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Extended-Spectrum-Beta-Lactamases, AmpC Beta-Lactamases and Plasmid Mediated Quinolone Resistance in Klebsiella spp. from Companion Animals in Italy

Valentina Donati¹, Fabiola Feltrin¹, Rene S. Hendriksen², Christina Aaby Svendsen², Gessica Cordaro¹, Aurora García-Fernández³, Serena Lorenzetti¹, Raniero Lorenzetti¹, Antonio Battisti¹,², Alessia Franco¹

¹Istituto Zooprofilattico Sperimentale delle Regioni Lazio e Toscana, Rome, Italy, ²Technical University of Denmark, National Food Institute (DTU-Food), Kongens Lyngby, Denmark, ³Istituto Superiore di Sanità, Department of Infectious, Parasitic and Immune-Mediated Diseases, Rome, Italy

Abstract
We report the genetic characterization of 15 Klebsiella pneumoniae (KP) and 4 isolates of K. oxytoca (KO) from clinical cases in dogs and cats and showing extended-spectrum cephalexin resistance (ESC) resistance. Extended spectrum beta-lactamase (ESBL) and AmpC genes, plasmid-mediated quinolone resistance (PMQR) and co-resistances were investigated. Among KP isolates, ST101 clone was predominant (8/15, 53%), followed by ST15 (4/15, 27%). ST11 and ST340, belonging to Clonal Complex (CC)11, were detected in 2012 (3/15, 20%). MLST on KP isolates corresponded well with PFGE results, with 11 different PFGE patterns observed, including two clusters of two (ST340) and four (ST101) indistinguishable isolates, respectively. All isolates harbored at least one ESBL or AmpC gene, all carried on transferable plasmids (IncR, IncFII, IncI1, IncN), and 16/19 were positive for PMQR genes (qnr family or aac(6’)-Ib-cr). The most frequent ESBL was CTX-M-15 (11/19, 58%), detected in all KP ST101, in one KP ST15 and in both KP ST340. blaCTX-M-15 was carried on IncI1 plasmids in all but one KP isolate. All KP ST15 isolates harbored different ESC resistance genes and different plasmids, and presented the non-transferable bld3512 gene, in association with blaCTX-M-15, blaCTX-M-2 (on IncR, or on IncN), bld2812a (on IncR) or bld2812b genes (on IncI1). KO isolates were positive for blaCTX-M-15 gene (on IncI2), or for the bld3512 and bld281H-1 genes (on IncL/M). They were all positive for qnr genes, and one also for the aac(6’)-Ib-cr gene. All Klebsiella isolates showed multiresistance towards aminoglycosides, sulfonamides, tetracyclines, trimethoprim and amphenicols, mediated by strA/B, aadA2, aadB, ant (2’)-Ia, aac(6’)-Ib, sul, tet, dfr and cat genes in various combinations. The emergence in pets of multidrug-resistant Klebsiella with ESBL, AmpC and PMQR determinants, poses further and serious challenges in companion animal therapy and raise concerns for possible bi-directional transmission between pets and human, especially at household level.


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Introduction
Klebsiella are bacterial pathogens that can cause a variety of severe infections in humans, mainly due to K. pneumoniae (KP) [1], [2] and to a lesser degree to K. oxytoca (KO) [3], [4]. KP is also a well-known causal agent of mastitis in cattle and Bacteremia in calves, cervicitis and metritis in mares, pneumonia and septicemia in foals, pneumonia, urinary tract infection (UTI) and septicemia in dogs [5], [6], [7].

Increasing antimicrobial resistance, especially towards aminoglycosides, (fluoro)quinolones, third and fourth generation cephalosporins, cephamycins, and carbapenems have been reported in the last decade [8], [9], [10], and poses serious therapeutic problems when treating Klebsiella infections in humans. In veterinary medicine, scarce information is reported on the occurrence of extended spectrum beta-lactamase (ESBLs), AmpC beta-lactamases and plasmid mediated quinolone resistance (PMQR) in Klebsiella isolates from companion animals [11], [12]. The aim of the study was to provide molecular characterization of extended-spectrum cephalexin resistance (ESC) resistance and PMQR in Klebsiella isolates from clinical cases or lesions in necropsied animals of canine and feline origin in Italy. A further aim was to determine phenotype and genotype of co-resistances, and to provide plasmid identification and genetic relatedness by Multi-locus Sequence Typing (MLST) and Pulsed Field Gel Electrophoresis (PFGE) among the isolates, to evaluate potential clustering of ESC, PMQR, and other resistance genes among clones.

Materials and Methods
Origin of ESC-resistant Klebsiella
Between 2006 and 2012, the Istituto Zooprofilattico Sperimentale delle Regioni Lazio e Toscana (IZSLT) investigated samples from 1555 dogs and 429 cats of clinical cases and necropsy specimens with suspicious bacterial infections, submitted by veterinarians practising mainly in central Italy, and some practising in northern Italy. Presumptive positive Klebsiella isolates were identified using the API 20E identification system (bioMérieux, Grapponne, France). For species-level identification of isolates...
with phenotypic inconclusive results 16S rDNA sequencing technique was employed, by means of the MicroSeq Full Gene system (Applied Biosystems, USA) as described previously [13].

Genotypic characterization

Multilocus Sequence Typing on KP isolates was performed as previously described [14], and interpreted according to the KP MLST database (www.pasteur.fr/mlst).

In addition, all isolates were genotyped by PFGE using XbaI according to the previously published protocol [15].

Antimicrobial susceptibility testing

Antimicrobial susceptibility testing was performed as minimum inhibitory concentrations (MIC) by micro-broth dilution in 96-well microtitre plates [Trek Diagnostic Systems, Westlake, OH, USA]. The following antimicrobials were tested: ampicillin, cefotaxime, cefazidime, ciprofloxacin, chloramphenicol, florfenicol, gentamicin, kanamycin, nalidixic acid, streptomycin, sulfonamides, tetracycline, and trimethoprim. The results were interpreted according to the European Committee on Antibiotic Susceptibility Testing (EUCAST) epidemiological cut-offs (www.eucast.org) and to Clinical Laboratory Standard Institute [16] or EUCAST clinical breakpoints for those drugs for which epidemiological cut-offs have not been made available (kanamycin, chloramphenicol, sulfamethoxazole, trimethoprim). For streptomycin, a cut-off of 16 mg/L was used, according to EUCAST MIC distributions.

Confirmatory test for the detection of ESBLs were performed on isolates resistant to cefotaxime or ceftazidime according to Clinical Laboratory Standard Institute (CLSI) recommendations [16].

Detection of genes encoding beta-lactamase and PMQR

For the confirmed ESBL-producing isolates, the encoding genes belonging to the beta-lactamase and PMQR families were further analyzed for the presence of blaCTX-M [17], blaoxI [18], blafem [19], blaOXA [20], blampC families [21], as well as for genes of the qnr family, qep-A, and aac(6)-Ib-cr encoding for PMQR [22], [23], [24], [25], [26]. The isolates were further screened by PCR for genes encoding carbapenemases [27]. Amplicons were sequenced by BigDye Terminator chemistry (Applied Biosystems, Foster City, CA, USA) and migrated with an automated sequencer (ABI Prism 310; Applied Biosystems). Sequence data analysis was performed using CLC DNA workbench software version 5.7.1 (CLC Bio, Aarhus, Denmark) and evaluated against the GenBank nucleotide databases.

Detection of plasmid replicons

Identification of plasmids was performed by PCR-based replicon typing as previously described [28], [29], [30], and using the PBRT kit (Diatheva, Fano, Italy).

Plasmid analysis

Plasmid DNA preparations were performed using the NucleoSpin Plasmid/Plasmid (NoLid) kit (Macherey-Nagel, Düren, Deutschland) and used to transform Max Efficiency DH5α Competent Cells (Invitrogen, Life Technologies, U.S.A.). In order to identify the plasmid carrying the ESBLs and AmpC genes, the selection of the transformants was performed on LB agar plates containing 100 mg/ml ampicillin.

Additionally, the isolates were tested according to the manufacturer’s instructions using an array hybridization kit for DNA-based detection of the most common resistance genes, and for the integrase gene (intI1) of class 1 integrons of Gram negative bacteria (Alere Technologies GmbH, Jena, Germany) and the results interpreted by the ArrayMate, Alere.

Results

Isolation rates

The samples (n = 1984; dogs and cats) yielded a total 70 (3.53%, 95% CI: 2.72%–4.34%) KP and 23 (1.16%, 95% CI: 0.69%–1.63%) KO among the isolates, respectively. Of these, 15 (21.4%) KP and four (17.4%) KO revealed resistance to ESC and were investigated in this study.

Genetic relatedness

The 15 KP isolates investigated by MLST were assigned to four different Sequence Types (ST): ST11 (n = 1), ST340 (n = 2), ST101 (n = 8), and ST15 (n = 4) (Figure 1). ST11 and its single-locus (tudB) variant (SLV) ST340 (3/15, 20%), both belonging to CC11, were detected in 2012. The separation of the isolates based on MLST corresponded well with PFGE results grouping the same isolates (Figure 1). A total of 11 different PFGE patterns were observed including two clusters of two and four indistinguishable isolates, respectively (Figure 1). The cluster of the two isolates both belonged to ST340 and was related (80% similarity) to a single isolate exhibiting a unique PFGE pattern and belonging to ST11. The other cluster of four indistinguishable isolates was highly related (99% to 80% similarity) to additional four isolates within the same PFGE group, all belonging to ST101 (Figure 1). No clustering was observed related to time, animal origin, nor infection, but some to the presence of resistance genes (Table 1). No MLST was assigned to the four KO isolates. However, the four isolates revealed three different PFGE patterns of which one was a cluster of two identical isolates (Figure 2). The three patterns seemed not to be related, indicating a similarity of 45% and 55% to the pattern of the two clustering isolates. Interestingly, the two isolates of the same PFGE pattern were both from dogs and isolated within the same year, but it could be the result of a random effect (Figure 2).

Antimicrobial susceptibility testing

All isolates showed microbiological resistance to third-genera
cephaplosporins, and also clinical resistance, either when the MIC results were interpreted according to clinical breakpoints set by CLSI [16] or by EUCAST (e.g. MIC cefotaxime ≥ 4mg/L), except for 9KP (MIC 1 mg/L). The phenotype for PMQR was evident (ciprofloxacin MIC 0.25mg/L, nalidixic acid 8 mg/L) in KO isolates only, because in all KP isolates it was masked by the presence of resistance genes (Table 1). No MLST was assigned to the four KO isolates. However, the four isolates revealed three different PFGE patterns of which one was a cluster of two identical isolates (Figure 2). The three patterns seemed not to be related, indicating a similarity of 45% and 55% to the pattern of the two clustering isolates. Interestingly, the two isolates of the same PFGE pattern were both from dogs and isolated within the same year, but it could be the result of a random effect (Figure 2).

Genes encoding ESBL-, AmpC-, and PMQR

All Klebsiella isolates investigated showed the presence of at least one ESBL or AmpC gene encoding ESC resistance. Additionally, 16 out of 19 isolates harbored a PMQR gene (qnr family or aac(6)-Ib-cr, single or in combination). The most frequent ESC gene harbored by KP isolates was blaoxI-M-TX-M-15 (n = 11, 58%), detected in all eight ST101, in one ST13 and in both two ST340 isolates, respectively. All four ST15 KP isolates carried the blaoxI-M-20 gene, single or in combination with the ESC resistance genes blaoxI-M-TX-M-15.
**Discussion**

The paper provides evidence of heterogeneity of determinants conferring ESC resistance in clinical *Klebsiella* isolates from dogs and cats in Italy. To the best of our knowledge, all the ESC-resistant *Klebsiella* investigated were from sporadic clinical cases, although two clusters of KO (n = 2) and KP ST101 (n = 4) showed 100% similarity. Apparently, isolates were epidemiologically unrelated, with the exception of 6KP and 1KP (both ST101), with a DNA restriction profile showing 80% similarity, isolated from two cases of necropsies requested by the same veterinary practitioner in 2010 and 2011, respectively.

We also document the novel finding of the co-existence of the ESBL *blaSHV*28 and the AmpC *blaCMY*2 gene in one KP ST15 isolate from a cat and the first report of *qnrS* and *qnrA* and *aac(6’)-Ib-cr-PMQR associated with *Klebsiella* infections in companion animals from Europe. *Klebsiella* with ESBL phenotype were described in dogs and cats from China [12], although it was mediated in those isolates by the presence of the CTX-M-9 and CTX-M-1 group beta-lactamases, while a CTX-M-15 positive KP clone was reported from hospital-acquired infections in pets from France [31]. In our study, the ESBL gene *blaCTXM15* accounts for the majority of CTX-M genes detected in KP, but it was harbored mainly by the predominant KP ST101 lineage.

Interestingly, all KP ST15 isolates showed the association of the *blaCTXM15*, *blaCMY2*, or *blaCMY9* gene, both associated with the PMQR gene *qnrA1*, with one the two also positive for the *aac(6’)-Ib-cr* gene. The ST101 KO isolates exhibited the *blaSHV12*, *blaTEM1*, *blaDHA1*, and *qnrB4* genes. All isolates under study were negative for the presence of carbapenemase genes.

**Plasmid analysis**

Plasmids detected from all the strains and classified by the PCR-based replicon typing method, are reported in Table 1.

In our study, the *blaCTXM15* gene was successfully transferred by transformation from several MLST prototypic strains, demonstrating its plasmid localization. All but one transformant strains were positive for plasmids belonging to the IncR incompatibility group, and in the 18KP transconjugant the *blaCTXM15* gene was co-transferred with the *blaDHA1* gene. Other ST15 KP strains also presented the IncR plasmid carrying different ESBL genes such as *blaSHV2*, and *blaCTXM1*. The 2KP ST15 strain harbored a IncI plasmid carrying the *blaCMY2*-gene, while an IncN plasmid carrying both the *blaCTXM1* and *qnrS1* gene was detected in the ST11 16KP strain.

The KO transformants carried the ESBL genes, *blaCTXM9* and *blaSHV12* in two incompatibility groups, IncHI2 and IncL/M, respectively (Table 1).
Table 1. Sequence Types, plasmid incompatibility groups, and antimicrobial resistance phenotypes and genotypes in ESC-resistant *Klebsiella pneumoniae* (KP) and *Klebsiella oxytoca* (KO).

<table>
<thead>
<tr>
<th>Sequence Type</th>
<th>Key</th>
<th>Antimicrobial Resistance profile</th>
<th>Plasmid</th>
<th>ESBL and AmpC genes</th>
<th>PMQR genes</th>
<th>Other resistance and Integron genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>2KP</td>
<td>AMP,CTX,CFT,NAL,CIP,STR,KAN,CLO,SULFA,TRI,TET</td>
<td>IncI1; IncR</td>
<td>blaCTX-M-15, blaARM-28</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; catA1; dfrA12; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>15</td>
<td>9KP</td>
<td>AMP,CTX,CFT,NAL,CIP,STR,KAN,CLO,SULFA,TRI,TET</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaARM-28, blaTEM-1</td>
<td>aac(6')-Ib-cr</td>
<td>strA/B; catA1; dfrA12; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>15</td>
<td>7KP</td>
<td>AMP,CTX,CFT,NAL,CIP,STR,KAN,CLO,SULFA,TRI</td>
<td>IncR</td>
<td>blaCTX-M-15, blaARM-28, blaTEM-1</td>
<td>aac(6')-Ib-cr</td>
<td>strA/B; dfrA14; sul1; intI1</td>
</tr>
<tr>
<td>15</td>
<td>13KP</td>
<td>AMP,CTX,CFT,NAL,CIP,STR,KAN,CLO,SULFA,TRI</td>
<td>IncN; IncFIA; IncFIB; IncR</td>
<td>blaCTX-M-15, blaSHV-28, blaTEM-1, blaOXA-1</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; strA; catA1; dfrA12; sul1; sul1</td>
</tr>
<tr>
<td>101</td>
<td>10KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,SULFA,TRI,TET</td>
<td>IncR; IncFIIk; IncFII</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA2; dfrA1; dfrA14; sul1; tet(D)</td>
</tr>
<tr>
<td>101</td>
<td>3KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA2; dfrA1; dfrA14; sul1; tet(D)</td>
</tr>
<tr>
<td>101</td>
<td>5KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA2; dfrA1; dfrA14; sul1; tet(D)</td>
</tr>
<tr>
<td>101</td>
<td>8KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA2; dfrA1; dfrA14; sul1; tet(D)</td>
</tr>
<tr>
<td>101</td>
<td>6KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncHII2; IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>101</td>
<td>4KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>101</td>
<td>1KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,TRI</td>
<td>IncHII2; IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-1, blaTEM-1, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>340</td>
<td>17KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,SULFA,TRI,TET</td>
<td>IncHII2; IncR; IncFIIk, IncFII</td>
<td>blaCTX-M-15, blaSHV-11, blaTEM-5, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>340</td>
<td>18KP</td>
<td>AMP,CTX,CFT,NAL,CIP,KAN,GEN,SULFA,TRI,TET</td>
<td>IncR; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-11, blaTEM-5, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>11</td>
<td>16KP</td>
<td>AMP,CTX,CFT,NAL,CIP,SULFA,TRI,TET</td>
<td>IncN; IncFIIk</td>
<td>blaCTX-M-15, blaSHV-11, blaTEM-5, blaOXA-1, blaOXA-2</td>
<td>aac(6')-Ib-cr</td>
<td>aadA2; dfrA21; dfrA17; sul1; tet(A); intI1</td>
</tr>
<tr>
<td>NA</td>
<td>3A KO</td>
<td>AMP,CTX,CFT,CIP,STR,KAN,GEN,SULFA,TRI,TET</td>
<td>IncHII2; IncLM</td>
<td>blaCTX-M-15, blaSHV-11, blaTEM-5, blaOXA-1, blaOXA-2</td>
<td>qnrS1</td>
<td>aadA2; dfrA2; dfrA1; sul1; tet(A)</td>
</tr>
<tr>
<td>NA</td>
<td>4A KO</td>
<td>AMP,CTX,CFT,CIP,STR,KAN,GEN,SULFA,TRI,TET</td>
<td>IncHII2; IncLM</td>
<td>blaCTX-M-15, blaSHV-11, blaTEM-5, blaOXA-1, blaOXA-2</td>
<td>qnrS1</td>
<td>aadA2; dfrA2; dfrA1; sul1; tet(A)</td>
</tr>
<tr>
<td>NA</td>
<td>1A KO</td>
<td>AMP,CTX,CIP,STR,SULFA,TRI,TET</td>
<td>IncHII2; IncP</td>
<td>blaCTX-M-15</td>
<td>qnrA1</td>
<td>aadA2; dfrA2; sul1</td>
</tr>
<tr>
<td>NA</td>
<td>6A KO</td>
<td>AMP,CTX,CIP,SULFA,TRI,TET</td>
<td>IncHII2</td>
<td>blaCTX-M-15</td>
<td>qnrA1; aac(3')-Ib-cr</td>
<td>aadA2; aadB; sul1</td>
</tr>
</tbody>
</table>

Legend:
NA: Not Applicable; AMP = Ampicillin; CFT = Ceftazidime; CIP = Ciprofloxacin; CLO = Chloramphenicol; CTX = Cefotaxime; GEN = Gentamicin; KAN = Kanamycin; NAL = Nalidixic Acid; STR = Streptomycin; SULFA = Sulfamethoxazole; TET = Tetracycline; TRI = Trimethoprim.

Note: When underscored, plasmids and their content of beta-lactamase and PMQR genes were detected in transformant strains.

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time in clinical cases of pets from Italy the clone KP ST11 and its SLV ST340, harboring ESC and qnrST-PMQR resistance. Among these CC11 isolates, the ST11 harbored IncN plasmid, which has been frequently involved in the transmission of the blaCTX-M-1 gene, a feature suggesting an animal reservoir for this ESBL, since this Inc plasmid types have been demonstrated to be highly prevalent in zoonotic enterobacterial pathogens [29]. The same animal origin reservoir is proposed for the IncI plasmids harboring the blaKPC-2 gene found in E. coli avian commensal strains [34]. It is noteworthy that ST11 and ST340 carried transferable ESBL resistance but not resistance to carbapenems. ST(CC)11 and ST15 and ST101 are among human epidemic clones, carrying both ESBLs and carbapenemases, which have been increasingly detected worldwide, in Europe and in Italy in the last years [36], [37], [38], [39], [40]. These infections are worrisome, since the antimicrobial treatment options for these multidrug-resistant strains are very limited. In Italy, during the last years the rapid emergence of the carbapenemase KPC-producing KP, belonging to the ST101, CC11, and predominantly to a single Sequence Type ST258, has become a serious problem in health-care settings [41], [42], [43]. As for CTX-M and SHV-12 ESBLs in Italy, a high occurrence in KP isolated from humans has been demonstrated, being the ST11, ST37, ST147 and ST273 the prevalent clones [44], [45], [46], [47]. In two KO isolates, the ESBL-encoding \( \text{bla}_{\text{SHV-12}} \) gene co-existed with the AmpC gene \( \text{bla}_{\text{CMY-2}} \) in accordance with phenotype of resistance to cefotaxime and cefoxitin observed in the ESBL phenotypic confirmatory test. These two isolates also carried the PMQR gene \( \text{qnrB1} \). In these two KO transformants the IncL/M plasmid harbored both the \( \text{bla}_{\text{SHV-12}} \) and the \( \text{bla}_{\text{CMY-2}} \) genes but not the \( \text{qnrB1} \) gene. To our knowledge, this feature has never been described before.

The other KO presented the \( \text{bla}_{\text{CTX-M-9}} \) gene, located in an IncHI2 plasmid as described worldwide but associated to a \( \text{qnrA1} \) gene, a feature previously described in Spain in E. coli and KP of human origin [48], [49] and in KO in clinical specimens from Japan [50]. Similarly to what has been observed in other human and canine KP isolates [34], [12], the association of \( \text{bla}_{\text{CTX-M}} \) genes with the the \( \text{aac(6’)-Ib-cr} \) encoding an aminoglycoside acetyl transferase determining PMQR, was demonstrated in all ST101 and ST340 isolates, but only in out of four ST15, but these PMQR genes were not located in the same plasmid in our strains. Conversely, PMQR encoded by different \( \text{qnr} \) genes of the \( \text{qnrC} \) or \( \text{qnrB} \) groups were observed in all the KO isolates studied (Table 1). In the case of the \( \text{qnrB} \) gene, the two KO isolates also harbored the ESBL \( \text{bla}_{\text{CTX-M-9}} \) gene, a feature reported previously in association with \( \text{bla}_{\text{CMY-2}} \) and IncHI2 plasmids in KO of human origin [51].

Multidrug-resistance in the ESC resistant and PMQR isolates studied is of further concern from a therapeutic perspective, for a possible impact on clinical outcome of affected animals. In many isolates, the demonstration of the integrase \( \text{intI1} \), accounts for the presence of resistance gene cassettes with \( \text{aai} \), \( \text{andA} \), \( \text{cat} \), \( \text{diff} \) genes, associated with Class I integrons, similarly to what has been described in KP of human origin [52], [53]. As for streptomycin resistance, MIC>16 mg/L correlated in 100% isolates with the presence of \( \text{strA/B} \) genes.

Fortunately, the absence of carbapenemases offers so far a better scenario for antimicrobial therapy in companion animals, although a possible circulation, within a short time, of these carbapenemase-producing epidemic strains, is of concern also in veterinary medicine.

In conclusion, monitoring and characterization of multidrug-resistant \( \text{Klebsiella} \) in companion animals by means of phenotypic and molecular methods proved to be useful for providing a picture of mechanisms of resistance that may further spread clonally or by horizontal gene transfer, at regional or even at international level. Sharing this kind of information appears essential for building awareness in companion animal therapy, also in view of preventing and controlling the spread of multidrug-resistant strains in veterinary hospital settings. Indeed, the bi-directional exchange between owners and pets of \( \text{Klebsiella} \) carrying resistance to critically important antimicrobials for human health, raise some concerns also for the possibility of a spill back to humans, especially at household level.

The emergence of the PMQR and, above all, the emergence of concurrent transferable cephamycin, oxymino-cephalosporin, and beta-lactamase inhibitor resistance in multidrug-resistant \( \text{Klebsiella} \) isolates in pets, a recent issue even in human therapy [54], may pose in the next future further and serious therapeutic challenges also in bacterial infections of companion animals.

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Author Contributions

Conceived and designed the experiments: AF AB RSH FF AGF. Performed the experiments: VD SL FF GC RL CAS AGF. Contributed reagents/materials/analysis tools: AB AF RSH. Wrote the paper: AF AB RSH AGF.

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