Full Length Research Paper

Shiga toxigenic, enteroinvasive and enteropathogenic Escherichia coli in fish from experimental fish farm (Layo), Côte d’Ivoire

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Received 23 March, 2019; Accepted 24 May 2019

The objective of this study was to know the sanitary quality of fish coming from Layo farm. Twenty fishes (Oreochromis niloticus), were selected in four ponds and Escherichia coli were isolated in gills and viscera according to microbiological methods. One hundred and twenty strains of E. coli were isolated, and their virulence was performed by polymerase chain reaction (PCR) using specific primers. Eleven (11) strains (9.16%) including 7 strains of gills (11.66%) and 4 strains of viscera (6.66%) had virulence genes eae, Stx1, Stx2 or iai. Atypical Enteropathogenic E. coli (EPEC, eae+, lack of bfp) was isolated from gills (5%) and viscera (1.66%). Shiga toxigenic E. coli (STEC) with genes eae + Stx2, Stx1 and Stx2 were isolated in viscera (5%) and gills (3.33%). For Enteroinvasive E. coli (EIEC), iai gene was isolated in gills (3.33%) but no ipah gene. Enterotoxigenic E. coli (ETEC) with lt gene and Enteroaggregating E. coli (EAEC) with aggA gene were not detected in this study. This study revealed that some fish from Layo farm are carriers of virulent E. coli that can cause serious human diseases and can lead to consumer death if cooking is insufficient or by cross-contamination. This therefore poses a real public health problem.

Key words: Escherichia coli pathogens, fish, ponds, public health.

INTRODUCTION

Fish is one of the main sources of animal protein in human diet. In Côte d’Ivoire, production of fish ranges from 50,000 to 80,000 tons annually. This production covers 30% of the needs which went from 300 000 tons

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in 2005 to 850 000 tons in 2010. The deficit is filled by fish imports which represent 268 333 tons (67%) of national need for halieutic products (Coulibaly, 2010). To solve deficit problems, overfishing and extinction of some species, several aquaculture stations have emerged, including Layo. Layo aquaculture station located in Dabou department is an experimental site. Some ponds are fed directly by Ebrié lagoon or groundwater. Several studies have shown that waters of Ebrié lagoon are polluted because of human activities taking place around them (Kouassi et al., 1990; Adingra, 2007; Tuo et al., 2013). Recent work on waters in ponds of this experimental farm have revealed their strong contaminations with fecal coliforms and Vibrio (Toulé et al., 2017). But there is not enough information about sanitary quality of fish in these aquaculture sites. Relative information are about physico-chemical parameters and microbiological qualities of water. Also, this study was conducted to investigate the presence of *Escherichia coli* (indicator of faecal contamination) in fish *Oreochromis niloticus* in ponds at the Experimental Fish Farm (Layo) and to detect by PCR method, five pathogroups of *E. coli* like Enteropathogenic *E. coli* (EPEC), Shiga toxigenic *E. coli* (STEC), Enteroinvasive *E. coli* (EIEC), Enterotoxinogenic *E. coli* (ETEC) and Enteroaggregating *E. coli* (EAEC) causing diarrhea, haemolytic uremic syndrome in human.

### MATERIALS AND METHODS

#### Sampling sites

This study was conducted in fish farm Layo. This site was located between 05°13’50.9” N and 004°26’25.1” W, contained 18 ponds whose sizes varied from 200 to 900 m², 0.70 to 1.30 m of water depth and fed by Ebrié lagoon and groundwater. Four ponds, E5, E6, E11 and E13 were selected according to water sources (Table 1).

#### Water quality sampling and measurement

Water quality parameters like, pH, salinity, temperature, and dissolved oxygen were measured *in situ* using a multiparameter YSI 6920 V2.1S (USA). Water samples were taken approximately at 20 cm of depth using 1000 mL borosilicate bottles and stored in a cooler containing ice before transported to laboratory within 4 h. The determination of suspended solids (SS) was made according to the centrifugation method (Rodier et al., 2009). Nitrate (NO₃⁻) was measured by the cadmium reduction method (HACH method 8039) and the nitrite method (NO₂⁻) by the diazotisation method (HACH method 8507).

#### Fish sampling

Sampling was performed in April 2015. In four ponds selected, five fish of *O. niloticus* were taken per ponds, placed in individual labeled sterile polypropylene plastic bags, kept on ice and transported to laboratory within 4 h. Their body weight ranged from 300 to 500 g, and their length between 20 and 30 cm. In total, 20 gills and 20 viscera were analyzed.

### Table 1. Characteristics of ponds.

| Pond | Water source   | Depth (m) | Area (m²) |
|------|----------------|-----------|-----------|
| E5   | Groundwater    | 0.70-1.20 | 887.4     |
| E6   | Groundwater    | 0.70-1.20 | 922.2     |
| E11  | Groundwater    | 0.70-1.20 | 240.0     |
| E13  | Ebrie lagoon   | 0.70-1.20 | 864.0     |

#### Isolation and identification of *E. coli* in viscera and gills of fish

For analysis, 25 g of viscera and gills from each sample were added to 225 ml of sterile buffer peptone water contained in a sterile plastic stomacher bag and mixed well and incubated at 44°C for 24 h. After, 0.1 mL of solution was inoculated on desoxycholate agar (Becton Dickinson, GmbH) and Petri dishes were incubated at 44°C for 24 h. Red colonies were used as presumptive *E. coli*. Three colonies of *E. coli* per Petri dishes were purified and confirmed by positive indole, negative citrate and urea. *E. coli* strain of American Type Culture Collection 25922 (ATCC 25922) was used as control.

#### Detection of virulence genes by PCR

Detection of virulence genes (Table 2) were made in 120 strains of *E. coli* from viscera (60 strains) and gills (60 strains). DNA of each isolate was extracted according to boiling method. Three colonies of an overnight bacterial culture were taken and suspended in 500 μL of distilled water. DNA was purified according the modified method described by Ausubel et al. (1992). The mixture was stored at -20°C for 10 min and then boiled at 100°C for 10 min. 400 μL of phenol/chloroform mixture (24:1) were added to 150 μl of supernatant. The tube was vortexed for 2 min and centrifuged at 13000 rpm for 2 min. Top phase was recovered into a new tube and the lower phase was discarded. On the upper phase obtained, 1/10th volume of sodium acetate to 3 M and 200 μL of absolute ethanol were added and stored at -20°C for precipitation and the mixture was incubated for 1 h at -20°C. After incubation, they were centrifuged again at 13000 rpm for 20 min at +4°C and then supernatant is removed by flipping the tubes. 1 ml of 70% ethanol stored at -20°C was added to the pellet and tubes were centrifuged again at 13000 rpm for 5 min. Then the supernatant was removed by flipping the tubes. The pellet was dried at the heating block for 15 min at 95°C and 100 μL of nuclease-free water were added to each tube.

According to modified methods of multiplex PCR previously described by Dadie et al. (2014), eight genes were screened in this study.

#### Multiplex PCR for *Stx1, Stx2 and it genes detection*

The PCR was performed in a final volume reaction of 25 μL containing 8.25 μL nuclease-free water (Ambion), 5 μL PCR buffer (5X), 1.5 μL magnesium chloride (MgCl₂, 25 mM) (Promega Corporation, Madison, USA), 0.5 μL Deoxyxynucleotide Triphosphates (dNTPs, 10 mM), 0.75 μL of each primer (20 mM) (Table 2), 0.25 μL Go Tag®G2 Flexi DNA polymerase 5 U/μL (Promega Corporation, Madison, USA) and 5 μL of template DNA. For positive control of gene, *E. coli* strains previously known were used (Dadie et al., 2014). The amplification program used for *Stx1, Stx2 and it genes*, included an initial denaturation (94°C, 3 min), followed by 35 cycles each composed of initial denaturation (94°C, 30 s), primer annealing (57°C, 45 s) and extension (72°C, 30 s) and a final extension (72°C, 30 s). After these cycles, a final extension
Table 2. Genes, sequences, size of fragments and reference for each primer used in this study.

| Gene  | Sequences                                      | Size (pb) | References, Genbank/EMBL number |
|-------|------------------------------------------------|-----------|---------------------------------|
| eaeA  | eae f: 5'-CACACGAATAAACTGACTAAAATG-3'  
eae r: 5'-AAAAACGCTGACCCGCACCTAAAT-3' | 376       | AE005595                        |
| bfpA  | bfp f: 5'-TTCTTGGTGCTTGGCTTCTTTT-3'  
bfp r: 5'-TTTTTTGGTGTATTTGGTA-3' | 367       | Yatsuyanagi et al. (2002)       |
| ipah  | ipah f: 5'-TGAAAAAATCCTATGCGCTCT-3'  
ipah r: 5'-CCAGTCCGTAATTAATCTCT-3' | 423       | Luscher and Athwegg (1994)      |
| ial   | ial f: 5'-CTGGATGTATGGGTAGG-3'  
ial r: 5'-GGAGGCCAACAATTATTTCC-3' | 320       | Svenungsson et al. (2000)       |
| aggA  | aggA f: 5'-AGACTCTGGCCGAAAGACTGTATC-3'  
aggA r: 5'-ATGGCTGTCTGTAATAGAGA-3' | 194       | Schmidt et al. (1995)           |
| Stx1  | Stx1 f: 5'-GAAAGTCCGTGGGATTACG-3'  
Stx1 r: 5'-AGCGATGCAGCTATTAATAA-3' | 130       | AF461172                        |
| Stx2  | Stx2-5'-ACCCTTTTTCAGATTTTACATA-3'  
Stx2-5'-TACACAGGAGCAGTTCAGACGT-3' | 298       | AY143337                        |
| Lt    | Lt-5'-TCTCTATGTCATACGGAGC-3'  
Lt-5'-CCATACGTATTGCCGCAAT-3' | 322       | Frankel et al. (1989)           |

of 5 min at 72°C is realised.

**Multiplex PCR for invasive and adhesine genes (eae, bfpA, ial, aggA, ipah)**

This second PCR was performed in a final volume reaction of 25 μL containing 5.25 μL nuclease-free water (Ambion), 5 μL PCR buffer (5X), 1.5 μL magnesium chloride (MgCl₂, 25 mM) (Promega Corporation, Madison, USA), 0.5 μL Deoxynucleotide Triphosphates (dNTPs, 10 mM), 0.75 μL of each primer (20 μM) (Table 2), 0.25 μL Go Tag®G2 Flexi DNA polymerase 5 U/μL (Promega Corporation, Madison, USA) and 5 μL of template DNA.

For invasive and adhesine genes (eae, bfpA, ial, aggA, ipah), the program was an initial denaturation step at 94°C for 3 min, followed by 35 cycles of denaturation (94°C for 30 s), primer annealing (56°C for 20 s), and extension (72°C for 30 s), with a final extension at 72°C for 5 min. Primers used in these PCR were reported in Table 2. PCR amplification products were revealed on a gel Doc EZ® imager (Bio-Rad) after electrophoresis in 2% agarose gel containing Syber safe (Invitrogen).

**RESULTS AND DISCUSSION**

**Physico-chemical parameters in ponds**

Physico-chemical results are presented in Figure 1. Water temperature in four selected ponds ranged from 35 to 36°C. This variability may be the fact that Côte d'Ivoire has a tropical climate (Inza et al., 2009) and these high temperatures could be explained by solar radiation according to Lwamba et al. (2015). High temperature is also an important predisposing factor for the growth of harmful bacteria for fish in aquaculture (Zhang et al., 2016). Salinity in all ponds was low and ranged from 0.25
(E11) to 0.65\% (E5) certainly because they are fed by inland waters and Ebrié lagoon is under influence of Agneby River. Agneby River after having crossed its watershed, discharges its effluents in Ebrie lagoon (Toulé et al., 2017) and is the seat of a strong anthropic pressure (Kamagate et al., 2017). Suspended solids (SS) ranged from 44.2 (E13) to 57.2 mg L\(^{-1}\) (E6). These values are higher than those observed (7.14 mg L\(^{-1}\)) in floating cages in Ebrié lagoon at Jacqueville aquaculture station (Toulé et al., 2017). These high values in Layo ponds could be explained by the fact that they are closed ecosystems where water renewal is rare. In rearing structures, fish are fed by artificial food. As a result, SS could be attributed to the enormous amounts of organic matter produced from uneaten food and fish metabolite waste. The resuspension of SS during aquaculture activities (sexing, fishing, sorting, transfer, etc.), rain penetration and runoff of soil and plant particles to ponds and pens could result in increased levels in SS.

The smallest dissolved oxygen values were identified in ponds E5 (5.32 mg L\(^{-1}\)) and E6 (7.43 mg L\(^{-1}\)), while the highest values were found in ponds E11 (16.19 mg L\(^{-1}\)) and E13 (17.14 mg L\(^{-1}\)). The higher dissolved oxygen in pond E13 where suspended matter is the lowest could be due to the fact that it is directly under the influence of Ebrié lagoon. pH varies from 6.29 (E11) to 8.5 (E6).

**Virulence genes detected among E. coli isolated**

The results of the genetic analysis of *E. coli* isolates from fish samples are presented in Table 3. Of the 120 *E. coli* isolates, 11 strains (9.16%) were positive for virulence genes. In gills, the pathogenic strains of *E. coli* were more isolated with a frequency of 11.66% (7 strains) than in viscera with 6.66% (4 strains). EPEC represented by *eae* gene were identified in 5% (3 strains) and in 1.66% (1 strain) in gills and viscera strains, respectively. The lack of *bfpA* gene in the present study suggests that EPEC strains are probably atypical. This kind of EPEC (*eae*, *bfpA*, *Stx*) was isolated from acute diarrhea (Vieira et al., 2001), and would be predominant in strains from developed countries (Paciorek, 2002; Trabulsli et al., 1996). EPEC is the leading cause of childhood diarrhea in developing countries (Food and Drug Administration, 2012). It damages the epithelial cells of the small intestine by producing typical lesions (Kaper et al., 2004).

Shigatoxigenic gene *Stx1* was identified only in viscera for 1.66% (1 strain) while *Stx2* was identified in viscera at 3.33% (2 strains) and in gills at 3.33% (2 strains). The positive isolate for both *Stx2* and *eae*, characteristic of Enterohemoragique *E. coli* (EHEC) but included in STEC, was identified in gills at 1.66% and in viscera at 1.66% in 1 strain. The presence of Shigatoxigenic genes has also been identified in intestines of fish from ponds in northeast of Sao Paulo, Brazil (Ribeiro et al., 2016). The presence of these genes is predominant in patients with Hemolytic Uremic Syndrome. Although the detection of strains carrying both *eae* and *Stx* genes in aquaculture environments is low (Zschock et al., 2000; Iritno et al., 2005; Alagarsamy et al., 2009), it has been detected in the present work and in approximately 7.69% of intestinal fish strains from ponds in northeast of Sao Paulo (Ribeiro et al., 2016). Invasive gene *ial* was identified only in gills (3.33%) but *ipah* invasive gene was not detected in the present study. Regarding EIEC, previous studies revealed the rarity of this pathotype in environment or in food (Barbosa et al., 2014). It is more often isolated in fish from developing countries (Peng et al., 2009). This pathotype is also responsible for infantile diarrhea (Moreno et al., 2010; Nguyen et al., 2005). The genes encoding ETEC (*lt*, *st*) and EAEC (*agg A*) were not identified in this study (Table 3).

All pathogenic genes identified in this study are likely to cause diarrhea, dysentery, haemorrhagic colitis, haemolytic uremic syndrome and chronic post-infection sequelae in men (Ribeiro et al., 2016). Their detections in gills and viscera of fish could come from direct contact with water (Fouz et al., 2000). These results are also identified in the viscera of *O. niloticus* from aquaculture stations in Sao Paulo, Brazil (Ribeiro et al., 2016). In contrast to this work, pathogenic strains were not detected in fish caught in Aby Lagoon, Côte d’Ivoire by Kambire et al. (2017).

The majority (80%) of the genes sought, *Stx2*, *eae*, *ial*,
Table 3. Prevalence of virulence genes of Shigatoxigenic (Stx1, Stx2, eae+Stx2), Enteropathogenic (eae, bfpA) and Enteroinvasive (ial) Escherichia coli in gills and viscera of fish from Layo ponds, Abidjan, Côte d’Ivoire.

| Origin | E. coli isolates | Stx1 (%) | Stx2 (%) | Lt (%) | Eae (%) | Bfp A (%) | ial (%) | Agg A (%) | IpA H (%) | eae + Stx2 (%) | Total (%) |
|--------|------------------|----------|----------|--------|---------|-----------|---------|-----------|-----------|---------------|-----------|
| Gills  | 60               | 0 (0)    | 2 (3.33) | 0 (0)  | 3 (5)   | 0 (0)     | 2 (3.33)| 0 (0)     | 0 (0)     | 1 (1.66)      | 7 (11.66) |
| Gut    | 60               | 1 (1.66) | 2 (3.33) | 0 (0)  | 1 (1.66)| 0 (0)     | 0 (0)  | 0 (0)     | 0 (0)     | 1 (1.66)      | 4 (6.66)  |
| Total  | 120              | 1 (0.83) | 4 (3.33) | 0 (0)  | 3 (2.5) | 0 (0)     | 2 (1.66)| 0 (0)     | 0 (0)     | 2 (1.66)      | 11 (9.16) |

Figure 2. Prevalence of Enteropathogenic (eae), Shigatoxigenic (Stx1, Stx2, eae+Stx2) and Enteroinvasive (ial) E. coli in fish ponds.

and eae + Stx2, were detected in fish strains from E6 pond. The Stx2 gene was detected in E5, E6 and E11 ponds. The ial gene was present in strains from E13 and E6 ponds. At least 1 gene was detected per pond (Figure 2).

The aquaculture station of Layo is close to a village where there is no septic system and animals such as chicken, sheep, goats and other domesticated animals move around freely. Bovine faeces have been identified as the main reservoir of E. coli and are a vehicle of transmission to the environment, cattle and food (Wang et al., 1996).

Environmental conditions surrounding crop sites may affect the quality of water and farmed fish. The prevalence of these pathotypes reflects the bacterial compositions of the living environments (water and sediments) and the health status of the fish according to Pakingking et al. (2015). In aquaculture activities, few studies have been conducted on the presence of microorganisms responsible for human diarrhea. In addition, the presence of E. coli in environment is generally quantitatively assessed as an indicator of faecal contamination with respect to the quality of irrigation water, without considering that the presence of this microorganism, even at low concentration, indicates a risk of transmission of pathogens to humans. It is true that E. coli bacteria do not cause losses in aquaculture production, but it can cause human diseases. Therefore, fish farmers do not see the need to apply appropriate health control measures to ensure product quality. However, infected fish used as a food source can serve as means of transmission of these agents to humans, and even contaminate other surfaces.
Conclusion

This study revealed the presence of E. coli pathotypes like STEC, EPEC, EIEC in gills and viscera of O. niloticus in ponds of Layo fish farm. However, ETEC and EAEC pathotypes were not detected. Although these pathogenic bacteria do not cause losses in fish production, they cause serious human diseases that can lead to death. There is therefore a real public health problem that should be of concern and brought to the attention of the appropriate government authorities.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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