053.
[27] Elihani D, Bakir L, Aouni M, Passet V, Arlet G, Brisse S, Weill FX: Molecular epidemiology of extended-spectrum beta-lactamase-producing *Klebsiella pneumoniae* strains in a university hospital in Tunis, Tunisia, 1999–2005. Clin Microbiol Infect 2010, 16(2):157–164.

[28] Mshana SE, Hain T, Domann E, Lyamuya EF, Chakraborty T and Imirzalioglu C (2013) Predominance of *Klebsiella pneumoniae* ST14 carrying CTX-M-15 causing neonatal sepsis in Tanzania. BMC Infectious Diseases 13:466.

[29] Cattoir V, Poirel L, Rotimi V, Soussy C and Nordmann P (2007) Multiplex PCR for detection of plasmid-mediated quinolone resistance *qnr* genes in ESBL-producing enterobacterial isolates. J Antimicrobial Chemother 60, 394–397

[30] Minghua W, Qinglan G, Xiaogang X, Xiaoying W, Xinyu Y, Shi W, Hooper D, and Minggui W (2009) New Plasmid-Mediated Quinolone Resistance Gene, *qnrC*, Found in a Clinical Isolate of *Proteus mirabilis*. Antimicrob Agents Chemother 1892–1897.

[31] Cavaco LM, Hasman H, Xia S, Aarestrup FM (2009) QnrD, a novel gene conferring transferable quinolone resistance in *Salmonella enterica Serovar Kentucky* and *bovismorbificans* strains of human origin. Antimicrob Agents Chemother 58: 603–608.

[32] Park C, Robicsek A, Jacoby GA, Sahm D, and Hooper D (2006) Prevalence in the United States of *aac(6\_)-Ib-cr* Encoding a Ciprofloxacin-Modifying Enzyme. Antimicrob Agents Chemother 3953–3955.

[33] Yamane K, Wachino J, Suzuki S, and Arakawa Y (2008) Plasmid-Mediated *qepA* Gene among *Escherichia coli* Clinical Isolates from Japan. Antimicrob Agents Chemother 1564–1566.

[34] C. Eckert, V. Gautier, M. Saladin-Allard, N. Hidri, C. Verdet, Z. Ould-Hocine, G. Barnaud, F. Delisle, A. Rossier, T. Lambert, A. Philippon, and G. Arlet (2004) Dissemination of CTX-M-Type -Lactamases among Clinical Isolates of Enterobacteriaceae in Paris, France. Antimicrob Agents Chemother, p. 1249–1255

[35] David S. Hammond, Jacqueline M. Schooneveldt, Graeme R. Nimmo, Flavia Huygens, and Philip M. Giffard (2005) *blaSHV* Genes in *Klebsiella pneumoniae*: Different Allele Distributions Are Associated with Different Promoters within Individual Isolates. Antimicrob Agents Chemother p. 256–263

[36] Weill FX, Lailler R, Praud K, Ke’rouanton A, Fabre L, Brisabois A, Grimont PA, and Cloeckaert A (2004) Emergence of Extended-Spectrum–Lactamase (CTX-M-9)-Producing Multiresistant Strains of *Salmonella enterica* Serotype Virchow in Poultry and Humans in France. J Clin Microbiol p. 5767–5773.

[37] Lascols C, Robert J, Cattoir V, Bébéar C, Cavallo J, Podglajen I, Ploy M, Bonnet R, Soussy C, Cambau E (2007) Type II topoisomerase mutations in clinical isolates of *Enterobacter cloacae* and other enterobacterial species harbouring the *qnrA* gene. Int J Antimicrob Agents 402–409.

[38] Cassettari VC, da Silveira IR, Dropa M, Lincopan N, Mamizuka EM, Matte MH, Menezes PR (2009) Risk factors for colonization of newborn infants during an outbreak of extended-spectrum β-lactamase-
producing *Klebsiella pneumoniae* in an intermediate-risk neonatal unit. J Hosp Infect 71:340–347.

[39] Miranda M, Natividad C, Leanos B, Valenzuela A, Garza-Ramos U, Rojas T, Solorzano F, Chihu L, Silva J (2004) Clonal and Horizontal Dissemination of *Klebsiella pneumoniae* Expressing SHV-5 Extended-Spectrum β-Lactamase in a Mexican Pediatric Hospital. J Clin Microbiol 42:30–35.

[40] Crivaro V, Bagattini M, Salza MF, Raimondi F, Rossano F, Triassi M, Zarrilli R (2007) Risk factors for extended-spectrum β-lactamase-producing *Serratia Marcescens* and *Klebsiella pneumoniae* acquisition in a neonatal intensive care unit. J Hospital Infect 67:135–141.

[41] Robicsek A, Jacoby GA, Hooper DC (2006) The worldwide emergence of plasmid-mediated quinolone resistance. *Lancet Infect Dis* 6:629–640.

[42] Ruiz E, Sa´enz Y, Zarazaga M, Rocha-Gracia R, Marti´nez-Marti´nez L, Arlet G and Torres C (2012) qnr, aac(6’)-lb-cr and qepA genes in Escherichia coli and *Klebsiella* spp.: genetic environments and plasmid and chromosomal location. J Antimicrob Chemother 67: 886–897

[43] Dzierzanowska D, Kaminiska W, Semczuk K, Borowiec D, Matysiak M, Szumala-Kakol A, Gierczynski R, Patzer JA (2010) Carriage of genes for various Extended-Spectrum β-Lactamases: a novel resistance strategy of *Klebsiella pneumoniae* in Poland. Int J Antimicrob Agents 35:392–395.

[44] Damjanova I, Toth A, Paszti J, Hajbel-Vekony G, Jakab M, Berta J, Milch H,Fuzi M (2008) Expansion and countrywide dissemination of ST11, ST15 and ST147 ciprofloxacin –resistant CTX-M-15-type β-Lactamase-producing *Klebsiella pneumoniae* epidemic clones in Hungary in 2005-the new MRSA. J Antimicrobial Chemother 62:978–985.

**Figures**
Figure 1

Plasmid profile extracted from transconjugants.
Figure 2

Plasmid profile extracted from transformants.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- supplementaryinformationfile.docx