Antimicrobial agents are crucial for the treatment of many bacterial diseases in pigs, however, the massive use of critically important antibiotics such as colistin, fluoroquinolones and 3rd–4th-generation cephalosporins often selects for co-resistance. Based on a comprehensive characterization of 35 colistin-resistant Escherichia coli from swine enteric colibacillosis, belonging to prevalent Spanish lineages, the aims of the present study were to investigate the characteristics of E. coli clones successfully spread in swine and to assess the correlation of the in vitro results with in silico predictions from WGS data. The resistome analysis showed six different mcr variants: mcr-1.1; mcr-1.10; mcr-4.1; mcr-4.2; mcr-4.5; and mcr-5.1. Additionally, blaCTX-M-14, blaCTX-M-32 and blaSHV-12 genes were present in seven genomes. PlasmidFinder revealed that mcr-1.1 genes located mainly on IncHI2 and IncX4 types, and mcr-4 on ColE10-like plasmids. Twenty-eight genomes showed a gyrA S83L substitution, and 12 of those 28 harbored double-serine mutations gyrA S83L and parC S80I, correlating with in vitro quinolone-resistances. Notably, 16 of the 35 mcr-bearing genomes showed mutations in the PmrA (S39I) and PmrB (V161G) proteins. The summative presence of mechanisms, associated with high-level of resistance to quinolones/fluoroquinolones and colistin, could be conferring adaptive advantages to prevalent pig E. coli lineages, such as the ST10-A (CH11-24), as presumed for ST131. SerotypeFinder allowed the H-antigen identification of in vitro non-mobile (HNM) isolates, revealing that 15 of the 21 HNM E. coli analyzed were H39. Since the H39 is associated with the most prevalent O antigens worldwide within swine colibacillosis, such as O108 and O157, it would be probably playing a role in porcine colibacillosis to be considered as a valuable subunit antigen in the formulation of a broadly protective Enterotoxigenic E. coli (ETEC) vaccine. Our data show common features with other European countries in relation to a prevalent clonal group (CC10), serotypes (O108:H39, O138:H10, O139:H1, O141:H4), high plasmid content within the isolates and mcr location, which would support global alternatives to the use of antibiotics in pigs. Here, we report for first time a rare finding so far, which is the co-occurrence of double colistin-resistance mechanisms in a significant number of E. coli isolates.

Keywords: Escherichia coli, colistin, mcr, ESBL, fluoroquinolones, ST10, colibacillosis, swine
INTRODUCTION

Multidrug-resistant Enterobacteriaceae, such as Escherichia coli, represent a threat to both human and veterinary health. E. coli has a great capacity to accumulate resistance genes, mostly through horizontal gene transfer. The major problematic mechanisms correspond to the acquisition of genes coding for extended-spectrum beta-lactamas (ESBL), carbapenemases, 16S rRNA methylases, plasmid-mediated quinolone resistance (PMQR) and mcr genes conferring resistance to polymyxins (Poirel et al., 2018).

Colistin has been widely used in Spain for the control of neonatal and post-weaning diarrhoea (PWD) in pigs caused by certain E. coli pathotypes: Enterotoxigenic E. coli (ETEC), defined by the presence of genes encoding enterotoxins (eltA, and/or estA, and/or estB); atypical Enteropathogenic E. coli (aEPEC), carriers of eae but negative for bfpA (aEPEC); Shiga toxin–producing E. coli (STEC), positive for stx2a; STEC/ETEC, positive for both shiga toxin type 2e and enterotoxin-encoding genes (stx2a and estB and/or estA) (García-Meníño et al., 2018). PWD results in significant economic losses for the pig industry due to costs derived of treatment and handling, decreased weight gain, and mortality. These circumstances have promoted the use and abuse of antibiotics in intensive farming (Luppi, 2017; Rhouma et al., 2017). However, specific regulations have been set up in Europe due to the concern that colistin resistance could be transmitted from food-production animals to humans which makes necessary the investigation of sustainable alternatives to antimicrobials (EUROPEAN COMMISSION, 2018).

In Spain, the rates of antibiotic resistance in pig farming were recently analyzed in a collection of 499 E. coli isolates from 179 outbreaks of enteric colibacillosis occurred during a period of 10 years (2006–2016) (García et al., 2018; García-Meníño et al., 2018). The results revealed a prevalence of colistin-resistant E. coli implicated in PWD in Spanish farms as high as 76.9% within 186 ETEC, STEC and STEC/ETEC isolates. Besides, PCR and sequencing identified the presence of mcr-4 in 102 isolates, mcr-1 in 37 isolates and mcr-5 in five isolates. Interestingly, almost all mcr-4 isolates belonged to the clonal group ST10-A (CH11-24) (García et al., 2018), which was shown to be highly present (more than 50%) within the mcr-1 diarrheagenic isolates of a second study (García-Meníño et al., 2018). Both studies reinforced other countries’ findings that the pig industry is an important reservoir of colistin-resistant E. coli, as well as being carriers of other additional risk genes such as blaESBL genes (García et al., 2018; García-Meníño et al., 2018; Magistrali et al., 2018; Manageiro et al., 2019). Based on reported evidences (Beyrouthy et al., 2017; Gilrane et al., 2017), there is great concern about the in vivo acquisition of mcr- and blaESBL-bearing plasmids by human E. coli isolates following treatment with colistin, or via animal transmission through direct contact or via food chain. Particular attention is given to those named as high-risk clones of (ESBL)-producing bacteria, worldwide spread within humans and animals, including Escherichia coli sequence types ST10, ST131, ST405, and ST648 (Mathers et al., 2015; Sellera and Lincopan, 2019).

The aims of this study were (i) the characterization of resistances and plasmid profiles of successfully spread mcr-1, mcr-4, and mcr-5 E. coli in Spanish pig farming; (ii) the assessment of WGS-based approaches for the characterization of pathogenic E. coli, through the correlation of the in vitro results with in silico predictions using the bioinformatics tools of the Center for Genomic Epidemiology (CGE).

MATERIALS AND METHODS

E. coli Collection

Thirty-five swine E. coli, positive by PCR for mcr-genes, were fully sequenced. Specifically, the 35 E. coli were selected from 499 diarrheagenic isolates of different geographic areas of Spain (2006–2016) (García et al., 2018; García-Meníño et al., 2018), taking into account the results of prevalence and significant association observed between pathotypes, presence of mcr and certain serogroups. In brief, the serogroups O108, O138, O141, O149, O157 were found significantly associated with ETEC; serogroups O26, O49, O80, O111 with aEPEC; serogroups O138 and O141 with STEC/ETEC; serogroup O139 with STEC; and serogroups O2, O15, O26, O45, O111, O138, O141, O157 with mcr-positive isolates (García-Meníño et al., 2018). Therefore, the collection analyzed here included 27 ETEC isolates (of serogroups O7, O8, O15, O45, O108, O138, O141, O149, O157, O157, O158); four STEC (O2, O139); three STEC/ETEC (O138 and O141) and one aEPEC (O111). The 35 representative isolates were carriers of the three mcr-types (mcr-1, mcr-4, and mcr-5) detected so far in our E. coli collection of porcine origin. Conventional pheno- and geno-typing was performed to complete classical characterization of serotypes, phylogroups, pathotypes and resistance profiles.

Conventional Typing

The H antigen was established for motile isolates by serotyping using H1 to H56 antisera, while non-motile isolates (HNM) were analyzed by PCR to determine their flagellar genes as described elsewhere (García-Meníño et al., 2018). The phylogroup was assigned by means of the quadraplex PCR of Clermont et al. (2013). Antimicrobial susceptibility was determined by minimal inhibitory concentrations (MICs) using the MicroScan WalkAway®-automated system (Siemens Healthcare Diagnostics, Berkeley, CA, United States) according to the manufacturer’s instructions for: amikacin, ampicillin-sulbactam, aztreonam, cefepime, cefazidime, ciprofloxacin, colistin, fosfomycin, gentamicin, imipenem, levofloxacin, meropenem, minocycline, nitrofurantoin, piperacillin-tazobactam, tigecycline, and tobramycin. Additionally, resistance to ampicillin, amoxicillin-clavulanate, cefazolin, cefotaxime, cefoxitin, cefuroxime, chloramphenicol, doxycycline, nalidixic acid and trimethoprim-sulfamethoxazole was determined by disk (Becton Dickinson, Sparks, MD, United States) diffusion assays. All results were interpreted according to the CLSI break points (Clinical and Laboratory Standards Institute, 2019). Genetic identification of the ESBLs was performed by PCR using
the TEM, SHV, CTX-M-1 and CTX-M-9 group-specific primers followed by amplicon sequencing (García-Meniño et al., 2018).

Whole Genome Sequencing (WGS) and Sequence Analysis

The libraries for sequencing were prepared following the instructions provided by the TruSeq Illumina PCR-Free protocol. Mechanical DNA fragmentation was performed with Covaris E220, and the final quality of the libraries assessed with Fragment Analyzer (Std. Sens. NGS Fragment Analysis kit 1-6000 bp). Lastly, the libraries were sequenced in an Illumina HiSeq1500, obtaining 100–150 bp paired-end reads which were trimmed (Trim Galore 0.5.0) and filtered according to quality criteria (FastQC 0.11.7). The reconstruction of the genomes and plasmids in the genomes was carried out using the methodology PLAsmid Constellation NETwork (PLACNETw)1 (Lanza et al., 2014). The assembled contigs, with genomic size ranging between 4.9 and 5.9 Mbp (mean size 5.5 Mbp), were analyzed using the bioinformatics tools of the Center for Genomic Epidemiology (CGE)2 for the presence of antibiotic resistance (ResFinder V2.1), virulence genes (VirulenceFinder v1.5.3), plasmid replicon types (PlasmidFinder 1.3./PMLST 1.4.), and identification of clonotypes (CHTyper 1.0), sequence types (MLST 2.0) and serotypes (SerotypeFinder 2.0). All the CGE predictions were called applying a select threshold for identification and a minimum length of 95 and 80%, respectively. Phylogroups were predicted using the ClermonTyping tool at the iame-research center web3. The mcr gene location was determined using PlasmidFinder/ResFinder prediction, together with PLACNETw references, and automatic annotation with Prokka v1.13 (Seemann, 2014).

RESULTS AND DISCUSSION

The phenotypic and genotypic traits of the 35 mcr-positive E. coli of swine origin, as well as their resistome and mobiome are summarized in Table 1. ResFinder confirmed that all genomes were mcr carriers. Likewise, VirulenceFinder predicted the acquired virulence genes encoding for the enterotoxins (sta1, stb, icaA), for fimbriae (fecF, k88), verotoxin (stx2) and intimin (eae), correlating in all cases with the pathotype assignment previously determined by PCR (García et al., 2018; García-Meniño et al., 2018).

Serotype Identification

In most studies, there is lack of information on E. coli serotypes since serotyping is performed by very few laboratories worldwide, hindering epidemiological comparisons. Here, we not only proved that there is a very good correlation between serotyping and SerotypeFinder predictions, but also the advantage of in silico H-antigen identification for those non-mobile (HNM) isolates. It is of note that 15 of the 21 HNM isolates were predicted as H39 (Table 1), namely O108:H39, O157:H39 and O45:H39 (five genomes, each). Given that the H39 is associated with the most prevalent O antigens within swine colibacillosis, such as O108 and O157, as well as ONT (García-Meniño et al., 2018), it would be probably playing a role in porcine colibacillosis to be considered as a valuable subunit antigen in the formulation of a broadly protective ETEC vaccine (Roy et al., 2009). The remaining six HNM isolates showed different O:H combinations: O138:H14, ONT:H5, O8:H20, O50/O2:H32, and O182:H19. SerotypeFinder also allowed the O45-antigen determination of two non-typeable (ONT) isolates (LREC-141 and LREC-146) and O182 of LREC-172; while LREC-147, belonging to O157 serogroup (Table 1), was predicted as ONT, probably due to the limitation of the assembly based on Illumina short reads (100–150 bp paired-end reads here) (Wick et al., 2017).

Phylogroups, Sequence Types and Clonotypes

The phylogroups established for the 35 genomes were the common ones reported for porcine E. coli isolates (A, B1, D-E) (Shepard et al., 2012; Bosak et al., 2019). However, we found discrepancies in the assignation obtained with the quadruplex PCR of Clermont et al. (2013) in comparison with that predicted by ClermonTyping for seven isolates: phylogroup E by PCR, while phylogroup D in silico (Table 1).

MLST and CHTyper tools determined 12 different STs, but mostly belonging to CC10 (21 genomes) and clonotype CH11-24 (18 genomes) (Table 1). The predominance of CC10, and specifically ST10, is in accordance with published data on E. coli isolates of swine origin, independently of the pathogenicity or antibiotic-resistance/susceptibility status (Shepard et al., 2012; Kidsley et al., 2018; Magistrati et al., 2018).

Resistome, Plasmidome and Phenotypic Expression of Resistances

The resistome analysis revealed that 34 of the 35 genomes encoded mechanisms of antibiotic resistance for ≥three different antimicrobial categories (Table 1). Seven E. coli were carriers of blaKSLR, namely blaCTX-M-14 (four genomes), blaCTX-M-32 (one) and blaSHV-12 (two). Besides, six different mcr variants were identified within the 35 E. coli: mcr-1.1 (in 18 genomes, including two mcr-4.2 carriers); mcr-1.10 (one); mcr-4.1 (one); mcr-4.2 (13 genomes, including the two mcr-1.1 carriers); mcr-4.5 (two) and mcr-5.1 (two).

PlasmidFinder revealed a high plasmid diversity based on the identified replicons, with two to seven different plasmid types per genome (Table 1). Within this heterogeneity, mcr-1.1 genes were found mainly on plasmids of the IncH2 and IncX4 types (six and four of the 12 mcr-1.1 plasmid-located genes, respectively); however, mcr-1.1 was also found integrated in the chromosome of LREC-145, LREC-148, LREC-149 and LREC-164 genomes. The mcr-1.10 gene of LREC-151 was located on the chromosome, while mcr-4 and mcr-5 variants were on Col8282-like (mcr-4.1), ColE10-like (for all 13 mcr-4.2 and two mcr-4.5 carriers) and pKP13a-like (mcr-5.1) plasmids. Furthermore, we found that

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1https://castillo.dicom.unican.es/upload/
2https://cge.dtu.dk/services/
3http://clermontyping.iame-research.center/
| Code       | Year of Isolation^1 | Serotype^2 | Phylo Group^3 | CHType^4 (CC) | ST | Plasmid content | Inc group (pMLST)^6 | Acquired resistances (in black) and point mutations (in blue)^7 | mcr/location^8 | Virulence genes^9 | Phenotypic resistance profile^10 | Pathotype-associated VF^11 |
|------------|---------------------|------------|---------------|----------------|-----|----------------|---------------------|---------------------------------------------------------------|----------------|-----------------|----------------------------------|------------------------|
| LREC-144   | 2010                | O141:H4    | A             | 11-24          | 5786 (10) | IncF (F30:A-B:) | IncX1 IncHII2 (ST4) | aadA1, aph(3“)-ib, aph(6)-id; mdf(A); sul1; tet(A); dfrA1; mcr-1.1 | gyrA D87G | mcr-1.1/IncHI2 | sta1, stb, fedF, astA, fedA, iha, iroA, iss | NAL^*, SXT, MIN^*, DOX, FOF, CST |
| LREC-145   | 2014                | O50/O2:H32 | A             | 11-23          | 10 (10)   | IncF (F89:A-B6) | IncI1 (ST90) IncII pO111-like | blaTEM-1β; aadA1, aadA2, aadA24, aph(3“)-la; cmlA1; erm(B); mdf(A); sul3; tet(A); dfrA1; mcr-1.1 | gyrA S83L | mcr-1.1/chromosome | stx2. | iha |
| LREC-147   | 2008                | *ONT:H5    | B1            | 29-38          | 156 (156) | IncF (F110:A-B:42) | IncHI2 (ST4) | aadA1, aadA2, acc (6“)-ib3; catA1, catB3, cmlA1; acc(6“)-ib-cr; mdf(A); sul1, sul3; tet(B); dfrA1; mcr-1.1 | gyrA S83L, gyrA D87N, parC S80L, parC E84G | mcr-1.1/IncHI2 | stb, astA, iss, IptA, gad | TIC, AMP, SAM, AMC, TOB^*, NAL, CIP, LVX, SXT, MIN, DOX, CHL, CST |
| LREC-148   | 2013                | O157:H59   | A             | 11-24          | 10 (10)   | IncF (F12/08:A-B42) | IncB/O/K2 Col156-like | blaCTX-M-14, blaCTX-M-15; mdr(A); tet(B); mcr-1.1 | gyrA S83L; pmrB V161G | mcr-1.1/chromosome | lacA, stb, K88, astA, cba, celB, cma, gad, iha, sepA | TIC, AMP, SAM, AMC^*, CFZ, CXM, CTX, FEP, NAL^*, MIN^*, DOX, CST |
| LREC-149   | 2010                | O138:H10   | A             | 27-0           | 100 (165) | IncF (F110/108:A-B:42) | IncI1 (STunknown) IncII IncQ | blaTEM-1β; aadA1, aac(3)-Iia; mdr(A); tet(A); dfrA1; mcr-1.1 | gyrA S83L | mcr-1.1/chromosome | lacA, stb, K88, astA, capU, iha | LT, STb, K88 |
| LREC-164   | 2009                | O111:H9    | B1            | 4-24           | 29 (29)   | IncHI2 (STunknown) | IncX1 Col8282-like | aadA1, aadA2, aph(3“)-ib, aph(6)-id; catA1, catB3, cmlA1; mdf(A); sul1, sul3; dfrA1; tet(A); mcr-1.1 | gyrA S83L | mcr-1.1/chromosome | eae, espA, espB, espF, espJ, tcpP, tee, cil, efa1, astA, celB, iha, IptA, ileA, ileB | TIC, NAL, SXT, DOX^*, CHL, CST |
| LREC-165   | 2006                | O8:H20     | A             | 7-0            | 398 (398) | IncF (F2:A-B71) | IncHI2 (ST4) IncII | aadA1, aadA2; catA1, cmlA1; mdr(A); sul3; tet(A); mcr-1.1 | mcr-1.1/ND | | astA, capU, gad | DOX^*, CHL, CST |

(Continued)
| Code      | Year of Isolation | Serotype | Phylogroup | ST (CC) | Plasmid content | CHType | Acquired resistances (in black) and point mutations (in blue) | mcr/location | Virulence genes | Phenotypic resistance profile | Pathotype-associated VF |
|----------|-------------------|----------|------------|---------|----------------|--------|---------------------------------------------------------------|-------------|----------------|-----------------------------|--------------------------|
| LREC-166 | 2010              | O7:H4    | A          | 11-27   | 93             | IncF   | blaTEM-1,2, bilaTEM-12, apm(3’)Ll, apm(3’)Ib, apm(6)Id; catA1; mdf(A); sul3; mcr-1.1 | mcr-1.1/ ND | stb, astA, iss | TIC, AMP, SAM, CFZ, CXM, CTX, CAZ, ATM, CHL, CST | Stb                     |
| LREC-169 | 2015              | O141:H4  | A          | 11-24   | 7323 (10)     | IncF   | blaTEM-1,2, bilaTEM-12, apm(3’)Ll, apm(3’)Ib, apm(6)Id; catA1; mdf(A); sul3; mcr-1.1 | mcr-1.1/ ND | stb, astA, iss | TIC, AMP, GEN*, NAL, CIP, LVX, SXT, CHL, CST | Sta, F18                 |
| LREC-170 | 2015              | O139:H1  | *D         | 2-54    | 1              | IncF   | mdf(A); mcr-1.1 | mcr-1.1/ IncX4 | stb2, fedF, elia, fedA, gad, lpa | TIC, AMP, SXT, MIN*, DOX, CXT | VT2e, F18               |
| LREC-177 | 2008              | O138:H14 | *D         | 28-65   | 42             | IncF   | blaTEM-1,2, aadA1, apm(3’)Ib, apm(6)Id; mdf(A); sul1; tet(B); mcr-1.1 | mcr-1.1/ ND | stb1, astA, bca, cma, fedA, gad, lpa | TIC, AMP, SAM, AMC*, NAL, CIP, LVX, MIN, DOX, CXT | LT, Sta, F18             |
| LREC-172 | 2014              | O182:H10 | A          | 11-94   | 10 (10)       | IncF   | blaTEM-1,2, aadA1, apm(3’)Ib, apm(6)Id; mdf(A); sul1; tet(B); mcr-1.1 | mcr-1.1/ IncX4 | stb1, astA, bca, cma, fedA, gad, lpa | TIC, AMP, SAM, GEN, TOB*, ATM, NAL, CIP, LVX, SXT, DOX*, CHL, CST | Sta, F18               |
| LREC-174 | 2010              | O15:H45  | *D         | 4-331   | 118           | IncF   | blaTEM-1,2, aadA1, apm(3’)Ll, apm(6)Id; mdf(A); sul3; mcr-1.1 | mcr-1.1/ IncX4 | stb1, astA, bca, cma, fedA, gad, lpa | TIC, AMP, SAM, GEN, TOB*, ATM, NAL, MIN, DOX*, CHL, CST | Stb                     |
| LREC-175 | 2009              | O45:H45  | E          | 550-400 | 4247          | IncF   | mcr-1.1 | mcr-1.1/ IncX4 | stb1, astA, bca, cma, fedA, gad, lpa | TIC, AMP, SAM, GEN, TOB*, ATM, NAL, MIN, DOX*, CHL, CST | Stb                     |
| Code   | Year of Isolation | Serotype | Phylo Group | CHType | ST (CC) | Plasmid content | Acquired resistances (in black) and point mutations (in blue) | mcr/location | Virulence genes | Phenotypic resistance profile | Pathotype-associated VF |
|--------|-------------------|----------|-------------|--------|---------|-----------------|---------------------------------------------------------------|--------------|----------------|-------------------------------|-----------------------|
| LREC-178 | 2009 | O141:H4 | A | 11-24 | 10 | IncF (F30A::B-); IncH1-II (ST9*); IncI1 (STuncknown) IncX1 IncX4 | blbTEM-16::aaddA1, aaddA2; cmIA1; mdf(A); mphB; su1, su3; dfrA1; mcr-1.1; gyrA D87G; pmrB V161G | mcr-1.1/IncX4 | sta1, stb, stx2, fddF, cma, fedA, iha | TIC, AMP, GEN, TOB*, SXT, CHL, CST | Sta, Stb, VT2e, F18 |
| LREC-181 | 2010 | O139:H1 | *D | 2-54 | 1 | IncI1 (STuncknown) IncX1 | blbTEM-16::aaddA1, aaddA2; cmIA1; mdf(A); mphB; su1, su3; dfrA1; mcr-1.1; gyrA D87G; pmrB V161G | mcr-1.1/IncX4 | sta1, stb, stx2, fddF, cma, fedA, iha | TIC, AMP, GEN, TOB*, SXT, CHL, CST | Sta, Stb, VT2e, F18 |
| LREC-136 | 2012 | O149:H1 | A | 27-0 | 100 | IncF (F108:A::B54) IncI1 (STuncknown) IncR ColI8282-Iike | blbTEM-16::aaddA1, aaddA2; cmIA1; mdf(A); su3; dfrA1; mcr-1.1 | ColI8282 | sta1, stb, astA, capU, iha | TIC, AMP, SAM, AMC, NAL, DOX*, CHL, CST | LT, Sta, Stb, K88 |
| LREC-131 | 2011 | O108:H59 | A | 11-24 | 10 | IncF (F111:A::B42) IncI1 (ST3) IncII ColI56-Iike ColE10-Iike | blbTEM-16::aaddA2; aph(3′)-Ia; cmIA1; mdf(A); su3; tet(B); dfrA1; mcr-4.1; gyrA S83L | mcr-4.1/ColE10 | itcA, sta1, fddF, colE, fedA, iha | TIC, AMP, CFZ, CFX, CTX, CAZ, ATM, NAL, CIP, LVX, SXT, MIN*, DOX, CHL, CST | LT, Sta, F18 |
| LREC-132 | 2016 | O108:H59 | A | 11-24 | 10 | IncF (F111:A::B42) IncH1-II (ST9) IncI1 (ST48) IncII ColI56-Iike ColE10-Iike | blbTEM-16::aaddA1, aaddA2, aph(3′)-Ia, aph(3′)-Ib, aph(4)-Ia, catA1; mdf(A), mphB; su1; tet(B); dfrA1; mcr-4.2; gyrA S83L, gyrA D87G, parC S80I | mcr-4.2/ColE10 | itcA, sta1, stb, fddF, astA, cma, gad, iha | TIC, AMP, AMC, GEN, TOB, NAL, CIP, LVX, SXT, MIN*, DOX, CHL, CST | LT, Sta, Stb, F18 |
| LREC-133 | 2008 | O138:H14 | *D | 28-41 | 42 | IncF (F14:A::B-); IncX1 ColE10-Iike | mdrA; mcr-4.2; gyrA S83L, parC S80R | mcr-4.2/ColE10 | sta1, stb, stx2, fddF, air, elA, fedA, gad, iha, iss, lptA | NAL, CIP, SXT, MIN, DOX, CHL, CST | Sta, Stb, VT2e, F18 |
| LREC-134 | 2013 | O139:H1 | *D | 2-54 | 1 | IncI1 (STunknown) IncX1 ColE10-Iike | aac(3)-Ia, aph(3′)-Ia, aph(4)-Ia, catA1; mdrA, mphB; su1; tet(B); dfrA1; mcr-4.2; gyrA D87N | mcr-4.2/ColE10 | sta1, stb, stx2, fddF, elA, fedA, gad, lptA | GEN, TOB, CST | VT2e, F18 |
| LREC-137 | 2009 | O157:H59 | A | 11-24 | 10 | IncF (F2::A::B42) IncII ColI56-Iike ColE10-Iike | blbTEM-16::aaddA1; mdrA; su3; tet(B); dfrA1; mcr-4.2; gyrA S83L, pmrB V161G | mcr-4.2/ColE10 | itcA, stb, K88, astA, cba, cma, gad, iha, sepA | TIC, AMP, GEN, TOB, NAL, CIP*, SXT, MIN, DOX, CHL*, CST | LT, Stb, K88 |
| Code      | Year of Isolation | Serotype | Phylo Group | CHType | ST (CC) | Plasmid content | Acquired resistances (in black) and point mutations (in blue) | mcr/location | Virulence genes | Phenotypic resistance profile | Pathotype-associated VF |
|-----------|-------------------|----------|-------------|--------|---------|-----------------|---------------------------------------------------------------|--------------|-----------------|-------------------------------|------------------------|
| LREC-138  | 2008              | O157:H39 | A           | 11-24  | 10(10)  | IncF (F2/111:A-B42) IncI1 (ST154) IncI2 ColE10-like | aac(3)-Iv, aph(3’)-Ib, aph(4’)-la, aph(6’)-Id; floR; mdf(A); tet(B); *mcr-4.2* gyrA S83L, gyrA D87G, parC S80I | mcr-4.2/ColE10 | itcA, sta1, stb, astA, cba, cma, iva | NAL, CIP, LVX, MIN*, DOX, CHL, CST | LT, STa, STb, F18 |
| LREC-139  | 2009              | O108:H39 | A           | 11-24  | 10(10)  | IncF (F89*C2:A-B42) IncN (ST1) IncI2 Col156-like ColE10-like | aac(3)-Iv, aph(3’)-Ib, aph(4’)-la, catA1; erm(B); mdf(A); mph(B); suI1; tet(B); tet(M); dfrA1; *mcr-4.2* gyrA S83L, gyrA D87N, parC S80I | mcr-4.2/ColE10 | itcA, astA, celB, gad, iva | NAL, CIP, LVX, CST | LT |
| LREC-140  | 2015              | O108:H39 | A           | 11-24  | 10(10)  | IncF (F111:A-B42) IncHI2 (ST9) IncN (ST1) IncI2 ColE10-like | blaCTX-M-32, blaTEM-18; aac(3)-Iv, aadA1, aph(3’)-Ib, aph(4’)-la; catA1; emr(B); mdf(A); mph(B); suI1; tet(B); tet(M); dfrA1; *mcr-4.2* gyrA S83L, gyrA D87G, parC S80I | mcr-4.2/ColE10 | itcA, sta1, stb, fedF, astA, fedF, cma, fedA, iva | TIC, AMP, CFZ, OXM, CTX, CAZ, FEP, GEN*, TOB*, ATM, NAL, CIP, LVX, SXT, MIN, DOX, CHL, CST | LT, STa, STb, F18 |
| LREC-142  | 2010              | O45:H39  | A           | 11-24  | 10(10)  | IncF (F89*C2:A8*:B42) IncHI1 (ST2*) IncX1 Col156-like ColE10-like | blaCTX-M-32, blaTEM-18; aac(3)-Iv, aadA1, aph(3’)-Ib, aph(4’)-la, catA1; erm(B); mdf(A); mph(B); suI1, suI3; tet(B); dfrA1; *mcr-4.2* gyrA S83L, gyrA D87G, parC S80I | mcr-4.2/ColE10 | itcA, stb, *K88*, astA, gad, iva | TIC, AMP, GEN*, TOB, NAL, CIP, LVX, SXT, DOX*, CHL, NIT*, CST | LT, STb, K88 |
| LREC-143  | 2006              | O138:H14 | *D          | 28-41  | 42(4)   | IncF (F14*A8:B*) IncHI1 (ST2*) IncX1 ColE10-like | aac(3)-Iv, aph(3’)-la; mdf(A); catA1; suI1; tet(B); dfrA1; *mcr-4.2* gyrA S83L, parC S80R | mcr-4.2/ColE10 | stat1, stb, stx2, fedF, air, fedA, gad, iva, lctA, iss | NAL, CIP, SXT, MIN, DOX, CHL, CST | STa, STb, VT2e, F18 |
TABLE 1 | Continued

| Code   | Year of Isolation1 | Serotype² | Phyla Group³ | CHType⁴  | ST⁵ (CC) | Plasmid content Inc group (pMLST)⁶ | Acquired resistances (in black) and point mutations (in blue)⁷ | mcr/location⁸ | Virulence genes⁹ | Phenotypic resistance profile¹⁰ | Pathotype-associated VF¹¹ |
|--------|-------------------|-----------|--------------|----------|---------|-----------------------------------|---------------------------------------------------------------|---------------|-----------------|-------------------------------|-----------------------------|
| LREC-156 | 2011             | O108:H39  | A            | 11-24    | 10 (10) | IncF (F111:A8*:B42)             | blaCTX-M-14; blaTEM-1B; aac(3)-Iv, aadA1, aadA2, aph(3’)-lb, aph(4)-la, aph(6)-id; catA1; mdf(A); sul1; sul3; tet(B); dirA1; mcr-4.2 gyrA SB38, gyrA D87G, parC SB30; pmrB V161G | mcr-4.2/ ColE10 | itoA, sta1, fedF, astA, celB, fedA, iha | TIC, AMP, CFZ, OX, CTX, FEP, GEN*, TOB, NAL, CIP, LVX, SXT, MIN, DOX, CHL, CST | LT, Stb, F18                |
| LREC-135 | 2008             | O45:H39   | A            | 11-24    | 10 (10) | IncF (F89*:C2:A*:B42) IncI1 (ST202*) IncX1 ColE10-like | blaTEM-1A; aac(3)-Iv, aadA1, aadA2, aph(3’)-lb, aph(4)-la, aph(6)-id; mdf(A); mcr-4.5 gyrA SB38, gyrA D87G, parC SB30; parE L416F; pmrB V161G | mcr-4.5/ ColE10 | itoA, stb, K88, astA, iha, sepA | TIC, AMP, GEN*, TOB, NAL, CIP, LVX, CST | LT, Stb, K88                |
| LREC-146 | 2008             | O45:H39   | A            | 11-24    | 10 (10) | IncF (F89*:C2:A*:B42) IncI1 (ST202*) IncX1 ColE10-like | blaOX-1, blaTEM-1A; aac(3)-Iv, aadA1, aph(3’)-lb, aph(4)-la, aph(6)-id; floR; mdf(A); sul1; sul2; mcr-4.5 gyrA SB38, gyrA D87G, parC SB30; pmrB V161G | mcr-4.5/ ColE10 | itoA, stb, K88, astA, iha, sepA | TIC, AMP, SAM, AMC, GEN, TOB, NAL, CIP, LVX, SXT, CHL, CST | LT, Stb, K88                |
| LREC-152 | 2008             | O157:H39  | A            | 11-24    | 10 (10) | IncF (F108*:A*:B42) IncI1 (ST290*) IncX1 | blaTEM-1A; aac(3)-Iv, aph(3’)-la, aph(3’)-lb, aph(4)-la, aph(6)-id; mdf(A); mcr-5.1 gyrA SB38, pmrB V161G | mcr-5.1/ ND (plasmid reference pKP13a) | itoA, stb, K88, astA, gad, iha | TIC, AMP, SAM, GEN, TOB, NAL, CST | LT, Stb, K88                |
| Code     | Year of Isolation | Serotype | Phylogroup | ST (CC) | Plasmid content | Acquired resistances (in black) and point mutations (in blue) | mcrr/location | Virulence genes | Phenotypic resistance profile | Pathotype-associated VF |
|----------|-------------------|----------|------------|---------|-----------------|--------------------------------------------------------------|--------------|-----------------|----------------------------|------------------------|
| LREC-177 | 2007              | O157:H59 | A          | 11-24   | IncI2 CoI56-ike | aadA1, aph(3′)-Ia; mdf(A); tet(B); dfrA1; mcr-5.1 | mcr-S.1/ND   | itCA, sta1, stb, fedF | NAL, SX7, MIN, DOX, CST   | LT, StA, StB, F18   |
| LREC-141 | 2007              | O45:H59  | A          | 11-24   | IncH12 (ST4)   | blaTEM-1A, aac(3)-Ia, aadA1, aph(3′)-Ia, aph(6′)-Ib, tetA; mfr(A); sul1, sul2, suxA, tetA; dfrA1, mcr-1.1, mcr-4.2 | mcr-1.1/ND   | itCA, fedF      | TIC, AMP, SAM, AMC, GEN, TOB, NAL, OIP, LXF, SX, DOX, CST | LT, F18     |
| LREC-163 | 2011              | O157:H39 | A          | 1               | IncI1 CoI56-ike | mcr-1.1/ND, itCA, fedF, tetA, gtd, lha | mcr-1.1/ND   | itCA, fedF      | TIC, AMP, SAM, AMC, GEN, TOB, NAL, OIP, LXF, SX, DOX, CST | LT, F18     |

1. Year of isolation of the WGS isolates recovered from pig colibacillosis. 2. Serotypes, 3. clonotypes, 4. sequence types, 5. rep68/clampID STs, 6. acquired antimicrobial resistance genes and/or chromosomal mutations, 7. virulence genes were determined using SerotypeFinder 2.0, CHtyper 1.0, MLST 2.0, pMLST 2.0, ResFinder 3.1 and VirulenceFinder 2.0 online tools at the Center of Genomic Epidemiology (https://cge.cbs.dtu.dk/services/), respectively; while 8. phylogroups were predicted using the ClermontTyping tool at the Iame-research Center web (http://clermonttyping.iame-research.center). 9. Serotypes: underlined are those antigens that were non-typeable (ONT or HNM) by conventional serotyping but determined by SerotypeFinder. 10. LREC-147 was solved as O157 by conventional typing. 11. Phylogroups: “D” indicates that LREC-133, LREC-134, LREC-143, LREC-151, LREC-170, LREC-171, LREC-174 revealed discrepancies between the assignation obtained with the quadruplex PCR of Clermont et al. (2013) and the in silico assignation using ClermontTyping tool, showing phylogroup E by PCR, but phylogroup D in silico.
there was no \textit{mcr} plasmid co-occurrence in LREC-141 and LREC-163, but rather the \textit{mcr-1.1} and \textit{mcr-4.2} genes were located in independent plasmids (IncHII2 and CoIE1-like types, respectively). The \textit{mcr} location remained undetermined for four isolates.

Since the \textit{mcr-1} plasmid gene was first described (Liu et al., 2016), it has been identified in members of the Enterobacteriaceae family encoded in different plasmid types, including IncI2, IncX4, IncHI1, IncHI2, IncFI, IncFII, IncP, IncK (Sun et al., 2018). Different authors corroborate that large conjugative plasmids of types IncHII2, IncX4 and IncI2 would be the maximum responsible for the dissemination of the \textit{mcr-1} gene among \textit{E. coli} isolates from different sources and geographical locations (Doumith et al., 2016; Li et al., 2017; Managerio et al., 2019). To date, other \textit{mcr} genes (2-9) have been described (Carroll et al., 2019); among them, the \textit{mcr-4} and \textit{mcr-5} genes appear mostly encoded in small and non-conjugative CoE-like type plasmids (Sun et al., 2018). Here we found similar results, since \textit{mcr-1.1} genes were located mainly on IncHII2 and IncX4 types, and \textit{mcr-4} on CoIE1-like plasmids. It is of note that the \textit{mcr-5.1} gene, predicted in LREC-152 and LREC-177, was linked to a Kp13-like plasmid (CP003996.1), location previously described by Hammerl et al. (2018) for one \textit{mcr-5} isolate recovered from a fecal pig sample at farm. Chromosomally-encoded \textit{mcr-1} location remains rare, however, it was described soon after the discovery of this plasmid-borne gene (Falgenhauer et al., 2016; Veldman et al., 2016). Here, we determined chromosomal location in five genomes by means of PLACNETw, and according to the predictive annotation of the \textit{mcr-contigs}, the only common element flanking the \textit{mcr-1} was a putative ORF, \textit{pap2}, which is part of the Tn6360 and encodes a Pap2 superfamily protein. Thus, Pap2 was detected in LREC-145, LREC-148, LREC-149, and LREC-164, while the ISApI1 element typically associated with the initial mobilization of \textit{mcr-1}, was missing within the five contigs (Snesrud et al., 2018).

Overall, our findings are in accordance with those reported by Magistrati et al. (2018) on 13 \textit{mcr-positive} \textit{E. coli} isolated from swine colibacillosis in Belgium, Italy and Spain. Both studies show common features in relation to a prevalent clonal group (CC10), serotypes (O108:H39, O138:H10, O139:H1, O141:H4), and \textit{mcr-plasmid} element. The confirmation of these similarities are of interest for the global design of alternatives to antibiotics that would curb the dissemination of specific clones in the pig farming.

The \textit{in vitro} analysis of resistances showed that 30 of the 35 \textit{E. coli} were multidrug-resistant (MDR) according to Magiorakos et al. (2012) definition (Table 1). Phenotypic results corresponded broadly to those predicted by ResFinder (Supplementary Table S1) as detailed below.

The quinolones/fluoroquinolones (FQ), together with polymyxins and 3rd-4th-generation cephalosporins, all are included in Category B of restricted antimicrobials in the EMA categorization, considering that the risk to public health resulting from its veterinary use needs to be mitigated by specific restriction (EMA/CVMP/CHMP, 2019). Two major mechanisms are implicated in the resistance to FQ, namely, mutations in the genes for DNA gyrase and topoisomerase IV, and decreased intracellular drug accumulation. In addition, plasmid-mediated quinolone resistances also play a role but usually conferring low-level FQ resistance (van Duijkeren et al., 2018). Phenotypically, 17 of the 35 isolates showed resistance to both nalidixic acid and ciprofloxacin, and other eight resistance to nalidixic acid only (Supplementary Table S1). In the majority of cases, phenotypic results correlated with those predicted by ResFinder. Particularly, 28 of the 35 genomes carried the \textit{gyrA} S83L substitution, with 12 of those 28 showing double-serine mutations (\textit{gyrA} S83L and \textit{parC} S80I). An additional substitution (\textit{gyrA} D87N) was detected in two of the 12 \textit{gyrA} S83L/parC S80I genomes. Thus, nalidixic acid resistance \textit{in vitro} corresponded to one single substitution (\textit{gyrA} S83L), and FQ resistance to double or triple substitutions (\textit{gyrA} S83L/parC S80I/\textit{gyrA} D87N). Plasmid-mediated quinolone resistant genes \textit{acc(6’)-Ib-cr} and \textit{qnrS1} were also present together with chromosomal mutations in LREC-147 and LREC-169, respectively. Double-serine mutations in specific positions of the \textit{gyrA} and \textit{parC} genes have been reported as a dominant feature of MDR lineages within \textit{E. coli}, \textit{S. aureus} and \textit{K. pneumoniae}, with favorable fitness balance linked to high levels of resistance to FQ (Fuzzi et al., 2017). This finding, in 12 out of the 28 in \textit{silico} predicted FQ-resistant could be conferring adaptive advantages to certain widely spread pig pathogenic clonal groups of \textit{E. coli}, such as the ST10-A (CH11-24) (Garcia et al., 2018). This hypothesis is presently assumed for ST131 and other risk clones linked to high FQ-resistance (Johnson et al., 2015; Fuzzi et al., 2017).

On the other hand, colistin has been widely used for the control of enteric diseases, mainly in swine and poultry (Rhouma et al., 2016; Hammerl et al., 2018). Several mechanisms of resistance due to chromosomal mutations or acquired resistance genes have been described so far (Olaitan et al., 2014; Poirel et al., 2018). The 35 colistin-resistant \textit{E. coli} of this study showed MIC values > 4 mg/L. As detailed above, ResFinder confirmed that all the analyzed genomes were \textit{mcr-carriers}. In addition to the plasmid mechanism (\textit{mcr}) of resistance, polymyxin resistance in \textit{E. coli} can be due to genes encoding LPS-modifying enzymes, particularly to mutations in the two-component systems PmrAB and PhoPQ, or in the MgrB regulator. Quesada et al. (2015) detected two colistin-resistant \textit{E. coli} recovered in 2011 and 2013 from the stools of two pigs, which showed mutations in PmrB V161G and PmrA S39I, reporting the finding as a rare event. Subsequently, Delannoy et al. (2017) analyzed 90 strains of \textit{E. coli} isolated from diseased pigs: 81 were phenotypically resistant to colistin and 72 \textit{mcr-1} carriers (including two colistin-susceptible). Although different mutations were found in the amino acid sequences of the MgrB, PhoP, PhoQ, and PmrB proteins of eight isolates, only two of them were \textit{mcr-1} positive (but colistin-susceptible). Surprisingly, we found here the double mechanism of colistin resistance in 16 \textit{E. coli}, harboring \textit{mcr}-genes together with one amino acid substitution: PmrB V161G (14 genomes) or PmrA S39I (two genomes). In a recent study on Parisian inpatient fecal \textit{E. coli} (Bourrel et al., 2019), the authors found 12.5% of colistin-resistant \textit{E. coli} carriers among 1,217 patients; however, \textit{mcr-1} gene was identified in only seven of 153 isolates, while 72.6% harbored mutations in the PmrA and PmrB proteins. According to the authors, their findings
indicate two evolutionary paths leading to colistin resistance in human fecal \textit{E. coli}, one corresponding to a minority of plasmid-encoded \textit{mcr-1} isolates of animal origin, and a second corresponding to a vast majority of human isolates exhibiting chromosomally encoded mechanisms (Bourrel et al., 2019). Thus, and given the limited data regarding the co-occurrence of double resistance mechanism, it is of note that 16 of the 35 \textit{mcr}-bearing genomes of our study showed mutations in the \textit{PmrA} and \textit{PmrB} proteins. Furthermore, two \textit{E. coli} (LREC-141 and LREC-163) shown to be carriers of two different \textit{mcr}-bearing plasmids together with \textit{PmrB} \textit{V161G} mutation. An explanation for this rare finding is that these isolates would be reflecting a cumulative evolution to antibiotic pressure and, as a consequence, enhancing the transmission (vertical and horizontal) of colistin resistance. In any case, further investigation is needed to evaluate the implication of chromosomal mutations and \textit{mcr} co-occurrence regarding colistin resistance phenotype.

In this study, 22 out of the 25 isolates showing phenotypic resistance to beta-lactams (Supplementary Table S1), were positive in the analysis \textit{in silico} for the presence of \textit{bla}_{TEM-1} genes, alone (14 genomes), or in combination with other \textit{bla} genes (\textit{bla}_{CTX-M-14}, \textit{bla}_{SHV-12}, \textit{bla}_{CTX-M-32} and \textit{bla}_{OXA-1}); additionally, two genomes showed the presence of \textit{bla}_{CTX-M-14} and \textit{bla}_{SHV-12}, respectively. With the exception of LREC-147, LREC-164, and LREC-170, which were phenotypically resistant to narrow-spectrum beta-lactamases but negative for the presence of genes, a good correlation was observed between genes predicted and resistance observed \textit{in vitro}. It is of note that \textit{bla}_{TEM-135}, determined in LREC-156 by conventional typing, was not identified \textit{in silico}. Beta-lactams are the most widely used family in current clinical practice. Numerous genes in \textit{E. coli} confer resistance to this group, being some of them, such as \textit{bla}_{TEM-1} widespread in \textit{E. coli} from animals coding for narrow-spectrum beta-lactamases that can inactivate penicillins and aminopenicillins. However, genes encoding ESBLs/AmpCs have increasingly emerged in \textit{E. coli} from humans and animals, including food-producing animals (Cortes et al., 2010).

Thirty out of the 35 genomes showed high frequency of resistance genes to aminoglycosides, specifically encoding AAC(3)-II/IV and AAC(6)-Ib, which are the most frequently encountered acetyltransferases among \textit{E. coli} of human and animal origins. The subclass AAC(3)-II, which is characterized by resistance to gentamicin, netilmicin, tobramycin, sisomicin, 2’-N-ethylnetrilmicin, 6’-N-ethylnetrilmicin and dibekacin (Shaw et al., 1993), and AAC(6’) enzymes that specify resistance to several aminoglycosides and differ in their activity against amikacin and gentamicin C1 (Ramirez and Tolmasky, 2010) seemed to correlate with the phenotypic detection of resistance to gentamicin and/or tobramycin (12 of the 17 resistant isolates) (Supplementary Table S1). We also detected high prevalence of genes encoding nucleotidyltransferases (\textit{aadA}), which specify resistance to spectinomycin and streptomycin, alone or together with phosphotransferases (APHs) (Ramirez and Tolmasky, 2010), but they were not tested in the phenotypic antimicrobial susceptibility tests.

It is noteworthy that the 35 genomes of our study were carriers of \textit{mdf(A)}. Edgar and Bibi (1997) described that cells expressing MdfA from a multicopy plasmid are substantially more resistant to a diverse group of cationic or zwitterionic lipophilic compounds. Besides, the authors found that MdfA also confers resistance to chemically unrelated, clinically important antibiotics such as chloramphenicol, erythromycin, and certain aminoglycosides and fluoroquinolones. This capability could correlate with the \textit{in vitro} resistance observed for some isolates to tetracyclines and aminoglycosides, in absence of other specific genes. In our collection, of the 24 isolates showing phenotypic resistance to minocycline and, or doxycycline (Supplementary Table S1), 20 showed carriage of \textit{tet} genes: 12 \textit{tet}(B), six \textit{tet}(A), one \textit{tet}(A) + \textit{tet}(B) and one \textit{tet}(B) + \textit{tet}(M). However, two \textit{tet}(A) isolates were susceptible to those antibiotics (LREC-163, LREC-169). Additionally, \textit{tet} genes were not detected \textit{in silico} in four phenotypically resistant isolates. In general, \textit{tet(A)} and \textit{tet(B)} are the most prevalent tetracycline resistance genes in \textit{E. coli} of animal origin, and specifically in isolates from pigs (Tang et al., 2011; Holzel et al., 2012; Jurado-Rabadan et al., 2014).

Although the use of chloramphenicol was banned in the European Union in food-producing animals in 1994, fluorinated derivative florfenicol is allowed for the treatment of bacterial infection in these animals (Schwarz et al., 2004; OIE, 2019). In the present study, all 19 chloramphenicol-resistant isolates (Supplementary Table S1) correlated with the presence of genes \textit{catA1} (12 genomes), \textit{catB3} (one genomes), \textit{cmlA} (ten genomes) or \textit{floR} (three genomes) detected \textit{in silico}. Travis et al. (2006) showed that chloramphenicol resistant genes are frequently linked to other antiobioresistance genes. Thus, through transformation experiments conducted with \textit{E. coli} from pigs demonstrated that \textit{aadA} and \textit{sul1} were located with \textit{catA1} on a large ETEC plasmid, and plasmids carrying \textit{cmlA} also carried \textit{sul3} and \textit{aadA}. According to the authors, this linkage might partly explain the long-term persistence of chloramphenicol resistance in ETEC despite its withdrawal years ago. In our study, ResFinder also showed an association of genes \textit{cmlA}, \textit{sul3} and \textit{aadA} present in the same contig (7 of the 10 genomes positive for \textit{cmlA}), and \textit{cmlA/aadA} in all cases. Additionally, \textit{aadA} and \textit{sul1} were located with \textit{floR} in LREC-146.

In \textit{E. coli} from food-producing animals, sulfonamide resistance is mediated by \textit{sul} genes (\textit{sul1}, \textit{sul2}, \textit{sul3}), widely disseminated, and frequently found together with other antimicrobial resistance genes, while \textit{dfr} genes confer trimethoprim resistance in \textit{E. coli} and other gram-negative bacteria (van Duijkeren et al., 2018). Within our collection, 20 of the 35 isolates were \textit{in vitro} resistant to trimethoprim/sulfamethoxazole (Supplementary Table S1), and most of them correlated with the presence of \textit{sul + dfr} genes in their genomes, with the exception of LREC-133 and LREC-170 (negative for the \textit{in silico} detection of \textit{sul}, \textit{dfr} genes) and LREC-146 (in which only \textit{sul1} and \textit{sul2} genes were predicted). Besides, ResFinder showed that \textit{sul1} (present in 16 genomes), \textit{sul3} (14 genomes) and \textit{sul2} (three genomes) were located together with \textit{dfr}A, and other resistance genes, as mentioned previously.

The fosfomycin resistance showed \textit{in vitro} by two isolates of the study collection, was not predicted for LREC-144 and LREC-145 (Supplementary Table S1) by ResFinder, which
analyzes the presence of *fos* genes encoding for fosfomycin-modifying enzymes. The use of this antibiotic has been limited to the treatment of infections by Gram-positive and negative pathogens, included *E. coli*, mainly in pig and poultry farming (Poirot et al., 2018). However, phosphonic acid derivates such as fosfomycin, have been recently categorized by the EMA (EMA/CVMP/CHMP, 2019) as Category A (antimicrobial classes not currently authorized in veterinary medicine in EU).

**CONCLUSION**

Swine colibacillosis control has been traditionally managed through the extensive use of antibiotics. Our results are a reflection of the situation within the industrial pig farming, where global hygiene procedures and vaccinations are essential for improvement in antimicrobial stewardship. The summative presence of antibioresistances could be conferring adaptive advantages to prevalent pig *E. coli* lineages, such as the ST10-A (CH11-24). Based on the different replications identified by PlasmidFinder (up to seven), it is of note the high plasmid diversity found within these isolates; further research is needed to know mechanisms of maintenance and advantages conferred to them.

Here, we report for first time a rare finding so far, which is the co-occurrence of double colistin-resistance mechanisms (*mcr*-genes and chromosonal mutations in the PmrA and PmrB proteins) in a significant number of *E. coli* isolates. This fact could be increasing the risk of colistin resistance-acquisition by means of food transmission. Globally, we found a very good correlation between resistances determined *in vitro* and genes predicted using CGE tools, and the same observation applies to the *E. coli* pathotype determination.

**DATA AVAILABILITY STATEMENT**

The nucleotide sequence of the 35 LREC genomes have been deposited in the NCBI sequence databases with accession codes SAMN11523829 to SAMN11523863. These sequences are part of BioProject ID PRJNA540146.

**AUTHOR CONTRIBUTIONS**

IG-M, DD-J, and SF-S undertook the laboratory work. AM and JB conceived and designed the study. IG-M, DD-J, VG, MT, and AM performed the data analysis. IG-M, DD-J, VG, MT, JB, and AM drafted the manuscript. All authors provided critical input and approved the final version.

**FUNDING**

This study was supported by projects PI16/01477 from Plan Estatal de I+D+I 2013–2016, Instituto de Salud Carlos III (ISCIII), Subdirección General de Evaluación y Fomento de la Investigación, and FEDER; AGL2016-79343-R from the Agencia Estatal de Investigación (AEI, Spain) and FEDER; ED431C 2017/57 from the Consellería de Cultura, Educación e Ordenación Universitaria (Xunta de Galicia) and FEDER. IG-M and VG acknowledge the Consellería de Cultura, Educación e Ordenación Universitaria, Xunta de Galicia for their pre-doctoral and post-doctoral grants (Grant Numbers ED481A-2015/149 and ED481B-2018/018, respectively). SF-S acknowledges the FPU programme from the Secretaríat General de Universidades, Ministerio de Educación, Cultura y Deporte, Gobierno de España (Grant Number FPU15/02644).

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.02469/full#supplementary-material
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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