

## Genetic Diversity and Quinolone Resistance in *Campylobacter jejuni* Isolates from Poultry in Senegal

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**We used the multilocus sequence typing (MLST) method to evaluate the genetic diversity of 46 *Campylobacter jejuni* isolates from chickens and to determine the link between quinolone resistance and sequence type (ST). There were a total of 16 ST genotypes, and the majority of them belonged to seven clonal complexes previously identified by using isolates from human disease. The ST-353 complex was the most common complex, whereas the ST-21, ST-42, ST-52, and ST-257 complexes were less well represented. The resistance phenotype varied for each ST, and the Thr-86-Ile substitution in the GyrA protein was the predominant mechanism of resistance to quinolone. Nine of the 14 isolates having the Thr-86-Ile substitution belonged to the ST-353 complex. MLST showed that the emergence of quinolone resistance is not related to the diffusion of a unique clone and that there is no link between ST genotype and quinolone resistance. Based on silent mutations, different variants of the *gyrA* gene were shown to exist for the same ST. These data provide useful information for understanding the epidemiology of *C. jejuni* in Senegal.**

*Campylobacter jejuni* is the leading cause of bacterial foodborne diarrheal disease throughout the world (2). In developing countries, the estimated incidence of *Campylobacter* in diarrhea is between 40,000 and 60,000 per 100,000 for children less than 5 years old and is 90 per 100,000 in the general population (3, 5, 19). Organisms are carried in the intestinal tracts of many animals, including birds, and epidemiological evidence has suggested that this site is a potential reservoir for human infections and an entry point into the food chain (12, 14, 17, 22). Most human infections with *C. jejuni* are self-limiting and do not require antimicrobial therapy. However, treatment with erythromycin or a fluoroquinolone is needed in individuals with invasive or severe disease, in immunocompromised patients, or in children less than 5 years old. But resistance to fluoroquinolone is increasing throughout the world (10). In *Campylobacter*, the predominant mechanism of fluoroquinolone resistance is the C-to-T transition at codon 86 in the quinolone resistance-determining region (QRDR) of the *gyrA* gene, which results in a threonine-to-isoleucine substitution in the functional protein (1, 24).

To study the epidemiology of *Campylobacter* infections, multiple typing methods have been developed in order to differentiate isolates below the species level (25). The multilocus sequence typing (MLST) method was developed by Dingle et al. (8) to characterize *C. jejuni* strains and to identify clonal lineages in this species (6, 7, 9, 16). This method uses genetic variations at multiple chromosomal locations and allows generation of sequence data, which may be available via the Internet for comparison with DNA sequences of other strains.

To date, there are little data on the epidemiology of *Campylobacter* in Senegal; one study was a study of the incidence of quinolone-resistant *C. jejuni* isolates in commercial chickens, which was 40% (4).

The aims of the present study were (i) to evaluate the genetic diversity of *C. jejuni* strains isolated from chickens in Senegal and (ii) to determine the link between quinolone resistance and sequence type (ST) genotypes.

### MATERIALS AND METHODS

***Campylobacter* isolates.** The *C. jejuni* isolates used originated from chicken carcasses and were collected during 2000 to 2002 in the Dakar region (Senegal) and suburbs (4). All of the bacterial isolates had been maintained as stock cultures in glycerol broth at  $-80^{\circ}\text{C}$ . A total of 46 isolates that were randomly picked from 99 *C. jejuni* isolates were included in the study, and these strains were obtained from 14 dispersed collection sites over a 3-year period. Prior to DNA extraction, cultures were removed from storage and allowed to thaw at room temperature. Isolates were cultured on Columbia agar plates containing 5% sheep blood and were incubated under microaerophilic conditions (5%  $\text{O}_2$ , 10%  $\text{CO}_2$ , 85%  $\text{N}_2$ ) at  $42^{\circ}\text{C}$  for 48 h.

**MLST.** Bacterial DNA was prepared by boiling as reported previously (1). MLST was performed as described by Dingle et al. (8) by using sequences obtained from seven housekeeping genes. PCR products were purified with a QIAquick gel extraction kit (QIAGEN, Germany), and the DNA sequences of both strands were determined with an ABI PRISM 310 DNA sequencer (Applied Biosystems). MLST alleles and sequence types were assigned to the isolates by comparing data to the *Campylobacter* MLST database (<http://campylobacter.mlst.net>). The isolates were organized into lineages with the program BURST (based on related STs), which is part of the START (sequence type assignment recombination tests) group of programs. Isolates were grouped together if they shared five, six, or all seven MLST loci. eBURST was used to examine patterns of evolutionary descent (11). Analysis of fixed differences, polymorphic sites, and the ratio of nonsynonymous substitutions to synonymous substitutions was performed with the DnaSP software package (version 4.00), available at <http://www.ub.edu/dnasp/> (20). The index of association ( $I_A$ ) was calculated using the program START (15).

**Antibiotic susceptibility.** The antibiotic susceptibility of each isolate was determined by the disk diffusion method. The MICs of ciprofloxacin and nalidixic

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acid were determined by the Etest method and were interpreted according to CLSI (formerly NCCLS) guidelines (18). Isolates were inoculated onto Mueller-Hinton blood agar, and Etest strips (AB Biodisk, Solna, Sweden) were applied to each plate. *C. jejuni* subsp. *jejuni* ATCC 33560 (= NCTC 11351) was included as a quality control strain. The plates were incubated at 42°C under microaerophilic conditions (5% O<sub>2</sub>, 10% CO<sub>2</sub>, 85% N<sub>2</sub>) for 24 h. The breakpoint for ciprofloxacin was considered to be an MIC of  $\geq 4$   $\mu$ g/ml.

**Determination of mutations in the QRDR of *gyrA*.** A 220-bp fragment covering the QRDR of *gyrA* was amplified by PCR by using the primers and conditions described previously, except that the annealing temperature was 50°C (24). After purification (QIAquick gel extraction kit; QIAGEN, Germany), sequencing of the amplified fragment was performed with the same primers using an ABI PRISM 310 DNA sequencer (Applied Biosystems). DNA sequences were compared to the previously published *C. jejuni* sequence (GenBank accession number L04566) using the CLUSTALW program available at <http://bioweb.pasteur.fr/seqanal/interfaces/clustalw.html>.

## RESULTS

**Diversity of MLST alleles.** The number of unique alleles at each locus varied from five for the Asp locus to nine for the Gln locus, and the percentage of variable sites in the data set ranged from 0.6% for the *uncA* locus to 4.8% for the *pgm* locus (Table 1). For all seven loci, the ratio of nonsynonymous substitutions to synonymous substitutions was much less than 1.

**Diversity of STs and lineages.** There were a total of 16 STs among the 46 isolates examined. Several of these STs (ST1035, ST1036, ST1037, ST1038, ST1039, ST1040, ST1041, ST1081, ST1358, ST1359, and ST1370) had not been described previously at the *Campylobacter* MLST database (<http://campylobacter.mlst.net>). Using the program BURST, the 46 isolates were grouped into four lineages (Table 2), and seven singleton sequence types were unrelated to any other sequence types based on the fact that the isolates in one group were identical at five, six, or all seven of the MLST loci. The most common ST was ST1036, which comprised 11 isolates (23.91% of the data set), followed by ST1035, which comprised 8 isolates (17.39%), and ST1039 with 5 isolates (10.87%). The seven singleton STs (ST1370, ST1359, ST1358, ST1211, ST1035, ST824, and ST52) included 17 isolates and accounted for 36.95% of the data set. A total of seven clonal complexes were identified (ST-21, ST-22, ST-42, ST-52, ST-257, ST-353, and ST-354 complexes). Within each clonal complex, the number of isolates ranged from 1 (ST-21, ST-42, and ST-257 complexes) to 12 (ST-353 complex). eBURST analysis allowed us to determine the relationships between ST1040 (unassigned) and ST1036 (ST-353 complex) on the one hand and ST1211 (ST-353 complex) on the other hand (Fig. 1). Indeed, the BURST analysis revealed that both ST1040 and ST1036 belong to group 1, whereas ST1211 was a singleton ST. ST1039 and ST1041 were grouped in the same lineage (group 2) but were not assigned to any complex based on the data in the *Campylobacter* MLST database. The other STs that were not assigned to any complex were ST1035 and ST1358. The  $I_A$  for the complete data set was 2.028, and the  $I_A$  was 0.371 when only one representative of each lineage was included.

**Resistance phenotypes and mutations of the *gyrA* QRDR.** Twenty-four isolates were susceptible to both ciprofloxacin and nalidixic acid, 3 isolates were susceptible to ciprofloxacin but had intermediate susceptibility to nalidixic acid, and 19 isolates were resistant to both antibiotics (Table 2). The great majority of quinolone-resistant isolates had a change from C to T in codon 86 of the *gyrA* gene, resulting in a threonine-to-isoleucine substitution in the functional protein. Another substitu-

TABLE 1. Allelic diversity

Locus	Fragment size (bp)	No. of alleles	No. of variable sites	% Variable sites	$d_N/d_S^a$
<i>aspA</i>	477	5	6	1.3	0.022
<i>glnA</i>	477	9	13	2.7	0.006
<i>gltA</i>	402	6	6	1.5	0.062
<i>glyA</i>	507	6	22	4.3	0.001
<i>pgm</i>	498	8	24	4.8	0.000
<i>tkt</i>	459	7	17	3.7	0.002
<i>uncA</i>	489	4	3	0.6	0.043

<sup>a</sup>  $d_N/d_S$ , ratio of nonsynonymous substitutions to synonymous substitutions.

tion in the same codon was the Thr-86-Ala substitution observed in nalidixic acid-resistant isolates and isolates with intermediate susceptibility. Forty-three isolates had silent mutations at His-81 (CAC→CAT) and Ser-119 (AGT→AGC), and 10 isolates had an additional silent mutation at Gly-110 (GGC→GGT).

**Relationships between ST, resistance phenotypes, and *gyrA* mutations.** In all groups except group 3, the resistance phenotype varied within each ST (Table 2). However, certain phenotypes were predominant. In group 1, the most common phenotype was double resistance to ciprofloxacin and nalidixic acid, whereas groups 2 and 4 contained a majority of susceptible isolates. In singleton STs, the resistance phenotypes were diverse. Three phenotypes were identified among isolates assigned to ST1035, while ST52 contained only susceptible isolates. Eleven of the 14 isolates having the Thr-86-Ile substitution belonged to group 1. Based on silent mutations, different variants of the *gyrA* gene were shown to exist for the same ST genotype; the first variant had a silent mutation at His-81 and Ser-119, and the second variant had an additional silent mutation at Gly-110.

## DISCUSSION

The isolates analyzed in this study originated only from chickens, in contrast to other studies in which isolates having diverse origins were examined (8, 16). This study is the first study in which the MLST approach was used to examine the clonal relationships of *C. jejuni* isolates in Senegal. Although the number of isolates was relatively small, the diversity of the isolates was established by identification of various genotypes and clonal complexes.

The allelic diversity observed here was similar to the diversity observed previously in farm animals by Colles et al. (6) with regard to the number of alleles and nucleotide variability; the only exception was the *uncA* locus, for which Colles et al. found a higher degree of diversity.

In this study, we identified seven clonal complexes, and all of these complexes have been associated with human disease (9, 21). Consistent with our results, Dingle et al. (8) identified six clonal complexes in 34 isolates obtained from chickens. In contrast to previous studies in which the ST-21 complex was the largest complex (6, 7, 8), this complex appears to be uncommon in Senegalese chickens since its ST1359 genotype occurred only once in the data. The other complexes that were less well represented were ST-42, ST-52, and ST-257. In Europe, these complexes were identified for various isolates from

TABLE 2. MLST lineages, ST, *gyrA* mutations, and resistance phenotypes

BURST lineage	Clonal complex	ST	Isolate	Year	<i>gyrA</i> mutations <sup>a</sup>	Phenotype <sup>b</sup>	
						Ciprofloxacin	Nalidixic acid
Group 1	353	1036	83B	2001	His-81, Ser-119	S	S
			104C	2002	His-81, Ser-119	S	S
			116B	2002	His-81, Ser-119, Thr-86-Ile, Gly-110	R	R
			65B	2001	His-81, Ser-119, Thr-86-Ile	R	R
			48A	2000	His-81, Ser-119, Thr-86-Ile	R	R
			48B	2000	His-81, Ser-119, Thr-86-Ile	R	R
			65A	2001	His-81, Ser-119, Thr-86-Ile	R	R
			65C	2001	His-81, Ser-119, Thr-86-Ile	R	R
			F19	2001	His-81, Ser-119, Thr-86-Ile	R	R
			118C	2002	His-81, Ser-119, Thr-86-Ile, Gly-110	R	R
			82A	2001	His-81, Ser-119, Thr-86-Ile	R	R
			113B	2002	His-81, Ser-119, Thr-86-Ile	R	R
			93A	2001	His-81, Ser-119	S	S
			112A	2002	His-81, Ser-119, Thr-86-Ile	R	R
			Group 2	Unassigned	1039	50C	2000
75A	2001	His-81, Ser-119				S	S
95B	2001	His-81, Ser-119				S	S
95A	2001	His-81, Ser-119				R	R
57A	2001	His-81, Ser-119				S	S
62A	2001	His-81, Ser-119				S	S
64C	2001	His-81, Ser-119				S	S
F18	2001	His-81, Ser-119, Thr-86-Ala, Gly-110				R	R
Group 3	22	1081	118B	2002	None	S	S
			660	2002	None	S	S
			22	2002	None	S	S
Group 4	354	1037	106B	2002	His-81, Ser-119, Thr-86-Ile	R	R
			1038	2002	His-81, Ser-119	S	S
			115B	2002	His-81, Ser-119	S	S
			V34A	2002	His-81, Ser-119	S	S
Singleton STs	52	52	44B	2000	His-81, Ser-119	S	S
			47C	2000	His-81, Ser-119	S	S
			72B	2001	His-81, Ser-119	S	S
			55A	2001	His-81, Ser-119	S	S
			103C	2002	His-81, Ser-119	S	S
			90B	2001	His-81, Ser-119	S	S
			115A	2002	His-81, Ser-119	S	S
			41B	2000	His-81, Ser-119	S	S
			84C	2001	His-81, Ser-119	R	R
			130A	2000	His-81, Ser-119, Gly-110	S	S
			76C	2001	His-81, Ser-119, Thr-86-Ala, Gly-110	S	I
			110B	2002	His-81, Ser-119, Thr-86-Ile, Gly-110	R	R
			76A	2001	His-81, Ser-119, Thr-86-Ala, Gly-110	S	I
			76B	2001	His-81, Ser-119, Thr-86-Ala, Gly-110	S	I
			78C	2001	His-81, Ser-119, Thr-86-Ala, Gly-110	R	R
			78A	2001	His-81, Ser-119, Thr-86-Ala, Gly-110	R	R
110A	2002	His-81, Ser-119, Thr-86-Ile	R	R			

<sup>a</sup> Silent mutations are His-81 (CAC→CAT), Ser-119 (AGT→AGC), and Gly-110 (GGC→GGT).

<sup>b</sup> R, resistant; S, susceptible; I, intermediate.

farm animals (6, 16, 21) and also for human isolates (6, 21), suggesting that they have a wide distribution. The ST-353 complex, the most common complex in this study, and the ST-22 and ST-354 complexes have all been found previously in poultry (16). Other studies have shown that these complexes contain isolates associated with human gastroenteritis (7, 9, 16). Indeed, a search of the *Campylobacter* MLST database revealed that these complexes contained a majority of the isolates obtained from human disease. Despite the lack of MLST

data for humans in Senegal, one could envisage that these complexes represent pathogen strains.

A number of new STs were identified for the first time in this study (ST1035, ST1036, ST1037, ST1038, ST1039, ST1040, ST1041, ST1081, ST1358, ST1359, and ST1370), and some of them were not assigned to any known complex (ST1035, ST1039, ST1040, ST1041, and ST1358). In group 1, for instance, MLST showed that isolates with the ST1040 genotype are single-locus variants of ST1036, which is a member of the

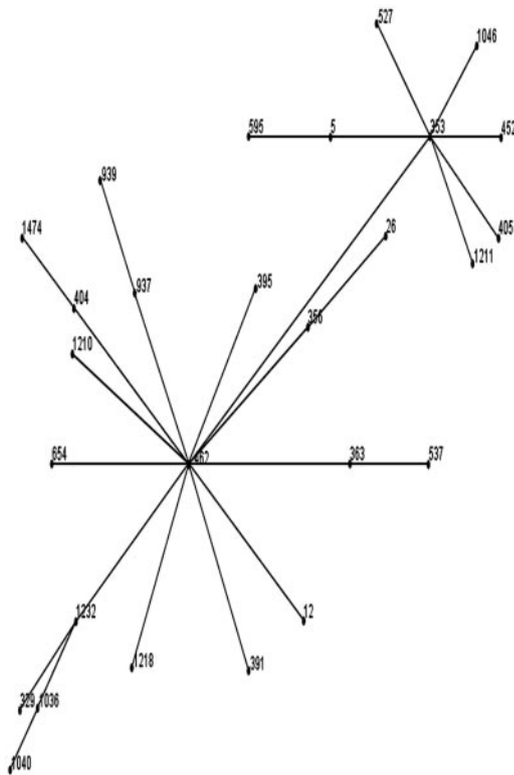


FIG. 1. Relationship between isolates of the ST-353 complex from the *Campylobacter* MLST database and ST1040 as determined by eBURST analysis. ST1036 is the ancestor of ST1040.

ST-353 complex. eBURST analysis of all ST-353 complex isolates in the MLST database with ST1040 indicated that ST1040 was derived from ST1036 (Fig. 1); the only difference was a single nucleotide polymorphism in the *gltA* gene sequence. Therefore, ST1040 probably diverged from ST1036 very recently. Interestingly, based on eBURST and BURST analyses of all ST-353 complex isolates in the MLST database it also appeared that ST462 rather than ST353 was the founder of the complex. As more MLST data are added to the database, more new links should be seen, and it is also likely that conclusions concerning the clonal complex founder will have to be modified. Also, eBURST analysis showed the relationship between ST1211 and ST1036. ST1211 was grouped among the singleton STs by BURST analysis, even though both ST1211 and ST1036 belonged to the ST-353 complex, and we explained this classification by the absence of intermediate STs (ST1232, ST462, and ST353, not present among our isolates) between ST1211 and ST1036 and also by the fact that only three of the seven MLST loci are shared by the two STs. Singleton STs were not negligible in our study (36.95% of the data set), and they represent distant genotypes not related to any other genotypes. Thus, we believe that with more MLST data, new linkages (not seen in this study) will be discovered. Another finding was that group 2 contained only STs (ST1039 and ST1041) that were not assigned to any complex. Perhaps these genotypes were the result of recombination events (23). Based on the complete data set, linkage analysis revealed that the  $I_A$  was 2.028, which is comparable to the value found by Dingle et al. (8) (2.016). In

our analysis, the  $I_A$  was 0.371 when only one representative of each lineage was included, whereas Dingle et al. (8) obtained an  $I_A$  of 0.56 and concluded that this indicated that there was a weakly clonal population. In addition, Suerbaum et al. (23) obtained an  $I_A$  of 0.256, which was comparable to our value, and suggested that there was a limited amount of recombination. Although the number of isolates analyzed in this study was too small to draw any definite conclusions, certain STs (ST1035, ST1036, and ST1039) and clonal complexes (ST-353, ST-354, and ST-52 complexes) seem to be predominant, even though other genotypes and clonal complexes may still be present in Senegalese chickens.

As previously reported (1, 24), the Thr-86-Ile substitution in the GyrA protein was the major mechanism of quinolone resistance, although a small number of isolates had the Thr-86-Ala substitution. In Senegal, fluoroquinolones (enrofloxacin and norfloxacin) were introduced into poultry production in 1996 (4), and their use became an important selective factor in the spread of quinolone-resistant clones. The fact that quinolone resistance was observed in STs of different MLST lineages (except the ST-22 complex) suggests that the emergence of quinolone resistance is not related to the diffusion of a unique clone but rather is the result of quinolone selection pressure on strains from different clones. Nevertheless, there might be a relationship between isolates carrying the Thr-86-Ile substitution and the ST-353 complex. It also appeared that there were different variants of the *gyrA* gene for the same ST genotype. This illustrates the variability of the *C. jejuni* genome outside the MLST loci. Previous studies have argued that *gyrA* polymorphism could be useful in epidemiological studies (13, 26). We believe that this polymorphism may provide information that supplements the data obtained by the MLST approach for understanding the epidemiology of this pathogen. Generally speaking, isolates that had the same ST genotype had variable resistance phenotypes, suggesting that there was no link between ST genotype and quinolone resistance.

To date, there is little information concerning the epidemiology of *Campylobacter* in Senegal, and in particular, there are no data on human isolates due to the absence of systematic investigations of *Campylobacter* in medical laboratories. This report provides more insight into the molecular biology of this bacterium in poultry production. Further studies with a larger number of isolates, including isolates from humans and several time frames, should provide more data on the epidemiology of *Campylobacter* in Senegal.

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