ABSTRACT:
The current study intended to isolate and identify *Shigella flexneri* via traditional bacterial isolation (TBI), PCR-sequencing identification (PSI), and phylogenetic tree analysis (PTA) from patients attended Al-Diwaniyah General Teaching Hospital, Al-Diwaniyah City, Iraq. A total of 25 stool samples were collected and subjected to the above mentioned techniques. The results of the TBI revealed the presence of *S. flexneri* in these samples. The PCR results confirmed the identification of these bacteria in these samples after a piece, 1256 bp, of the gyrB gene, encodes the subunit B protein of DNA gyrase, was targeted. For the sequencing of 2 samples, the results showed the complete confirmation revealing the correctness of the TBI and the PCR results in identifying these bacteria in the tested samples. The PTA revealed separate distinctive clustering of the sequenced samples in one branch of the phylogenetic tree. The current study enhances the documentation that *S. flexneri* is one of the infectious agents that causes GIT-related infections in the current tested city.

KEYWORDS: *Shigella flexneri*, traditional, isolation, PCR, phylogenetic tree,

1. INTRODUCTION

*Shigella flexneri* is a bacterium characterized by Gram-negative staining and its behaviour as a facultative anaerobe which is a member of the family *Enterobacteriaceae* (In et al., 2002; Wei et al., 2003). It belongs to a genus that has 4 species, which are *S. flexneri*, *S. boydii*, *S. sonnei*, and *S. dysenteriae*. Moreover, these members are classified according to their biochemical reactions and their O-antigen variations into serotypes leading to the identification of 13 serotypes for the *S. flexneri* alone (Jennison and Verma, 2004). The communicable disease, shigellosis, caused by these bacteria demonstrates itself with dysenteries in patients. The epithelial layers of the colon and rectum are the main targets for these bacteria leading to acute mucosal inflammation, severe tissue damage, abscess formation, ulceration, watery diarrhea, severe abdominal pain, and dysentery (Bennish, 1991; Taneja and Mewara, 2016). Lack of useful treatment may lead to the development of certain secondary health events such as septicemia, hemolytic uremic syndrome, pneumonia, rectal prolapse, arthralgia, seizures, toxic megacolon, intestinal perforation, electrolyte imbalance, and leukemoid reactions (Williams and Berkley, 2018). This disease is responsible for 1.1 million and over 164 million cases each year of mortalities and morbidities respectively; however, this disease attacks mostly children in countries with poor-health systems (Kotloff et al., 1999). The developing countries are known for the *S. flexneri* serotypes of 1b, 2a, 3a, 4a and 6; however, industrialized countries are known for the presence of 2a serotype in their patients (Kotloff et al., 1999). The disease is occurred via oral route and predisposed via contaminated water, poor hygiene, malnutrition, and lack of antibiotic useful treatment (Lee et al., 2017).

2. MATERIALS AND METHODS

2.1. Sampling

In the current study, 25 stool sample swabs were collected from GIT patients suffering from diarrhea and immediate cultivation in transporting media such as nutrient media. These samples were transferred to a microbiological Lab in the College of Veterinary Medicine, University of Al-Qadisiyah, Al-Diwaniyah, Iraq.

2.2. Traditional bacterial isolation

2.2.1. Cultivation

The samples were grown on Congo red tryptic soy agar plates (Payne and Finkelstein, 1977) that showed red-colored colonies. These colonies were inoculated in 50 ml of Pen assay, antibiotic medium 3, broth (Difco Laboratories, Detroit, Mich.) incubated at 37°C. After 4 hrs of cultivation, log-phase culture at 10ml was inoculated in 1L of Pen assay broth prewarmed to 37°C. This mixture was incubated overnight at 37°C in a shaking incubator (Turbyfill et al., 2000). MacConkey agar, xylose-lysine deoxycholate (XLD) agar, and Salmonella-Shigella (SS) agar were also used for this study.

2.2.2. Genomic DNA extraction

From the overnight bacterial broth culture, 1.5 ml were centrifuged to pellet the cells in an Eppendorf tube, DNA was extracted according to procedure of bacterial genomic extraction kit (Gene aid company, Taiwan).

2.2.3. PCR-based technique

The primers used in the current work were from (Yamamoto and Harayama, 1995), and they are F: GAGTCTCATATGACGGTTCTGCA and R: AGCAGGATCGATGTGCGAGCC that target the...
amplification of a piece in the gyrB gene at a length of 1256bp. The reactions involved the use of 100µl of total volume. The components were 5U of DNA polymerase, 50mM KCl, 10mM Tris-HCl (pH 8.3), 1.5mM MgCl₂, 0.001% (wt/vol) gelatin, 200 mM (each) of dNTPs, 10pM of each primer. The thermocycler conditions were 95°C for 5min as for the initial denaturation step (one cycle), 30 cycles of (96°C for 1min as for the main denaturation step, 63°C for 1min as for the annealing, and 72°C for 1min as for the extension), and 72°C for 7min as for one cycle for the final extension. Then, 3% agarose gel electrophoresis was used to run the PCR products.

2.2.4. Sequencing and Phylogenetic analysis.

The present work involved sequencing of 2 samples. The phylogenetic tree was generated using MegaX software and following A neighbor-joining analysis (Saitou and Nei, 1987; Tamura et al., 2013).

3. Results

3.1. Conventional detection of Shigella

The results of the TBI revealed the presence of S. flexneri in these samples. These bacteria appeared as red colonies on the Congo red tryptic soy agar plates. On the Kligler's Iron Agar and the Triple Sugar Iron Agar, the bacterial growth of the Shigella generated a gas-free alkaline slant and acid butt.

3.2. PCR-based results

The PCR results confirmed the identification of these bacteria in these samples after a piece, 1256 bp, of the gyrB gene, encodes the subunit B protein of DNA gyrase, was targeted, figure 1.

3.3. Sequencing results

For the sequencing of 2 samples, the results showed the complete confirmation revealing the correctness of the TBI and the PCR results in identifying these bacteria in the tested samples.

3.4. Phylogenetic-based results

The PTA revealed separate distinctive clustering of the sequenced samples in one branch of the phylogenetic tree. These isolates are registered in the NCBI-gene Bank under the numbers, MK242204.1 and MK342205.1, figure 2.

4. Discussion

Shigella flexneri is a bacterium characterized by Gram-negative staining and its behaviour as a facultative anaerobe which is a member of the family, Enterobacteriaceae. In et al., 2002; Wei et al., 2003). Failure to introduce sufficient management of the infection may add some detrimental incidents that might prevent required recovery within normal times or might lead to lose life (Williams and Berkley, 2018). The disease initiated by these bacteria is well known for its million-based morbidity and mortality occurrence especially in children in poor health system countries (Kotloff et al., 1999). The results in the current work revealed the presence of S. flexneri in the samples using the regular Lab techniques specific for the isolation of bacteria. These techniques are important in the isolation of bacteria providing vital primary detection before heading to downstream molecular techniques, and this agrees with (Faruque et al., 2002) who isolated these microorganisms from water samples using these primary techniques. The current work results also agree with (Jomezadeh et al., 2014) who isolated Shigella species from dysentery samples of children in Iran using the primary lab techniques. Using PCR-based techniques, the present study identified and confirmed the presence of S. flexneri in the samples tested for this purpose. These are important techniques to support the validity of the primary, TBI, methods. These results agree with (Vu et al., 2004) who used PCR in the detection of Shigella strains using ipaHas a target gene in Vietnam and suggested increasing prevalence of the disease in this country. The current work results also agree with (Ojha et al., 2013) who detected these bacteria using pentaplex PCR targeted 5 genes. Sequencing and the phylogenetic analyses revealed important identification and confirmation that supported the results obtained from the previous techniques used in the present study. These techniques reflect important information inferred by the phylogenetic analysis showing separate distinctive clustering of the sequenced samples in one branch of the phylogenetic tree, and this could indicate unique strains related to the current tested city (Fukushima et al., 2002). The current study enhances the documentation that S. flexneri is one of the infectious agents that causes GIT-related infections in the current tested city.
References


