

1 **Chromosome Mediated Fluoroquinolone and**
2 **Extended Spectrum Beta-lactamase Resistant Genes**
3 **in *E. coli* of Poultry Origin in Ekiti State**

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5 **Oluyege, A. O. ¹ and Ojo, K. O. ^{1*}**

6 ¹Microbiology Department, Ekiti State University, Ado-Ekiti, Nigeria.

7 ^{1*}Microbiology Department, Ekiti State University, Ado-Ekiti, Nigeria.

8 **ABSTRACT**
9

Background: One health approach aimed at solving global health crisis links human, animal, and environment health. This inclusive strategy has contributed to antibiotic classification in both human and animal medicine.

Aims: The aims of this research work are to determine the phylogenetic relationship of *E. coli* isolated from poultry and waste sources. The presence of chromosome mediated fluoroquinolone and extended spectrum beta-lactamase resistant genes will also be detected in the isolates.

Study design: Original research paper.

Methodology: Data on farming attitudes of poultry farmers were collected using a questionnaire. *E. coli* was isolated from fresh poultry droppings and waste disposal sites using eosine methylene blue agar. The antibiogram profile of the isolates was determined using the modified Kirby Bauer disc diffusion method. Phenotypic expression of fluoroquinolone (*qnrS*) and beta-lactamase (*blaCMY*) resistant traits were further detected using Polymerase Chain Reaction. The 16S rRNA gene sequencing was carried out followed by sequence alignment of *E. coli* genes with those from GenBank sources to determine the molecular identity of the isolates. Spearman's correlation coefficient (r_s) was run to determine the relationship between antibiotic treatment and resistant profile of the isolates. The phylogenetic relationship of the isolates was determined using Bio edit and Mega 6 software.

Results: Organic poultry farming was practiced by small-scaled, peasant farmers who raised free range birds while antibiotics were widely used on farms that adopted intensive mode of farming. The percentage occurrence of *E. coli* from waste disposal sources was lesser than that from fresh poultry droppings. Highest percentage of antibiotic resistance to the fluoroquinolones was found while the carbapenemase recorded the lowest. Statistical analysis shows that antibiotic treatment in poultry and resistant profile of isolates to antibiotics are directly related, ($r_s(3) = 0.866, p = .333$). The percentage similarity of gene sequence with those from Gene Data Bank ($\geq 99.29\%$) validates the identity of the isolates as *E. coli*. About, 60 % of the sampled population had the *qnrS* gene with a band size of approximately 322 base pair. Besides, 40 % of the sampled isolates possessed the *blaCMY* gene with a band size of approximately 460 base pair. Both genes co-existed in the chromosome of 15 % of the sampled isolates sourced from poultry droppings and waste sources. Phylogenetic classification links the origin of isolates from waste disposal sources to poultry production sites. Besides, phylogenetic analysis shows that variant strains of multiple antibiotic resistant *E. coli* from poultry with antibiotic treatment were more diverse compared to those obtained from birds bred without antibiotics.

Conclusion: The *qnrS* and *blaCMY* genes found in multiple antibiotic resistant *E. coli* mediated resistance to critically important antibiotics and co-existed in variants strains of *E. coli* occupying different clusters in the phylogenetic analysis. Crude antibiotic treatment regimen in poultry may be responsible for the expression of these antibiotic resistant genes harbored in the chromosome of variant strains of *E. coli*.

10 Keywords: [Poultry, antibiotic resistant genes]

11 Abbreviations: [M. A. R. B. - multiple antibiotic resistant bacteria]

12 1. INTRODUCTION

13 The development of one health approach linking the health of humans, animals, and the
14 environment has contributed to antibiotic classification in both human and animal medicine.
15 Critically important antibiotics include the aminoglycosides, third and fourth generation
16 cephalosporins, tetracyclines, sulfonamides, fluoroquinolones, macrolides and penicillins.
17 Veterinary highly important antibiotics are rifampicin, first generation cephalosporin,
18 lincomycins, bacitracin, colistin, and first generation quinolones. Also, there are veterinary
19 important antibiotics, virginiamycin, avilamycin, fusidic acid, novobiocin. Some of the
20 veterinary critically-important antibiotics such as the fluoroquinolone and newer generation of
21 cephalosporins are also considered to be critically important for both humans and animal
22 health. These two classes of antibiotics are not recommended as prophylaxis or for first line
23 treatment in poultry [1]. There are suggestions that antibiotic usage in poultry is responsible
24 for selection and transmission of antibiotic resistant genes. Antibiotics at sub-lethal
25 concentration can promote genetic changes through different pathways involving various
26 stress responses [2]. Bacterial strains harboring genes encoding resistance may be
27 transferred to the microbiota of an animal host without causing an infection. However, these
28 antibiotic resistant bacteria can potentially spread and cause infection to other host [3, 4].
29 Antibiotic resistant genes are often acquired among bacterial population through
30 spontaneous mutation (vertical evolution) and horizontal gene transfer. Vertical evolution
31 involves transfer of changes in genetic composition from parents to their offspring. Horizontal
32 gene transfer entails movement of genetic materials to another organism of different
33 offspring [5].

34 Antibiotic resistant genes encode gene products that genetically confer on bacterial cells the
35 ability to grow when antibiotics are administered on them [6]. These antibiotic resistant
36 genes include *qnrA*, *qnrB*, *qnrS*, *AcrA* (fluoroquinolone resistant gene); *aac(3)-IV*, *aac(3)-II*,
37 *aac(6)-ib-cr*, *ant(3)-I*, *aph(3)-II*-aminoglycoside resistant genes; *blaSHV*, *blaCMY*, *blaOXA*,
38 *blaTEM*, *blaCTX*-beta-lactamase resistant genes; *ermA*, *ermB*, *ermC*, *msrA*-erythromycin
39 resistant genes; *sul-I*, *sul-II*-sulfonamide resistant genes; *tetA*, *tetB*, *tet C*-tetracycline
40 resistant genes; *dfrA1*-trimethoprim resistant gene; *aadA1*- streptomycin resistant gene;
41 *cmIA*, *floR*, *cat-I*-chloramphenicol resistant genes; *Mcr-I* polymyxin resistant gene [7, 8]. The
42 genes that code for antibiotic resistance in bacteria can be plasmid mediated or located on
43 chromosomes [9]. It was observed that some strains of *E. coli* have plasmids with an intact
44 promoter, and genes that encode for resistance to streptomycin β -lactamase, sulfonamides
45 and tetracycline. But not all these genes are expressed due to a chromosomal transcriptional
46 control that silenced the expression of plasmid genes. Also, plasmid borne cells can be
47 eliminated in a bacterial population [10]. Besides, natural plasmids are usually large,
48 maintained at low copy number and prone to loss [11]. Apparently, the bacterial
49 chromosome may be considered a more stable reservoir and natural repository of antibiotic
50 resistant genes than any other known source(s).

51 2. MATERIALS AND METHODS

52 Research tools

53 A questionnaire was administered to farm managers, data on antibiotic treatment and poultry
54 management practices were collected.

55 **Place and Duration of Study:** Microbiology Department, Ekiti State University, Ado-Ekiti,
56 from February 2017 to December 2019.

57 Study population and site

58 The study population comprises 12 poultry farms with their managers. Poultry birds such as
59 broilers, turkeys and free range birds were reared in the selected poultry farms in Ido and
60 Usi Ekiti. The study population comprises layers, broilers, turkeys and free range birds. A
61 total of 204 fecal droppings, 12 feed samples, 12 water samples and 12 samples comprising
62 soil and waste water from disposal sites were collected for culturing.

63 Collection of samples

64 Fresh fecal droppings from poultry birds were randomly sampled with a sterile swab stick
65 and transferred into a freshly procured, factory-packed, sealed polythene bag. Farm feed,
66 water and soil from disposal sites were also collected in sterile universal containers and
67 immediately transferred to the Microbiology Laboratory, Ekiti State University, Ado-Ekiti [12].
68 The samples were cultured within 2 hours of collection.

69 **Isolation techniques and biochemical characterization**

70 Swab sticks containing fecal droppings from poultry birds were suspended in 5 mL of sterile
71 saline water, prepared as 10 % suspension. The suspension was streaked on E.M.B agar
72 plates with sterile wire loop and incubated at 37 °C for 18-24 hours. Distinct colonies with
73 green metallic sheen and dark centers from the primary culture were preliminarily identified
74 as *E. coli* [13, 12]. One gram of the collected soil sample was weighed and added to the test
75 tube containing 9 mL of sterile distilled water. A volume of 1 mL of the stock solution was
76 serially diluted in series of test tubes containing 9 mL of sterile distilled water. About 0.1 mL
77 of the tenth-fold dilution from the eighth tube was inoculated into plates and pour plating was
78 carried out with eosine methylene blue agar. The plates were incubated at 37 °C for 24
79 hours. A loopful of diluents from test tubes with dilution factor of 10⁻⁵ and 10⁻⁶ were streaked
80 on EMB agar. The plates were incubated as previous. The identified distinct colonies were
81 sub-cultured on sterile eosin methylene blue agar in order to obtain a pure secondary
82 culture. They were preserved on nutrient agar slant for biochemical characterization [12].

83 **Antibiotic sensitivity testing**

84 Antibiotic susceptibility test was carried out using the modified Kirby Bauer disc diffusion
85 method. Antibiotic discs (Oxoid) comprising ciprofloxacin (5 µg), Tetracycline (30 µg),
86 ofloxacin (5 µg), Trimethoprim/sulfamethoxazole (1.25/23.75 µg), gentamycin (10 µg),
87 amoxicillin-clavulanic acid (20/10 µg), ceftaxidime (30 µg), meropenem (10 µg) and
88 ceftriazone (30 µg) were used. Mac Farland standard of 0.5 which gives an inoculum size of
89 1.5 × 10⁸ CFU/mL was used to standardize the density of the suspension. The zones of
90 inhibition were recorded and interpreted as susceptible, intermediate and resistant based on
91 procedures of Clinical and Laboratory Standard Institute, 2013 [14].

92 **DNA extraction and molecular identification of isolates**

93 The DNA samples of twenty selected isolates were sub-cultured in 3 mL of Luria Bertani
94 broth and incubated at 37°C for 18 hours. The bacterial DNA samples were extracted and
95 using the phenol chloroform extraction protocol [15]. The quality of the extracted DNA was
96 estimated in 1.5 % agarose gel electrophoresis and the DNA bands were visualised in UV
97 light imaging system. The DNA fragments were amplified by PCR for bacterial identification
98 using previously reported universal 16S rRNA gene primers 27 F 5'- AGAGTTTGATCM
99 TGGCTCAG-3'- and 1525R 5'-AAGGAGGTGATCCAGCC-3' [16]. PCR was carried out in a
100 GeneAmp 9700 PCR System Thermal cycler (Applied Biosystem Inc., USA). The PCR
101 cocktail contained 10 µL of 5x GoTaq colourless reaction, 3 µL of 25 mM MgCl₂, 1 µL of 10
102 mM of dNTPs mix, 1 µL of 10 pMol each 27F 5'- AGAGTTTGATCMTGGCTCAG- 3' and
103 1525R 5'-AAGGAGGTGATCCAGCC-3' primers and 0.3 units of Taq DNA polymerase made
104 up to 42 µL with sterile distilled water and 8 µL DNA template [17]. The PCR profile consists
105 of an initial denaturation temperature at 94 °C for 5 min; followed by a 30 cycles of 94 °C for
106 30 s, 50 °C for 60 s, 72 °C for 1 minute 30 seconds; and a final termination at 72 °C for 10
107 mins [16, 18]. The integrity of the amplified 1.5 Mb gene fragment was checked on a 1.5 %
108 agarose gel electrophoresis. It was visualized by ultraviolet trans-illumination, photographed
109 and the sizes of the PCR products were estimated [19, 15]. The amplified fragments were
110 sequenced using a Genetic Analyzer 3130 XL sequencer from Applied Biosystems using
111 manufacturers' manual while the sequencing kit used was that of Big Dye terminator.
112 Sequence editing and phylogenetic analysis was done using bio-edit and mega 6 software.
113 The bacterial sources of DNA were identified by matching with validated sequences of
114 highest maximum identity score from the GenBank database [20].

115

116 **Molecular detection of *qnrS* and *blaCMY-2* genes in strains of *E. coli* via PCR**

117 Molecular detection of *qnrS* coding gene in the isolates was by simple PCR on the extracted
118 DNA template using *qnrS* coding regions specific primers. Previously reported primers
119 targeting *qnrS* coding genes, *qnrS*- F: 5'-GCAAGTTCATTGAACAGGGT 3' and *qnrS*-R: 5'
120 TCTAAACCGTCGAGTTCGGCG 3' with an amplicon size of about 428 bp were used [21].
121 The reaction cocktail at a volume of 25 μ L based on the manufacturer's prescription used for
122 PCR included 12.5 μ L of 2X PCR Master mix with dNTPS (400 μ M) and $MgCl_2$ (3 mM) at
123 final concentration of 1X and pH of 8.5, 2.5 μ L each of forward and reverse primers (1.0 μ M),
124 5 μ L of DNA template (100 ng), nuclease-free water was added to make up the reaction
125 volume [17]. The PCR profile used for the amplification of *qnrS* coding genes include an
126 initial denaturation at 94 $^{\circ}C$ for 45 s followed by 35 cycles each of denaturation temperature
127 of 94 $^{\circ}C$ for 45 s, annealing at 53 $^{\circ}C$ for 45 s, extension (72 $^{\circ}C$ for 5 min) and final extension
128 at 72 $^{\circ}C$ for 5 min [22]. Similarly, molecular detection *blaCMY-2* coding gene in the isolates
129 was also by simple PCR on the extracted DNA template using *blaCMY-2* coding regions
130 specific primers. Previously reported primers targeting *blaCMY-2* coding genes were
131 obtained from Promega, U.S.A. The Primers' coding genes were *blaCMY-F5*-GGCGTGTTG
132 GGCGGCGATG-3' and *blaCMY-R* 5' CAGCGG AACCGTAATCCA 3' with an amplicon size
133 of about 364 bp [23]. The reaction cocktail of volume 25 μ L used for PCR was based on the
134 manufacturer's prescription. It comprises 12.5 μ L of 2X PCR Master mix with dNTPS (400
135 μ M) and $MgCl_2$ (3 mM) at final concentration of 1X and pH of 8.5, 2.5 μ L each of forward and
136 reverse primers (1.0 μ M), 5 μ L of DNA template (100 ng), nuclease-free water was added to
137 make up the reaction volume [17]. The PCR profile used for the amplification of *blaCMY-2*
138 coding genes includes an initial denaturation at 95 $^{\circ}C$ for 2 minutes. Subsequently, 35 cycles
139 each of denaturation temperature of 95 $^{\circ}C$ for 45 s, annealing at 48 $^{\circ}C$ for 45 s, extension at
140 72 $^{\circ}C$ for 1 mins, followed by final extension at 72 $^{\circ}C$ for 5 mins were observed [23]. The
141 PCR products were analyzed in 1.5 % agarose gel [17, 19].

142 **Statistical Analysis**

143 Statistical analysis was carried out using SPSS version 20.0 for the analysis of Spearman's
144 correlation coefficient, percentages and frequency.

145 **3. RESULTS AND DISCUSSION.**

146 The antibiotic treatment profile of *E. coli* examined from poultry birds in Ido-Ekiti is shown in
147 Table 1. Data from farm records show that eleven antibiotics comprising enrofloxacin,
148 erythromycin, tetracycline, gentamycin, streptomycin, neomycin, colistin, cotrimoxazole,
149 metronidazole, tylosine and chloramphenicol were selectively used on the farms. Based on
150 the different antibiotic classes administered on the farm, sulfonamides were administered
151 most (50 %), while nitroimidazole and amphenicol were the least administered (8 %).
152 Diaveridine, an anticoccidiostat was administered in many of the farms (50 %). Carbapenem,
153 penicillin and cephalosporin were not administered.

154 The level of occurrence of *E. coli* in the samples is shown in Table 2. *E. coli* was found in
155 both poultry droppings and waste disposal sources were 44.1 % and 33 % respectively.

156 The antibiotic percentage resistant profile of *E. coli* isolated is shown in Table 3. Multiple
157 antibiotic resistant *E. coli* were isolated from 4 poultry waste disposal sites comprising both
158 soil and water sources and poultry droppings across the 12 farms. The average resistance of
159 the isolates to ciprofloxacin was the highest (87 %), followed by ofloxacin (83 %) and
160 cotrimoxazole.

161 The Spearman correlation coefficient (r_s) of variables is shown in Table 4. There was a
162 strong, positive correlation between antibiotics used for treatment and resistant profile of the
163 isolates, which was statistically significant ($r_s(3) = 0.866, p = .333$). Besides, there was also a
164 strong positive correlation between antibiotics on sale from market survey and resistant
165 profile of the isolates, which was statistically significant ($r_s(3) = 0.500, p = .667$).

166 The sequence alignment of isolates with strains from Gene data bank is shown in Table 5.
167 The percentage similarity of gene sequence with those from Gene Data Bank falls within the
168 range of 99.29 % - 100 %. The DNA sequences show all the sampled isolates were strains
169 of *E. coli*. The DNA sequences obtained in this research were deposited in GenBank under

170 the accession numbers SUB5294851KENECF3MK606083, SUB5294851KENECJ11MK60
171 6084, SUB529 4851KENECG3MK606085, SUB5294851KENECI1MK606086, SUB5294851
172 KENECL6MK606087, SUB5294851KENECH4MK606088, SUB5294851KENECJ12MK6060
173 89, SUB5294851KENECI2MK606090, SUB5294851KENECL5MK606091, SUB5294851KE
174 NECI14M K606092, SUB5294851KENECA2MK606093, SUB5294851KENECD1MK606094,
175 SUB5294851KENECH8MK606095, SUB5294851KENECK7MK606096, SUB5294851KENE
176 CE3MK606097, SUB5294851KENECC4MK606098, SUB5294851KENECEB5MK606099, SU
177 B5294851KENECK19MK606100, SUB5294851KENECK5MK606101, and SUB5294851KE
178 NECG9MK606102. The nucleotide sequences of the isolates were also documented.

179 The agarose gel electrophoresis of PCR product of amplified *qnrS* resistant genes from *E.*
180 *coli* isolates is shown in plate 1. Results from Polymerase Chain Reaction and agarose gel
181 electrophoresis of the PCR products show that *E. coli* in lane 1, 2, 4, 6, 7, 8, 10, 14, 17, 18,
182 19 and 20 comprising 60 % of the sampled population had the *qnrS* (fluoroquinolone
183 resistant) gene with a band size of approximately 322 bp on the chromosome.

184 Also, the agarose gel electrophoresis of the PCR products of extended spectrum beta
185 lactamase gene *blaCMY-2* amplified from the DNA of the isolates is represented in plate 2.
186 The results from molecular analysis show that *E. coli* in lanes 3, 7, 10, 13, 14, 16, 17 and 19
187 comprising 40 % of the sampled population possessed *blaCMY-2* (the extended spectrum
188 beta-lactamase gene) with a band size of approximately 460 bp on the chromosome. The
189 *blaCMY-2* gene was present in the two *E. coli* isolates from free range birds obtained from
190 the same farm without a record of antibiotic consumption. The gene was present in two
191 isolates obtained from layers. They were from two different farms. The gene was also
192 present in one isolate each from cockerel and broiler. Also, two isolates from different waste
193 disposal sites harbored the *blaCMY-2* gene. Besides, both the *qnrS* and *blaCMY-2* genes
194 co-existed in the genome of 15 % of the isolates from free range birds, layers, and soil
195 (disposal sites).

196 The phylogenetic analysis of the sequenced bacterial genes is represented in figure 1. The
197 phylogenetic tree constructed was a well rooted tree with *Lactobacillus plantarum*
198 NR_042254.1 used to create an out group (negative control) while *E. coli* MG913260.1 and
199 *Escherichia fergusonii* NR_027549.1 from Gene Data Bank were used as positive controls at
200 species and genus levels respectively. Isolate I2 out-grouped itself from the rest of the
201 samples. The constructed tree has 2 major clades. Isolates B5, C4, H4, I1, G3, L6, F3, A2,
202 and I14 formed a sub-cluster of the major clade. Also, isolates J12, H8, K5, K7, D1, E3, J11,
203 K19, L5, G9, I14 and A2 formed the second sub-cluster of the major clades. In addition,
204 isolates from free range birds where antibiotics were not administered were restricted to a
205 sub-cluster in one of the major clades while isolates from birds with record of antibiotic
206 consumption and those from disposal sites were distributed across the two major clades.

207 The findings of this research show that antibiotics are widely used in the business of poultry
208 management. The practice of organic farming that limits antibiotic use in poultry production
209 was only adopted among small-scaled, peasant poultry farmer(s) with free range birds.
210 Wide- spread use of antibiotics in poultry management is common with large scale poultry
211 management. Periodic use of antibiotics on the farm may be attributed to compromised bio-
212 security measures and failure of routine immunization programs on poultry farms. Besides,
213 antibiotics were used majorly to check high mortality rate, disease outbreak and declining
214 egg production. Knowledge based resources on organic farming may not be publicized in the
215 area of study due to prior belief on antibiotics as an antidote to infection control in poultry
216 management. The current level of antibiotic use in poultry management can be reduced by
217 raising highly disease resistant birds and implementing viable vaccination schedules.

218 *E. coli* was isolated from poultry droppings obtained from different birds. Fecal samples from
219 free range birds, without antibiotic treatment recorded the highest presence of *E. coli*. Also,
220 the presence of the bacteria in soil and waste water samples obtained from disposal sites is
221 an indication of bio-pollution. Antibiotic resistant bacteria constitute a bio-pollutant and the
222 environment is a reservoir of these hazardous agents. The level of occurrence of the

223 bacteria varies in different farms. Comparatively, *E. coli* was more in fresh poultry droppings
224 than in soil and water sourced from the waste disposal sites. Abiotic conditions in the
225 environment may limit the survival rate of the bacteria. However, the observed rates of
226 bacterial presence contrast that of [24] who recorded an isolation rate of 83 % in fecal
227 droppings from poultry sources. The highest level of *E. coli* in samples from free range birds
228 shows that broad antibiotic use on the sampled farm might be responsible for the observed
229 low isolation rates.

230 The presence of multiple antibiotic resistant *E. coli* in environmental (soil/water) samples
231 from disposal sites shows that the environment can be a secondary reservoir of multiple
232 antibiotic resistant bacteria. Also, there was spread of multiple antibiotic resistant profiles on
233 the farms than resistance to single antibiotics. This finding is consistent with the assertions
234 of [25] that antibiotic therapy eliminates susceptible bacterial population and selects resistant
235 strains that grow through a Darwinian selection process. The resistant variants multiply and
236 become the predominant bacterial population. Their resistant genes are successively
237 transmitted to their progenies during reproduction. Infection with multiple antibiotic resistant
238 bacteria that evade treatment may pose a threat to humans and health institutions if the
239 sources of spread are not checked

240 Highest percentage of antibiotic resistance to the fluoroquinolones was observed. This
241 observation might be associated with wide-use of enrofloxacin on farms. In contrast,
242 resistance to meropenem was the lowest and this may be due to non-use of the
243 carbapenemase class of antibiotics for antibiotic therapy in poultry. This result contrasts that
244 of [26] that the percentage resistance of *E. coli* from human origin to carbapenemase in most
245 European countries was less than 1 %, except Belarous with a higher range of 10 % - 25 %
246 [27]. Majority of the isolates were resistant to multiple classes of antibiotics. The resistance
247 shown to antibiotics such as meropenem, amoxycillin-clavulanate and the cephalosporins
248 that were not administered on the farms may be due to the use of an analogue to these
249 antibiotics for therapy. However, meropenem may be an antibiotic of choice presumably
250 because of proactive antibiotic formulation policy limiting its usage for poultry management.

251 The DNA sequences show that all the sampled isolates were strains of *E. coli*.

252 The statistical analysis suggests a direct relationship between antibiotic treatment on the
253 farm and bacterial resistance to antibiotics. This shows a direct proportional relationship
254 between the two variables. Both data on antibiotic treatment and that from market survey are
255 positively related to antibiotic resistant profile of isolates. However, data on antibiotic
256 treatment on the farm are more related to resistance of bacteria than those obtained from
257 market survey. This result contrasts the findings of [28] who recorded a negative Spearman
258 correlation coefficient, $r_s(8) = -0.243$ between antibiotic treatment and antibiotic resistant
259 profile of isolates. This may be attributed to the differences in sample size of the antibiotics
260 used for both treatment and antibiotic sensitivity test.

261 Results from molecular analysis assert that poultry droppings were the major reservoir of the
262 *qnrS* gene while waste water and soil samples from disposal sites are secondary reservoir.
263 These research findings agree with the assertions of [29] that the frequency of occurrence of
264 *qnrS* gene among poultry and their production sites in Nigeria is high. The *qnrS* gene
265 originated from the chromosome of an organism occupying a human, animal or
266 environmental reservoir [30]. The high level of occurrence of *qnrS* genes in *E. coli* from
267 poultry could be associated with the selection pressure constituted by frequent
268 administration of fluoroquinolone (enrofloxacin) on the farms under study. Resistance to
269 fluoroquinolones in enterobacteriaceae can be chromosomal or plasmid mediated.
270 Chromosome-mediated resistance majorly occurs due to accumulation of mutations primarily
271 in DNA gyrase (GyrA) then in topoisomerase IV [9]. In addition, quinolone resistance can be
272 associated with an over expression of efflux pump systems [31]. High presence of
273 fluoroquinolone resistant gene may constitute a threat to both human and animal health
274 because fluoroquinolones are effective antibiotics in health care delivery.

275 The presence of *bla*CMY-2 genes in chromosomes of isolates from free range birds and
276 waste sources shows that polluted soil and water are secondary reservoir of antibiotic
277 resistant genes. This is consistent with the works of [32] who reported that *bla*CMY-2 genes
278 may be of chromosomal origin. However, resistance to β -lactams in Enterobacteriaceae is
279 majorly due to the production of β -lactamases and it may be encoded either chromosomally
280 or on plasmids [33]. The presence of this gene without the administration of cephamycin (a
281 beta-lactam antibiotic) might be attributed to selection pressure imposed by other
282 antimicrobial substances with similar or broader mode of action. Though farm records
283 asserted that cephamycin, a beta-lactam antibiotic was not administered; a penicillin based
284 formulation was present in the market for treatment of poultry birds and could have been
285 administered. The expression of the *bla*CMY gene is traceable to the use of an analogue of
286 penicillin, a beta-lactam antibiotic likely used for antibiotic treatment on the farm.
287 Free range birds (animals) have the potential of acquiring antibiotic resistant bacteria from
288 the environment via the ecological food chain. Nomadic life style of free range birds
289 promotes acquisition, release and spread of acquired antibiotic resistant bacteria in
290 droppings to the environment. The fluoroquinolone resistant gene (*qnrS*) is more
291 predominant than the beta lactamase resistant gene (*bla*CMY) in sampled isolates from the
292 area of study though both genes co-existed on the bacterial chromosome.
293 Phylogenetic analysis shows that variant strains of *E. coli* from poultry with record of
294 antibiotic treatment and disposal sites were more diverse in their antibiotic consumption
295 profile and clade distribution compared to those from birds raised without antibiotics. The
296 evolution, divergence and spread of chromosomal DNA from certain strains of *E. coli* across
297 the clades is traceable to various responses of the DNA to selective pressure imposed by
298 antibiotic treatment. This phenomenon attributes antibiotic therapy in poultry as the major
299 source of these variant strains of multiple antibiotic resistant *E. coli*. Related strains could
300 have emerged and spread via a common genetic mechanism. Wrong antibiotic policies and
301 crude use of antibiotics in poultry can lead to emergence of phylogenetically diverse strains
302 of bacteria with potential to evade treatment both in human and veterinary medicine.
303 Polluted soil, water, and waste disposal sites are potential secondary reservoir of emergent
304 strains of multiple antibiotic resistant *E. coli*. Phylogenetically diverse strains of bacteria have
305 the potential to complicate treatment both in poultry and humans during infection.
306 Phylogenetic classification also links the origin of multiple antibiotic resistant *E. coli* from
307 waste disposal sites to poultry production sites. Similarly, it links the origin of chromosomal
308 antibiotic resistant genes in isolates from free range birds to their interaction with the
309 environment. This asserts the contribution of antibiotic treatment in poultry, improper waste
310 disposal and free range system of animal farming to increased presence of multiple antibiotic
311 resistant bacteria in the environment.
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Table 1: Antibiotic treatment profile of poultry birds examined

| Characteristics | | Frequency (%) |
|---|------------------------|---------------|
| Exposure to antibiotics | Yes | 11 (91.7) |
| | No | 1(8.3) |
| Purpose of usage | Disease prevention | 3 (27.3) |
| | Treatment | 7 (63.6) |
| | Enhance egg production | 1 (9.1) |
| | None | 1 (9.1) |
| Method of administration | Water | 11 (100) |
| | Feed | 0 (0) |
| | Injection | 0 (0) |
| Age of birds when first exposed | 2 weeks | 5 (45.5) |
| | 1 month | 3 (27.3) |
| | 2 months | 2 (18.2) |
| | 3 months plus | 1 (9.1) |
| Routine chart for antibiotic use | Yes | 3 (25) |
| | No | 9 (75) |
| Frequency of administration of antibiotics | Cotrimoxazole | 6 (50) |
| | Tetracycline | 5 (41.7) |
| | Gentamycin | 3 (25) |
| | Enrofloxacin | 2 (16.7) |
| | Erythromycin | 2 (16.7) |
| | Streptomycin | 2 (16.7) |
| | Neomycin | 2 (16.7) |
| | Colistin | 2 (16.7) |
| | Metronidazole | 1 (8.3) |
| | Chloramphenicol | 1 (8.3) |
| | Tylosine | 1 (8.3) |
| Frequency of administration of classes of antibiotics | Sulphonamide | 6 (50) |
| | Tetracycline | 5 (41.7) |
| | Aminoglycoside | 5 (41.7) |
| | Fluoroquinolone | 2 (16.7) |
| | Macrolide | 2 (25) |
| | Polymyxin | 2 (16.7) |
| | Nitroimidazole | 1 (8.3) |
| | Amphenicol | 1 (8.3) |
| Product availability from market survey | Tetracycline | 2 (100) |
| | Cotrimoxazole | 1 (50) |
| | Neomycin | 2 (100) |
| | Gentamycin | 2 (100) |
| | Erythromycin | 2 (100) |
| | Penicillin | 1 (50) |
| | Enrofloxacin | 1 (50) |
| | Streptomycin | 1 (50) |
| | Colistin | 1 (50) |
| | Chloramphenicol | 1 (50) |
| | Tylosine | 1 (50) |

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* Numbers in parenthesis are percentage values

Table 2: Isolation rate of *E. coli* from poultry in Ido-Ekiti

| Poultry site | No of Poultry droppings examined | No positive (%) | Poultry feed examined | No positive (%) | Poultry water examined | No positive (%) | Waste sources/samples examined | | | Total samples examined |
|--------------|----------------------------------|-----------------|-----------------------|-----------------|------------------------|-----------------|--------------------------------|-----------------------|------------------|------------------------|
| | | | | | | | Waste water samples examined | Soil samples examined | No. positive (%) | |
| A (Pullet) | 17 | 5 (29.4) | 1 | - | 1 | - | - | 1 | - | 20 |
| B(Layer) | 17 | 3 (17.7) | 1 | - | 1 | - | - | 1 | - | 20 |
| C(Broiler) | 17 | 3 (17.7) | 1 | - | 1 | - | - | 1 | - | 20 |
| D(Broiler) | 17 | 5 (29.4) | 1 | - | 1 | - | - | 1 | - | 20 |
| E(Turkey) | 17 | 6 (35.3) | 1 | - | 1 | - | - | 1 | - | 20 |
| F(Turkey) | 17 | 5 (29.4) | 1 | - | 1 | - | - | 1 | - | 20 |
| G(Layer) | 17 | 8 (47.1) | 1 | - | 1 | - | - | 1 | 1 | 20 |
| H(Layer) | 17 | 11 (64.7) | 1 | - | 1 | - | - | 1 | - | 20 |
| I(Layer) | 17 | 13 (76.5) | 1 | - | 1 | - | - | 1 | 1 | 20 |
| J(Cockerel) | 17 | 12 (70.6) | 1 | - | 1 | - | - | 1 | - | 20 |
| K(Local) | 17 | 14 (82.4) | 1 | - | 1 | - | 1 | - | 1 | 20 |
| L(Layer) | 17 | 5 (29.4) | 1 | - | 1 | - | - | 1 | 1 | 20 |
| Total (%) | 204 | 90 (44.1) | 12 | - | 12 | - | 1 | 11 | 4 (33.3) | 240 (100) |

* Numbers in parenthesis are percentage value

Table 3: Antibiotics percentage resistant profile of *E. coli* isolated

| Source code | N | Cephalosporin | | | Fluoroquinolone | | Aminoglycoside | Caberpenem | Sulfonamide | Tetracycline | n (%) |
|--------------|----|---------------|-----------|-----------|-----------------|-----------|----------------|------------|-------------|--------------|-----------|
| | | CRO (%) | CAZ (%) | AMC (%) | CIP (%) | OFX (%) | CN (%) | MEM (%) | SXT (%) | TET (%) | |
| A | 5 | 0 | 0 | 5 (100) | 4 (80) | 3 (60) | 4 (80) | 0 | 3 (60) | 4 (80) | 4 (80) |
| B | 3 | 1 (33.3) | 1 (33.3) | 3 (100) | 1 (33.3) | 1 (33.3) | 1 (33.3) | 0 | 1 (33.3) | 3 (100) | 3 (100) |
| C | 3 | 1 (33.3) | 1 (33.3) | 3 (100) | 1 (33.3) | 2 (66.7) | 3 (100) | 0 | 2 (66.7) | 2 (66.7) | 3 (100) |
| D | 5 | 1 (20) | 1 (20) | 5 (100) | 4 (80) | 5 (100) | 5 (100) | 0 | 2 (40) | 3 (60) | 5 (100) |
| E | 6 | 0 | 0 | 5 (83.3) | 5 (83.3) | 5 (83.3) | 3 (50) | 2 (33.3) | 4 (66.7) | 5 (83.3) | 6 (100) |
| F | 5 | 1 (20) | 1 (20) | 5 (100) | 5 (100) | 5 (100) | 1 (20) | 3 (60) | 4 (80) | 4 (80) | 5 (100) |
| G | 8 | 5 (62.5) | 5 (62.5) | 2 (25) | 8 (100) | 6 (75) | 3 (37.5) | 3 (37.5) | 6 (75) | 7 (87.5) | 8 (100) |
| H | 11 | 9 (81.9) | 8 (72.7) | 7 (63.6) | 11 (100) | 9 (81.9) | 8 (72.7) | 1 (8.7) | 11 (100) | 11 (100) | 11 (100) |
| I | 13 | 13 (100) | 12 (92.3) | 9 (69.2) | 11 (84.6) | 12 (92.3) | 13 (100) | 9 (69.2) | 10 (76.9) | 10 (76.9) | 12 (92) |
| J | 12 | 11 (91.7) | 10 (83.3) | 7 (58.3) | 12 (100) | 10 (83.3) | 7 (58.3) | 7 (58.3) | 10 (83.3) | 11 (91.7) | 12 (100) |
| K | 14 | 9 (64.3) | 10 (71.4) | 10 (71.4) | 12 (85.7) | 12 (85.7) | 12 (85.7) | 7 (50) | 11 (78.6) | 10 (71.4) | 12 (86) |
| L | 5 | 5 (100) | 4 (80) | 4 (80) | 5 (100) | 5 (100) | 4 (80) | 4 (80) | 5 (100) | 1 (20) | 5 (100) |
| F/W | - | - | - | - | - | - | - | - | - | - | - |
| DS | 4 | 4 (100) | 3 (75) | 3 (75) | 3 (75) | 3 (75) | 4 (100) | 2 (50) | 4 (100) | 2 (50) | 4 (100) |
| Total | 94 | 60 (63.8) | 56 (60) | 68 (72.3) | 82 (87.2) | 78 (83) | 68 (72.3) | 38 (40.4) | 73 (77.7) | 73 (77.7) | 90 (95.7) |

Keys: n=number of isolates, N- Number of isolates showing multiple antibiotic resistance, F/W- isolates from feed and water, DS-number of isolates from disposal site, OFX-Ofloxacin; CIP-Ciprofloxacin, GN-Gentamycin; AMC-Amoxicillin-clavulanate, CRO-Ceftriaxone; MEM-Meropenem, CAZ= Ceftaxidime TET= Tetracycline, SXT= Trimethoprim/Sulfamethoxazole. Source A- pullets, B- layers, C- broilers, D- broilers, E-,turkeys, F-,turkey, G-layers, H-layers, I-layers, J- cockerels, K-local birds, L-layer

Table 4: Spearman's statistical correlation of antibiotic consumption and resistant profile

| Antibiotics | Frequency of use (%) | r_s | Resistant profile (%) | r_s | Market survey (%) |
|-------------------------|----------------------|--------------|-----------------------|--------------|-------------------|
| Cotrimoxazole | 6 (50) | | 73 (77.7) | | 1 (100) |
| Tetracycline | 5 (41.7) | | 73 (77.7) | | 2 (100) |
| Gentamycin | 3 (25) | | 68 (72.3) | | 2 (100) |
| Enrofloxacin | 2 (16.7) | | NT | | 2 (100) |
| Erythromycin | 2 (16.7) | | NT | | 2 (100) |
| Streptomycin | 2 (16.7) | | NT | | 1 (50) |
| Neomycin | 2 (16.7) | | NT | | 2 (100) |
| Colistin | 2 (16.7) | | NT | | 1 (50) |
| Metronidazole | 1 (8.3) | $r_s (.866)$ | NT | $r_s (.500)$ | NS |
| Chloramphenicol | 1 (8.3) | | NT | | 1 (50) |
| Ceftriazone | NA | | 60 (63.8) | | NS |
| Ceftazidime | NA | | 56 (60) | | NS |
| Amoxicillin-clavulanate | NA | | 68 (72.3) | | NS |
| Ciprofloxacin | NA | | 82 (87.2) | | NS |
| Meropenem | NA | | 38 (40.4) | | NS |
| Ofloxacin | NA | | 78 (83) | | NS |
| Penicillin | NA | | NT | | 1 (50) |
| Tylosine | 1 (8.3) | | NT | | 1 (50) |

Keys: NA – not administered, NS not sold, NT- not tested, r_s - Spearman's correlation coefficient

Table 5: Sequence alignment of isolates with strains from Gene data bank

| Isolate code | Accession number | Matched <i>E. coli</i> strain from Gene data bank | Percentage identity (%) |
|--------------|------------------------------|---|-------------------------|
| F 3 | SUB5294851 KENECF3MK606083 | MH67149.1 | 99.71 |
| J 11 | SUB5294851 KENECJ11MK606084 | CP042934.2 | 99.86 |
| G 3 | SUB5294851 KENECG3MK606085 | CP044314.1 | 99.66 |
| I 1 | SUB5294851 KENECI1MK606086 | MK606086.1 | 100 |
| L 6 | SUB5294851 KENECL6 MK606087 | KM198100.1 | 99.79 |
| H 4 | SUB5294851 KENECH4MK606088 | JQ781559.1 | 100 |
| J 12 | SUB5294851 KENECJ12 MK606089 | EU420950.1 | 99.89 |
| I 2 | SUB5294851 KENECI2 MK606090 | MH656755.1 | 99.69 |
| L 5 | SUB5294851 KENECL5 MK606091 | KP789331.1 | 99.65 |
| I 14 | SUB5294851 KENECI14MK606092 | KJ477001.1 | 99.44 |
| A 2 | SUB5294851 KENECA2 MK606093 | KJ477001.1 | 100 |
| D 1 | SUB5294851 KENECD1 MK606094 | MG602206.1 | 99.65 |
| H 8 | SUB5294851 KENECH8 MK606095 | CP026641.1 | 99.42 |
| K 7 | SUB5294851 KENECK7 MK606096 | KY780353.1 | 99.29 |
| E 3 | SUB5294851 KENECE3 MK606097 | KY655103.1 | 100 |
| C 4 | SUB5294851 KENECC4 MK606098 | CP040269.1 | 99.79 |
| B 5 | SUB5294851 KENECB5 MK606099 | CP0462591 | 99.45 |
| K 19 | SUB5294851 KENECK19MK606100 | CP044315.1 | 99.65 |
| K 5 | SUB5294851 KENECK5 MK606101 | KU870317.1 | 99.86 |
| G 9 | SUB5294851 KENECG9 MK606102 | MG602205.1 | 99.58 |

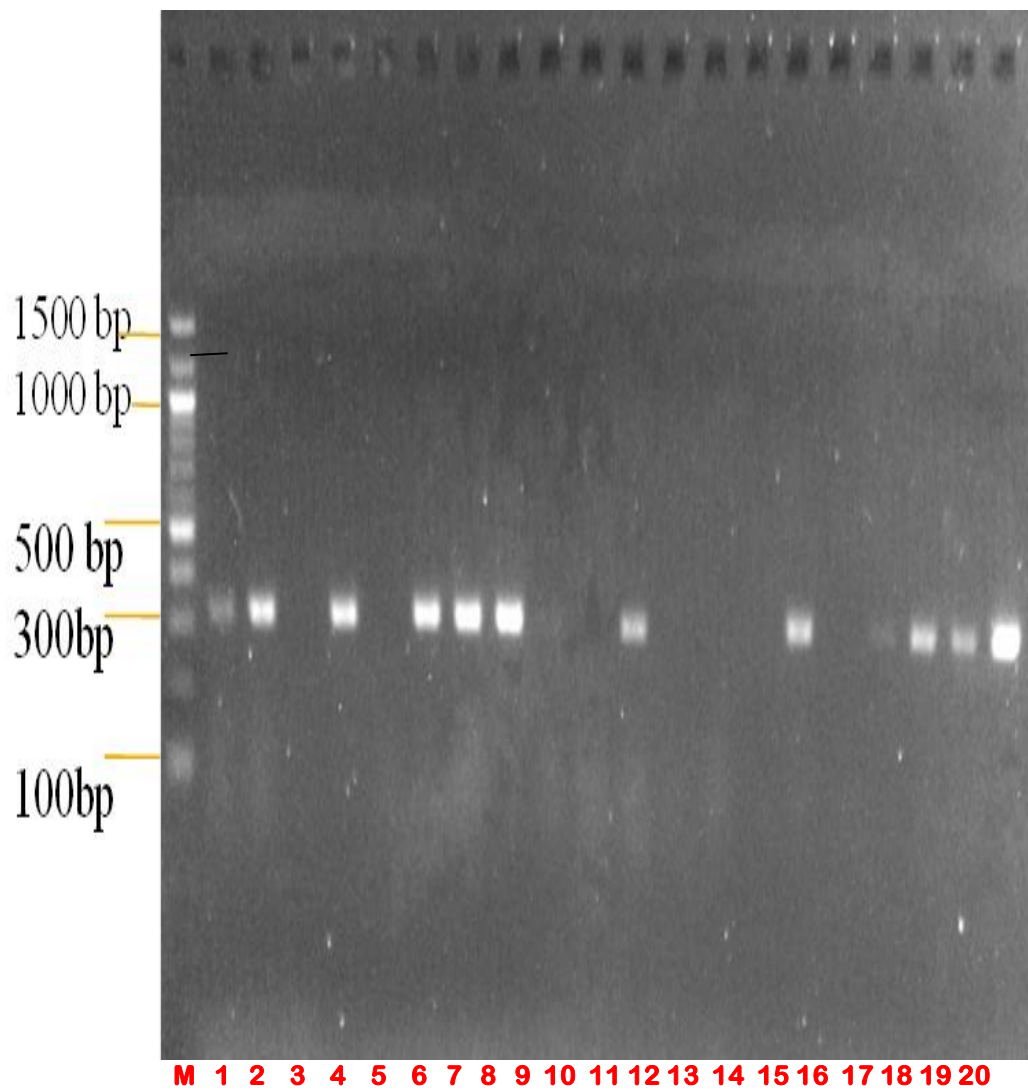


Plate 1: Agarose gel electrophoresis of the PCR products of *qnrS* resistant genes amplified from *E. coli* isolates (Band size approximately 322 bp).

Key: Lane M= Molecular Marker (100-1,500 bp), Lane 1= A2, Lane 2= B5, Lane 3= C4, Lane 4= D1, Lane 5= E3, Lane 6= F3, Lane 7= G3, Lane 8= G9, Lane 9= H4, Lane 10=H8. Lane 11= I1, Lane 12 = I2, Lane 13 = I14, Lane 14 = J11, Lane 15 = J12, Lane 16 =K5, Lane 17 = K7, Lane 18 = K19, Lane 19 = L5, Lane 20 = L6

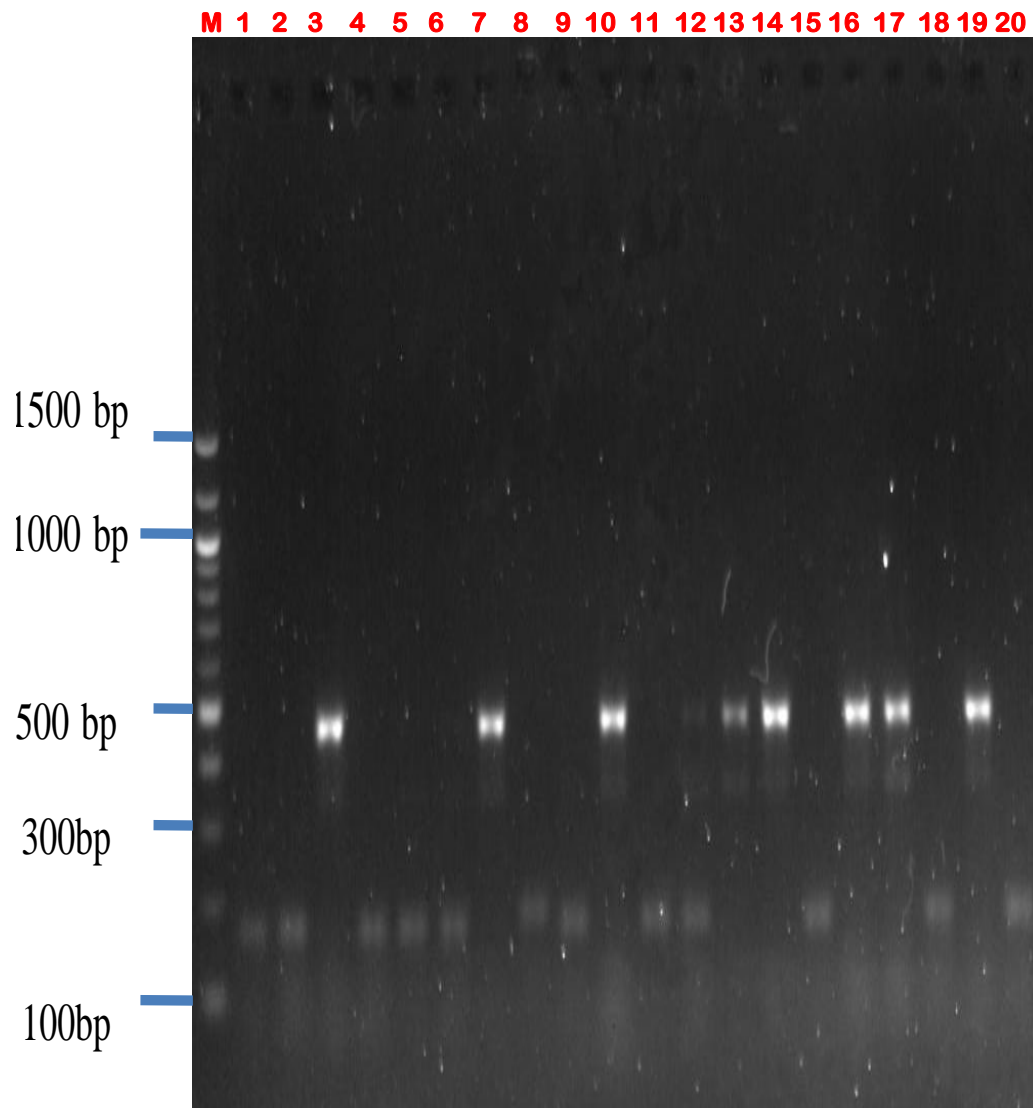
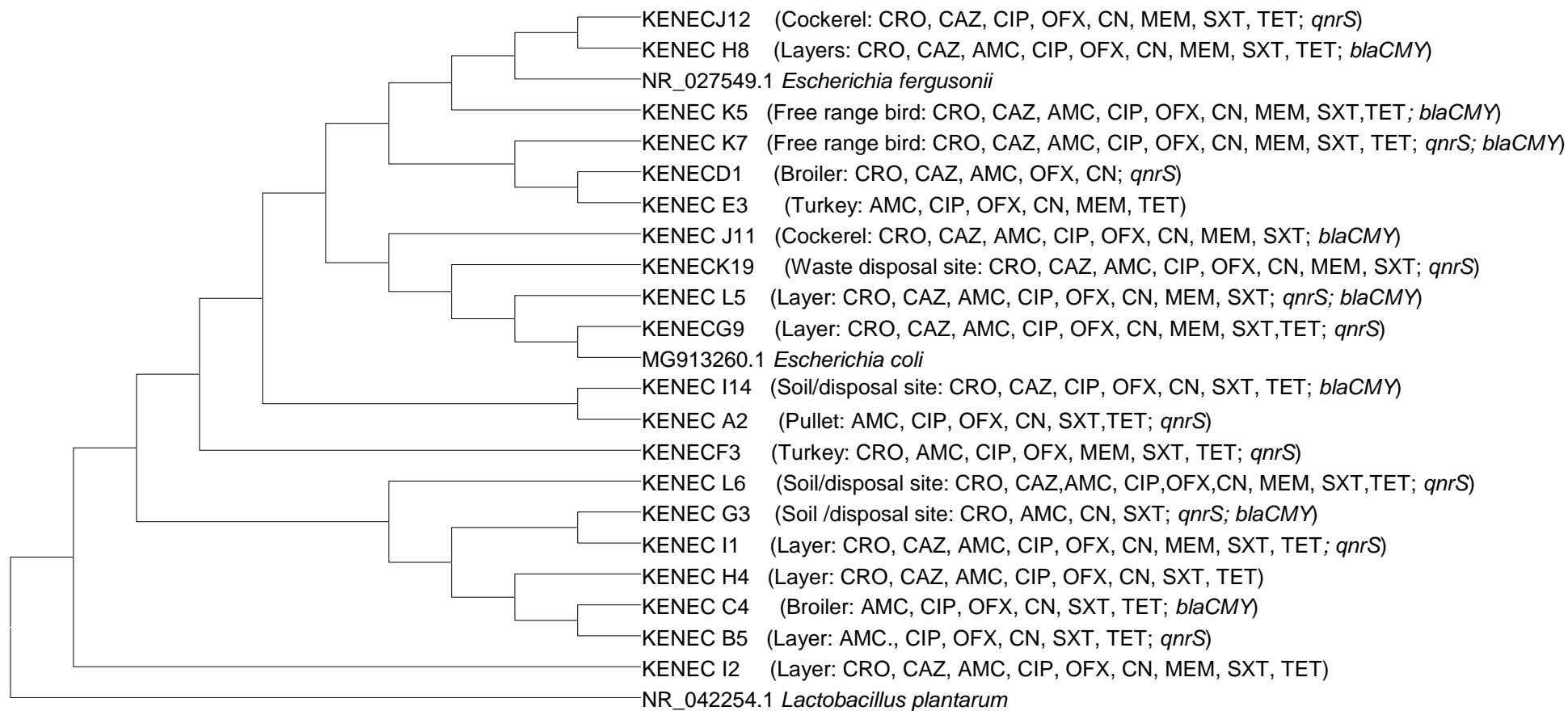


Plate 2: Agarose gel electrophoresis of the PCR products of the *bla*CMY-2 resistant gene amplified from *E. coli* isolates. (Band size approximately 460 bp)

Key: Lane M= Molecular Marker (100 -1,500 bp), Lane 1= A2, Lane 2= B5, Lane 3 = C4, Lane 4= D1, Lane 5 = E3, Lane 6= F3, Lane 7 = G3, Lane 8 = G9, Lane 9= H4, Lane 10=H8. Lane 11= I1, Lane 12 = I2, Lane 13 = I14, Lane 14 = J11, Lane 15 = J12, Lane 16 = K5, Lane 17 = K7, Lane 18 = K19, Lane 19 = L5, Lane 20 = L6



0.020

Figure 2: Phylogenetic analysis of selected *E. coli* isolates

1 **4. CONCLUSION**

2 Conclusion: The *qnrS* and *blaCMY* genes found in multiple antibiotic resistant *E. coli*
3 mediated resistance to critically important antibiotics and co-existed in variants strains of *E.*
4 *coli* occupying different clusters in the phylogenetic analysis. Wrong policies on antibiotic
5 formulation and crude treatment regime in poultry promote the expression of these genes
6 harbored in the genome of variant strains of *E. coli*.

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11 **COMPETING INTERESTS**

12 No competing interests exist.

13 **AUTHORS' CONTRIBUTIONS**

14 Author A designed the study managed the analyses of the study. Author B wrote the
15 protocols, conducted the experiments, wrote the first draft of the manuscript and managed
16 the literature searches. All authors read and approved the final manuscript.”

17 **CONSENT (WHERE EVER APPLICABLE)**

18 Not applicable

19 **ETHICAL APPROVAL (WHERE EVER APPLICABLE)**

20 Not applicable

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