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Full Length Research Paper

Phylogenetic analysis of *Campylobacter jejuni* from human and birds sources in Iraq

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The present study was designed for a phylogenetic tree analysis of *Campylobacter* species as molecular method for genetic identification of *Campylobacter jejuni* from human and birds sources and birds and amplified by polymerase chain reaction assay using specific primers for 16S rRNA gene of *Campylobacter jejuni* (GenBank: EF136575.1). In this study, the multiple sequence alignment analysis and neighbor joining phylogenetic tree analysis was performed by using Unweight Pair Group method with Arithmetic Mean (UPGMA tree) in MEGA 6.0 version that analyzes 827 bp for ribosomal 16S rRNA gene. *C. jejuni* was detected in 40% (16/40) of stool samples collected from patients suffering from gastroenteritis, while detection rates of *C. jejuni* were 15% (3 /20) and 10% (2/20) of fecal samples of domestic chicken and pigeon respectively by PCR assay. The phylogenetic analysis results revealed that all local isolates of *Campylobacter* spp. were closed related to NCBI-Blast *C. jejuni* strain No.Y19244.1, whereas other NCBI-Blast *Campylobacter* spp. were out of tree and more different to ten *Campylobacter* spp. Iraq isolates and also found the relationships between the local isolates of *Campylobacter* spp. (Human, Domestic chicken, and Pigeon). This study represents the first report on the use of molecular phylogeny to *Campylobacter* spp. obtained in Iraq and confirmed the zoonotic potential of *C. jejuni*.

Key words: Phylogenetic tree, Campylobacter species, 16S ribosomal gene, human, birds, Iraq.

INTRODUCTION

Campylobacteriosis is a common zoonotic disease that affect human and cause gastrointestinal disturbances (Barakat et al., 2013). *Campylobacter jejuni* is responsible for 90% of *Campylobacter* species human infections and they occur in sporadic way (Schielke et al., 2014). *Campylobacter* is one of the most frequently occurring bacterial agents of gastroenteritis in human (WHO, 2012). Most bird flocks are colonized within several days and still so until slaughter. The handling and ingesting of contaminated meat with *Campylobacter*, especially poultry meat is considered an important source of food-borne gastroenteritis in human (Hermans et al., 2011). Today, attention has turned to nucleic acid technology; the polymerase chain reaction (PCR) and related techniques are rapid, specific and sensitive as compared to other tests used in detection of

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Campylobacter spp. (Englen and Kelley, 2000). C. jejuni is the Campylobacter spp. predominantly found in infected humans and colonized broilers. Sequence analysis of the 16S rRNA gene is very useful for identification of bacteria to genus and species level (Hansson et al., 2008). The potential application of the 16S rRNA gene for determining phylogenetic relationships among all living organisms had attracted much interest and would play a major role in extensive rearrangement of Campylobacter taxonomy (Woese, 1987). Phylogenetic analysis may be used as a molecular tool in future studies in the surveillance of Campylobacter-like organisms (Nayak et al., 2014). C. jejuni is isolated from stool samples of diarrheic children and confirmed phenotypically on the basis of biochemical tests in many provinces in Iraq (Saliih and Al-Saad, 1994; Mohammad et al., 2004; Al-Ani et al., 2008). C. jejuni is identified by conventional PCR assay in human and domestic chicken in Al-Qadissiva province. Iraq (Al-Hisnaway, 2008). Abd (2014) proved that the detection rate of C. jejuni in human was 55.2% by Real-Time PCR Assay in Al-Muthanaa province, Iraq. The present study aimed at examining and analyzing the partial 16S ribosomal RNA gene sequence for construction of phylogenetic trees analysis of Campylobacter spp. Iraq isolates from infected humans, domestic chicken and pigeons in comparison to those of other NCBI-Blast Campylobacter spp.

MATERIALS AND METHODS

Samples collections

Human stool samples

A total of 40 stool samples of patients suffering from enteritis with ages ranging from 1 to 50 years were collected from general hospital in Al-Qadissiya province, Iraq during a period 6 months from October 2014 to March 2015 and after clinician consultation (included diarrhea, symptoms comprising vomiting, abdominal pain, fever) and microscopically examination in the hospitals where many samples contain motile bacteria, pus and few contain mucous and blood.

Bird samples

Fresh fecal samples were collected randomly from 20 flocks of domestic chicken from different farms, as well as 20 fecal samples of pigeon were collected from the same farms in Al-Qadissiya province, Iraq. The samples were collected during a period 6 months from October 2014 to March 2015.

All samples were placed in test tube containing 3 ml of peptone water in sterile condition and were immediately transported to the laboratory during 3 to 6 h in a cooler with ice packs. All the samples were frozen at -20°C for DNA extraction.

Genomic DNA extraction

Genomic DNA was extracted from stool samples by using AccuPrep® Stool DNA Extraction Kit, Bioneer, Korea. The

extraction was done according to company instructions. After that, the extracted gDNA was checked by Nanodrop spectrophotometer, store in -20°C in refrigerator until perform PCR.

Polymerase chain reaction (PCR)

PCR assay was carried out by using specific primer which was designed in this study from highly conserved regions of 16S ribosomal of C. jejuni (GenBank: EF136575.1). 16SrRNA forward primer (CGCACGGGTGAGTAAGGTAT) and 16SrRNA reverse primer (TAAACACATGCTCCACCGCT) were provided by Bioneer company, Korea and using DNA C. jejuni as positive control and it was provided by Genekam, Germany. PCR master mix was prepared by using AccuPower[®] PCR PreMix kit Bioneer, Korea. The PCR premix tube contains freeze-dried pellet of Taq DNA polymerase 1U, dNTPs 250 µM, Tris-HCI (pH 9.0) 10 mM, KCI 30 mM, MgCl₂ 1.5 mM, stabilizer, and tracking dye and the PCR master mix reaction was prepared according to kit instructions in 20 µl total volume by adding 5 µl of purified genomic DNA and 1.5 µl of 10 pmole of forward primer and 1.5 µl of 10 pmole of reverse primer, then complete the PCR premix tube by deionizer PCR water into 20 µl and briefly mixed by Exispin vortex centrifuge (Bioneer. Korea). The reaction was performed in a thermocycler (Mygene Bioneer. Korea) by set up following thermocycler conditions; initial denaturation temperature of 95°C for 5 min; followed by 30 cycles at denaturation 95°C for 30 s, annealing 58°C for 1 min, and extension 72°C for 1 min and then final extension at 72°C for 10 min. The 827 bp PCR products were examined by electrophoresis in a 1% agarose gel, stained with ethidium bromide, and visualized under UV trans-illuminator.

DNA sequencing method

The 827 bp PCR product was purified from agarose gel by using EZ EZ-10 Spin Column DNA Gel Extraction Kit, Biobasic. Canada. The purified 16S rRNA gene PCR product samples were sent to Bioneer Company in Korea to perform the DNA sequencing using 16SrRNA forward primer by AB DNA sequencing system. DNA sequencing method was performed for confirmative Phylogenetic tree relationship analysis of *Campylobacter* spp. based on 16S ribosomal RNA gene by Phylogenetic tree analysis using Unweight Pair Group method with Arithmetic Mean (UPGMA tree) in MEGA 6.0 version.

RESULTS AND DISCUSSION

Identification of *C. jejuni* in human and birds by PCR assay

Campylobacter is considered as human pathogen despite of it commensal organisms in domestic poultry and livestock. The present study describes a molecular method for detection C. jejuni from human and bird sources by using specific primer of 16S ribosomal of C. jejuni (Figure 1). Polymerase chain reaction (PCR) analysis using Campylobacter genus-specific partial 16S primers rRNA revealed the presence of Campylobacter spp. DNA in the faces (Turowski et al., 2014), where conventional PCR is rapid as nearly 2 times and sensitive method to determine *Campylobacter* spp. in comparison with culturing and this enhance its application

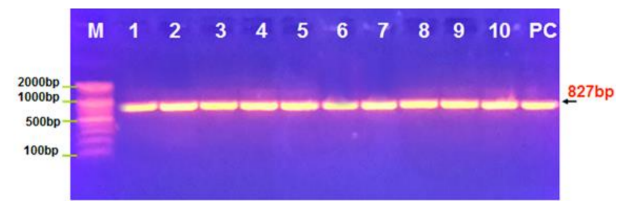


Figure 1. Agarose gel electrophoresis image that shown the PCR product of 16S rRNA gene that using in detection of *Campylobacter jejuni* Where M: Marker (1500-100bp), lane (1-3) domestic chicken, lane (4-5) pigeon, (6-10) Human were positive *Campylobacter jejuni* isolates and lane (PC) DNA Campylobacter genus positive control (Genekam, Germany) at 827bp PCR product size.

Host	No. of samples	Campylobacter jejuni positive	Percentage
Human	40	16	40
Chicken	20	3	15
Pigeon	20	2	10

as timesaving method of *Campylobacter* spp. by using 16SrRNA gene primer (Stoyanchev, 2004). Zhang et al. (2013) proved that PCR assay was sensitive (100%) in comparison with (49%) sensitivity of direct bacterial culture.

In present study, detection rate of C. jejuni in human was 16 (40%) out of (40) stool samples collected from infected patients which suffered from diarrhea and some of them suffered other symptoms such as fever, colic and vomiting. This result shows that detection rate of C. jejuni in human was relatively low, compared with the results of other studies reported in Iraq describing Campylobacter in human, which reported that the prevalence of campylobacteriosis was 55.2 and 66.7% (Abd, 2014; Al-Amri et al., 2007), while this result was higher than that record by Al-Hisnaway (2008) who found C. jejuni in 33.3% of stool samples of human by PCR assay in Al-Qadissiya province in Iraq. The different detection rate of the present study in comparison with other studies may influence many factors such as age, season, geography and immune state of human.

In this study, 3/20 (15%) fecal samples of domestic chicken were identified as *C. jejuni* by using specific primers of 16S ribosomal of *C. jejuni* by PCR assay and this result agree with AI-Hisnaway (2008) who detected *C. jejuni* with 17.6% from chicken fecal samples by conventional PCR assay, where *C. jejuni* has been reported to be the most frequent species recovered from poultry and poultry carcasses (Jorgensen et al., 2002).

The occurrence of *C. jejuni* in pigeon feces has been studied in several countries worldwide. In the present study, the detection rate of *C. jejuni* in pigeons fecal samples was 10% (2/20) (Table 1), this result was lower than that record by Casanovas et al. (1995) who found *Campylobacter* spp. in 26.2% of fecal pigeon samples and all of *Campylobacter* species isolated from pigeon fecal samples was *C. jejuni* (100%).

Sequencing analysis of 16S rRNA genes of Campylobacter spp. Iraqi isolates

The partial sequences for 16S ribosomal RNA genes of ten Iraq isolates *Campylobacter* spp. can be found under the accession numbers at NCBI-Gen Bank submission and they are shown in Table 2. Sequence analysis of ten samples positive for *Campylobacter* spp. was performed to confirm the PCR results in this study, the DNA sequencing analysis of 16S rRNA gene 827 bp PCR product by multiple sequence alignment Unweight Pair Group method with Arithmetic Mean (UPGMA tree) in MEGA 6.0 version showed specific detection of *C. jejuni*. These studies agreed with Dewhirst et al. (2005) who identified representing either *C. jejuni* or *Campylobacter coli* by 16S rRNA sequence analysis.

The nucleotide sequences of the 16S rRNA genes of ten *Campylobacter* spp. Iraq local isolates of human and birds were determined and compared with 16S rRNA

Campylobacter spp.	Sources/sample	Gen bank accession numbers			
IQDC-1	Domestic chicken/Feces	KR133485.1			
IQDC-2	Domestic chicken/Feces	KR133486.1			
IQDC-3	Domestic chicken/Feces	KR133487.1			
IQP-1	Pigeons/Feces	KR133488.1			
IQP-2	Pigeons/Feces	KR133489.1			
IQH-1	Human/Stool	KR133490.1			
IQH-2	Human/Stool	KR133491.1			
IQH-3	Human/Stool	KR133492.1			
IQH-4	Human/Stool	KR133493.1			
IQH-5	Human/Stool	KR133494.1			

 Table 2. Gen bank accession numbers of 16S ribosomal RNA gene, partial sequence for Campylobacter spp. Iraq isolates from Human and bird sources.

Table 3. The sequence identity for Campylobacter spp. Iraq isolates from human.

	• • • • • • • • •	Campylobacter spp. Iraq isolates of human									
Campylobacter spp. strains	Accession	IQ.H-1		IQ. H -2		IQ. H-3		IQ. H -4		IQ. H -5	
	number	Score	Identify (%)	Score	Identify (%)	Score	Identify (%)	Score	Identify (%)	Score	Identify (%)
C. Jejuni	Y19244.1	1495	99	1498	99	1469	99	1504	99	1495	99
C. faecalis	AJ.276874.1	1168	92	1194	93	1147	92	1170	92	1168	92
C. fetus	AJ.306569.1	1280	95	1273	95	1254	95	1184	95	1280	95
C. subantarctic	AM. 933373.1	1478	99	1482	99	1452	99	1509	99	1478	99
C. subantarctic	AM. 933374.1	1478	99	1482	99	1452	99	1509	99	1478	99
C. volucis	FM.883695.1	1439	98	1443	98	1413	98	1476	99	1439	98
C. hominis	AJ.251584.1	1114	91	1149	92	1098	92	1096	91	1114	91
C. coli	AM.042699.1	1210	99	1498	99	1258	99	1020	99	1210	99

sequences of eight strains of *Campylobacter* spp. The results showed that the sequence identity was 99% between ten *Campylobacter* spp. Iraq local isolates and *C. Jejuni* (Y19244.1), *Campylobacter subantarctic* (AM. 933373.1), *C. subantarctic* (AM. 933374.1) and *C. coli* (AM.042699.1) (Tables 3 and 4). The bacteria

with relatively small genomes, such as *C. jejuni* may undergo genetic variation to increase their potential to adapt to new environments; such genotypic variation could result in phenotypic changes. These variations are probably important in the transmission route from broiler to man, where *Campylobacter* spp. must survive several

hostile environments (Hansson et al., 2008).

Phylogenetic analysis

Phylogenetic tree analysis based on the clone 16S rRNA gene, partial sequence used for confirmative detection of *Campylobacter* spp. Iraq

	Accession ns number	Campylobacter spp. Iraq isolates of birds									
Campylobacter spp. strains		IQ DC -1		IQ DC -2		IQ DC -3		IQ P-1		IQ P-2	
		Score	Identify (%)	Score	Identify (%)	Score	Identify (%)	Score	Identify (%)	Score	Identify (%)
C. Jejuni	Y19244.1	1509	99	1495	99	1506	99	1511	99	1500	99
C. faecalis	AJ.276874.1	1158	92	1168	92	1218	93	1206	93	1157	92
C. fetus	AJ.306569.1	1168	95	1280	95	1303	95	1286	95	1264	94
C. subantarctic	AM. 933373.1	1515	99	1478	99	1489	99	1495	99	1483	99
C. subantarctic	AM. 933374.1	1515	99	1478	99	1489	99	1495	99	1483	99
C. volucis	FM.883695.1	1482	99	1439	98	1450	98	1456	98	1445	98
C. hominis	AJ.251584.1	1090	91	1114	91	1168	92	1162	92	1109	91
C. coli	AM.042699.1	1003	99	1210	99	1522	99	1506	99	1194	99

Table 4. The sequence identity for Campylobacter spp. Iraq isolates from domestic chicken and pigeon.

isolates that included this study where phylogenetic analysis of 16S rRNA gene sequences has become the primary method for determining prokaryotic phylogeny. Therefore, the validity of 16S rRNA gene based phylogenetic analyses is of fundamental importance for prokaryotic systematics (Dewhirst et al., 2005). However, studies have suggested that multiple strains should be investigated to evaluate the degree of sequence diversity within and between species (Clayton et al., 1995). In the present study, the phylogenetic tree was constructed based on the ten Campylobacter spp. Irag isolates included {(n=5) human, (n=3) chicken and (n=2) pigeons} and nine strains of NCBI-Blast Campylobacter spp.

The ten *Campylobacter* spp. Iraq isolates showed close relationship with NCBI-Blast *C. jejuni* (Y19244.1) compared to other strains of NCBI-Blast *Campylobacter* spp. (Figure 2). These results agreed with Weis et al. (2014) who used phylogenetic analyses of 16S rRNA sequence data to distinguish *C. jejuni* from other species and to map strains found in crows with strains previously isolated from humans, livestock, and poultry. Nayak et al. (2014) referred to phylogenetic analysis providing a rapid, accurate and effective method for identification of species within the *Campylobacter*.

Host relationship analysis of *Campylobacter* spp. Iraq strains

In present study, we have investigated putative specificity of the host using phylogenetic analysis of genetically closely related Campylobacter spp. from different sources where recent studies have suggested a potential role for birds in zoonotic transmission of Campylobacter spp., the leading cause of gastroenteritis in humans worldwide (Petersen et al., 2001; Broman et al., 2004; Weis et al., 2014). The results showed Campylobacter spp. IQH-2(KR133491.1) and Campylobacter spp.IQH-3(KR133492.1) isolates of human were more close relationship with Campylobacter spp.IQP-1 (KR133488.1) and Campylobacter spp.IQP-2 (KR133489.1) isolates of pigeons, as well as with Campylobacter spp.IQDC-1(KR133485.1) and Campylobacter spp.IQDC -3(KR133487.1) isolates of domestic chickens.

Campylobacter spp. IQH-1(KR133490.1), *Campylobacter* spp.IQH-4 (KR133493.1) and *Campylobacter* spp.IQH-5 (KR133494.1) of human were close related with *Campylobacter* spp.IQDC-2 (KR133486.1) of domestic chicken (Figure 3). These results agreed with Schouls et al. (2003) who record about 75% of the human strains were found to be most closely related to the patterns of the other human strains, and the patterns of 20% of the human strains were more similar to the patterns of the strains isolated from poultry.

Conclusion

This study suggested that phylogenetic tree analysis is based on 16S ribosomal RNA gene, partial sequence can be used for confirmative detection of *Campylobacter* spp. isolates and determine the close relationship between *Campylobacter* spp. isolated from human, domestic chicken and pigeon. These results highlighted the importance of domestic chicken and pigeon as a potential source of human

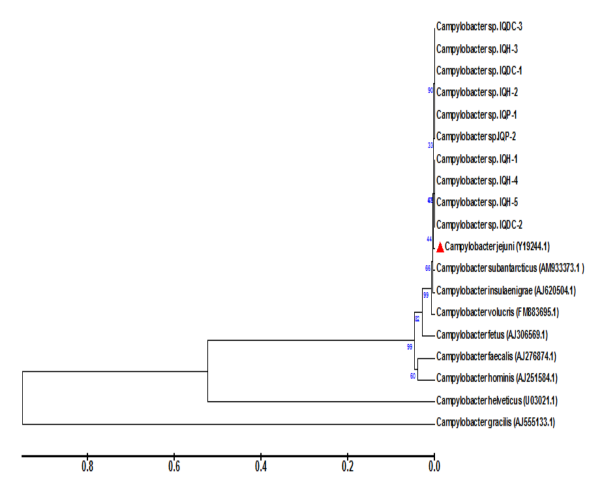


Figure 2. The phylogenetic tree analysis of *Campylobacter* spp. Iraq isolates based on partial 16SrRNA gene sequences using unweight pair group method with arithmetic mean (UPGMA tree) in (MEGA 6.0 version).

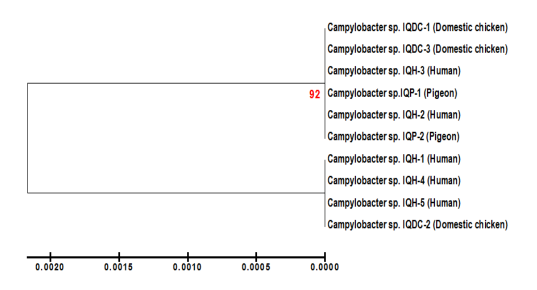


Figure 3. Phylogenetic tree analysis of *Campylobacter* spp. Iraq isolates based on the partial 16S ribosomal RNA gene sequence that used for host relationship analysis. The phylogenetic tree was constructed using Unweight Pair Group method with Arithmetic Mean (UPGMA tree) in (MEGA 6.0 version).

Campylobacter gastroenteritis.

Conflict of Interests

The author has not declared any conflict of interests.

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