

MOLECULAR CHARACTERIZATION AND ANTIBIOGRAM OF SELECTED PATHOGENS ISOLATED FROM STREAMS AND WELLS WATER SOURCES IN THREE LOCAL GOVERNMENT AREAS OF RIVERS STATE, NIGERIA

ABSTRACT

Aims: Reliance on unhealthy water sources such as hand dug wells and streams have led to the occurrence of waterborne diseases such as amoebic dysentery, gastroenteritis, typhoid fever and cholera among several communities in Rivers State. This study tends to identify selected multidrug resistant pathogens isolated from streams and wells water sources in three Local Government Areas of Rivers State using molecular techniques

Methodology: A total of 48 well and stream water samples were collected aseptically from 24 sampling sites in Ikwerre, Emohua and Etche Local Government Areas of Rivers State. Samples were analysed using membrane filtration technique for total heterotrophic bacterial count and total coliform count. The well and stream water samples were also cultured for *E. coli*, *Salmonella* and *Vibrio* sp. Antibiotic sensitivity test of the isolates was carried out using the disk diffusion method and multidrug resistant isolates were identified using PCR-based techniques (16S rRNA)

Results: The results of this study showed that total aerobic heterotrophic bacteria count occurred more in Emohua LGA for both well and stream water sources with bacterial count of 1.71×10^4 cfu/ml for well water samples and 2.35×10^4 cfu/ml for stream water samples. The mean fungal count of the stream water samples was highest in Emohua LGA with 2.55×10^4 cfu/ml followed by Ikwerre LGA which had 4.20×10^3 cfu/ml and Etche LGA with the least count of 1.80×10^2 cfu/ml. All the isolated bacteria species were found to be resistant to more than 50% of the antibiotics used with highest Multi Antibiotic Resistant Index of 0.625. multidrug resistant bacterial isolates identified include *Vibrio* sp strain 201707CJKOP-Y162 (MG593726), *Bacillus subtilis* clone N55 (JQ622582), *Chryseobacterium* sp strain CB2915-325-DE_0621 (MH512534), *Enterobacter* sp strain AB55 (MF407145), *Klebsiella pneumoniae* strain M792-16(MH680830), *Shigella sonnei* strain AR_0030(CP032523), *Escherichia coli* strain WECHEC025943(CP027205) and *Salmonella bongori* strain KC153129.116 (MG663494)

Conclusion: Access to good quality or potable drinking water and efficient sanitary practices are fundamental to human health and economic development. The occurrence of multidrug resistant pathogenic bacterial isolates in natural water sources requires serious routine evaluation in order to forestall the outbreak of waterborne disease epidemics.

Keywords: 16S rRNA; Public Health, Multidrug resistance; Stream and well water; Waterborne diseases

INTRODUCTION

Drinking water, defined as "suitable for human consumption", is free of disease-causing microorganisms. The possible consequences of microbial contamination for health are such that its control must be a primary objective and must never be compromised. Developing countries continue to struggle with the issue of food security, that is, the amount of food sufficient for consumption by the growing population; there is another dilemma in these countries. It is estimated that more than 200 types of diseases caused by pathogens are foodborne and waterborne causing problems in vulnerable groups of people such as, the elderly, pregnant women and young children. Therefore, ensuring the safety of food is an important challenge for public health [1].

Drinking water has been a major issue in many countries like Nigeria and majority of the rural populace in Nigeria do not have access to portable water [2]. Only few people can afford and rely on purified and treated water particularly for consumption therefore, underground water (well water) serve as

the major source of both drinking water and domestic water used in local population of Nigeria [3]. Majority of the disease-causing microorganisms transmitted via drinking water are predominantly faecal in origin and are commonly called enteric pathogens, Olatunji *et al.* [4]. With regards to the World Health Organization [5] standards, drinking water should not contain any microorganisms known to be pathogenic or any bacteria indicative of faecal pollution. The use of water contaminated with domestic and industrial waste, human and animal excreta leads to the development of waterborne diseases like gastroenteritis, typhoid fever, cholera, diarrhea and bacillary dysentery [6].

There is a growing public health interest relating to the uses of water and the outbreak of waterborne diseases. This concern is genuine and threatening considering the fact that water contaminated with pathogenic microorganisms when consumed or used in the preparation of food also leads foodborne disease. Rural communities in many developing countries obtain drinking water from untreated sources. In Nigeria for instance, water supply infrastructures are either still at the development stage or are totally absent in the rural communities [7]. According to a survey by Majuru *et al.* [8], about 6.5 million Nigerians have no access to safe source of water. The situation is made worse in the rural areas where only 24% of the population had access to safe water. So, the provision of clean, reliable and potable water in rural areas remains a major challenge considering the fact that majority of members of the population reside in the rural areas. When provision of **clean and safe water is inadequate, residents usually resort to use of contaminated water sources to certify their water needs** [9].

Most of the bacteria, viruses, parasites, and fungi that contaminate well and stream water comes from fecal material from humans and other animals, for example from on-site sanitation systems (such as pit latrines and septic tanks). Common bacterial contaminants include *E. coli*, *Salmonella*, *Shigella*, *Micrococcus* spp., *Pseudomonas* spp., *Streptococcus* spp., *Klebsiella* spp., and *Campylobacter jejuni*.

The use of antibiotics has helped a great deal in the treatment of waterborne diseases caused by the respective microorganisms. However, Stanley [10] noted that the major problem encountered in antibiotic treatment is the development of resistance by bacteria. The increasing use of antibiotics in food and animals leads to the wide spread occurrence of resistant enteric pathogens to humans. Gideon *et al.* [11] reported the antibiotic resistant profile of bacteria isolated from drinking water sources in Amai Kingdom, Delta State, Nigeria. Adekunle *et al.* [12] stated that the development of antibiotic resistance by enteric pathogens poses a great risk to citizens in areas where diarrhoea infection is problematic. Tolessa *et al.* [13] stated that bacteria are known to acquire antibiotic resistance through horizontal plasmids transfer mechanisms. Abui *et al.* [9] stated that in Nigeria and other many other developing nations, the apparent lack of potable water supplies **remains a major problem in most of the communities.**

The importance of the use of good quality water to satisfy the food, recreation, domestic and industrial needs of man have been emphasized in many reports including the United Nations Commission on Sustainable Development (2010) on water quality and WHO [5] guide line on drinking water quality. Although, numerous policies have been initiated and strategies adopted by different member countries of these world organisations, the problems of either unavailability, inadequate or poor quality sources of water supplies still abound and are confronting most cities and rural communities around the world. In Rivers State, some of the communities in the rural local government areas like Ikwerre, Emohua and Etche Local Government Areas are still shanties and largely underdeveloped. These communities are

very poor and basic social amenities like good quality source of water supply still remains a mirage. So, there is heavy dependence on hand-dug wells, shallow streams, rivers and creeks to provide for their water needs. Reliance on these unhealthy water sources have led to the occurrence of waterborne diseases such as amoebic dysentery, gastroenteritis, typhoid fever, cholera, mild/acute diarrhoea and bacillary dysentery among the residents of these local communities. This study tends to identify selected multidrug resistant pathogens isolated from streams and wells water sources in three Local Government Areas of Rivers State using molecular techniques.

MATERIALS AND METHODS

Study area and sampling sites

This research was carried out in selected villages in some communities from Ikwerre, Etche and Emohua Local Government Areas of Rivers State. Eight different sampling sites comprising of six hand-dug wells and 2 streams in each of the selected community from the three Local Government Areas were identified and sampled monthly for microbiological analysis. This made up a total sum of twenty-four sampling sites.

Collection of well and stream water samples

Six clean sterile glass bottles were used to collect well water samples from each of the six hand-dug wells in the community and transferred immediately into already labelled sterile two litre plastic bottle containers. Collection of stream water samples was done by entering into the water body up to the knee level and plunging the neck of the sterile glass bottle containers to about 30cm downward below the water to let the container fill with space left to allow for mixing. All the samples were collected in duplicate. Stream water samples collected with clean glass bottle containers were filtered immediately through a membrane of 0.45 μm .

Isolation of bacteria species from the samples

The APHA [14] membrane filter techniques was adopted. Using sterilized, the membrane filter disc (0.45 μm) was removed and placed on the surface of the porous support. Then, the funnel was placed and held infirm position on top of the porous support with clamps. 10ml of 10^{-2} serial dilution of water sample was transferred into the funnel and suction from vacuum pump was applied. The vacuum pump was terminated after the sample has passed through the membrane filter, and the funnel removed. The membrane filter disc was carefully removed using sterilized forceps and transferred on to the surface of freshly prepared Eosin Methylene Blue (EMB) Agar, *Salmonella Shigella* agar and Thiosulphate Citrate Bile Salts Sucrose (TCBS) agar plates. All the plates were incubated at 37°C for 24 hours

Purification and maintenance of isolates

Each discrete colony on a petri dish was transferred using a sterile inoculating wire loop into plates containing freshly prepared Nutrient agar and incubated at 37°C for 24-48hrs respectively. After incubation, the cultural characteristics of the isolates were recorded and compared with descriptive features contained in Holt *et al.* [15]. The isolates were then preserved on nutrient agar slants stored in the refrigerator at 4°C for biochemical characterization and identification.

Biochemical characterization and identification of isolates

The methods of Oranusi *et al.* [16] was employed for the identification of the bacteria isolates. The biochemical tests that were used to further characterize bacteria are: catalase, methyl-red, oxidase,

citrate utilization, and coagulase and indole tests. The identities of coliforms and other bacteria were then confirmed using the identification aid outlined in Bergey's Manual for Determinative Bacteriology [15] as well as that of known taxa as described by Cheesbrough [17].

Molecular Identification

DNA extraction (Boiling method)

Five milliliters of an overnight broth culture of the selected bacterial isolate in Luria Bertani (LB) was spun at 14000rpm for 3 min. The cells were re-suspended in 500ul of normal saline and heated at 95°C for 20 min. The heated bacterial suspension was cooled on ice and spun for 3 min at 14000rpm. The supernatant containing the DNA was transferred to a 1.5ml microcentrifuge tube and stored at -20°C for other downstream reactions.

DNA Quantification

The extracted genomic DNA was quantified using the Nanodrop 1000 spectrophotometer. The software of the equipment was launched by double clicking on the Nanodrop icon. The equipment was initialized with 2 ul of sterile distilled water and blanked using normal saline. Two microlitre of the extracted DNA was loaded onto the lower pedestal, the upper pedestal was brought down to contact the extracted DNA on the lower pedestal. The DNA concentration was measured by clicking on the "measure" button.

16S rRNA Amplification

The 16s rRNA region of the rRNA genes of the isolates were amplified using the 27F: 5'-AGAGTTTGATCMTGGCTCAG-3' and 1492R: 5'-CGGTTACCTGTTACGACTT-3' primers on a ABI 9700 Applied Biosystems thermal cycler at a final volume of 25 microlitres for 35 cycles. The PCR mix included: the X2 Dream taq Master mix supplied by Inqaba, South Africa (taq polymerase, DNTPs, MgCl), the primers at a concentration of 0.4M and the extracted DNA as template. The PCR conditions were as follows: Initial denaturation, 95°C for 5 minutes; denaturation, 95°C for 30 seconds; annealing, 52°C for 30 seconds; extension, 72°C for 30 seconds for 35 cycles and final extension, 72°C for 5 minutes. The product was resolved on a 1% agarose gel at 120V for 20 minutes and visualized on a blue light transilluminator.

Sequencing

Sequencing was done using the BigDye Terminator kit on on a 3510 ABI sequencer by Inqaba Biotechnological, Pretoria South Africa. The sequencing was done at a final volume of 10ul, the components included 0.25 ul BigDye® terminator v1.1/v3.1, 2.25ul of 5 x BigDye sequencing buffer, 10uM Primer PCR primer, and 2-10ng PCR template per 100bp. The sequencing condition were as follows 32 cycles of 96°C for 10s, 55°C for 5s and 60°C for 4min.

Phylogenetic Analysis

Obtained sequences were edited using the bioinformatics algorithm Trace edit, similar sequences were downloaded from the National Centre for Biotechnology Information (NCBI) data base using BLASTN. These sequences were aligned using MAFFT. The evolutionary history was inferred using the Neighbor-Joining method in MEGA 6.0 [18]. The bootstrap consensus tree inferred from 500 replicates [19] is taken to represent the evolutionary history of the taxa analyzed. The evolutionary distances were computed using the Jukes-Cantor method [20].

Antibiogram Profile Test

Antibiotic sensitivity test of the isolates was carried out using the disk diffusion method described by Bauer *et al.* [21]. The Sensitive Test Agar (LAB M) medium was used. The medium was prepared according to manufacturer's instruction. The medium was allowed to cool to 50°C and poured into sterile petri dish plates and allowed to solidify. When set, the agar plates were dried for 30 minutes at 35°C by placing them upright in the incubator with the lid tilted. The inoculums of each isolate prepared was from an 18hour broth culture adjusted to obtain turbidity comparable to 0.5 McFarland Standard. 0.5ml of 1.175% (w/v) barium chloride dehydrate (BaCl₂ · 2H₂O) solution was added to 99.5ml of 1% tetraoxosulphate (IV) (H₂SO₄) acid to obtain the turbidity standard and used to standardize the bacteria inoculums. 18hours broth culture diluted with sterile saline was used to make adjustments. Sterile cotton tipped applicator was dipped into standardized bacterial solution. The swab was then used to streak the entire dried surface of the medium. The inoculated plates were incubated for 5 minutes to remove excess moisture.

The antibiotic disc (Abtek Biological Limited, Liverpool) was then placed at equidistance from each other on the plate with the aid of a pair of sterile forceps. The discs were pressed firmly onto the agar medium with the sterile forceps to ensure adequate contact with the agar. The inoculated plates were incubated for 18-24 hours at 35°C. The antibiotics discs used for the test are; oflaxcin (5µ), gentamicin (10µ), nalixidic (30µg), nitrofurantoin (200µg), cotrimazole (25µ), amoxicillin (25µg), tetracycline (25µg) and augumentin (30µg). A ruler was used to measure the diameter of each zone of inhibition and recorded in millimeters (mm). The measurements were compared to a zone size interpretative chart [22], and the organisms were grouped as resistant, intermediate or susceptible to the test antibiotics according to the chart.

Multi Antibiotic Resistant Index (MARI)

This was carried out as described by Matyar *et al.* [23] with slight modification. MARI= resistant antibiotics ÷ total antibiotics tested. MARI value > 0.2 indicate existence of isolate(s) from high risk contaminated source with frequent use of antibiotic(s) while values ≤0.2 show bacteria from source with less antibiotics usage [24].

Results

The changes in the counts of the total heterotrophic bacterial population of well and stream water samples are presented in Table 1. The total heterotrophic bacterial counts ranged from 1.96 x 10² cfu/ml to 2.90 x 10³ cfu/ml for Ikwerre LGA, and 2.75 x 10² cfu/ml in to 2.35 x 10⁴ cfu/ml for Emohua LGA. While Etche had 1.93 x 10² cfu/ml to 3.57 x 10² cfu/ml. Total aerobic heterotrophic bacteria count occurred more in Emohua LGA for both well and stream water sources with bacterial count of 1.71 x 10⁴ cfu/ml for well water samples and 2.35 x 10⁴ cfu/ml for stream water samples. The mean fungal count of the stream water samples was highest in Emohua LGA with 2.55 x 10⁴ cfu/ml followed by Ikwerre LGA which had 4.20 x 10³ cfu/ml and Etche LGA with the least count of 1.80 x 10² cfu/ml. However, the highest fungal load of 2.25 x 10⁴ cfu/ml was obtained from well water source from Emohua LGA. Table 1 shows the mean total coliform count from well and stream water sources in Ikwerre, Emohua and Etche LGA's. the highest coliform count was recorded from well water sample obtained from Ikwerre LGA with coliform counts of 52 cfu/100ml while the least count was observed from stream water samples obtained from Emohua LGA with coliform counts of 1 cfu/100ml. From preliminary identification of the isolates, seven

genera of bacteria which include *Escherichia coli*, *Enterobacter* sp., *Salmonella* sp., *Shigella* sp., *Bacillus* sp., *Klebsiella* sp. and *Vibrio* sp were isolated from both the stream and well waters sources as presented in Table 2.

Table 3 shows the antibiogram profiles of the different test pathogens with selected types of antibiotics. While *E. coli* showed very high sensitivity to Ofloxacin (OFL), Nitrofurantoin and Nalidixic acid (NAL), it was totally resistant to Amoxicillin (AMX) and intermediate to Augmentin (AUG). On the other, *Salmonella* sp showed total resistance to Augmentin (AUG), Tetracycline (TET) and Amoxicillin (AMX) and high sensitivity to NAL, Ofloxacin, Nitrofurantoin and Gentamicin antibiotics. *Vibrio* sp were very resistant to Augmentin, Amoxicillin and Cotrimazole. It was highly sensitive to Nitrofurantoin, Nalidixic acid Ofloxacin, Gentamicin and intermediate to Tetracycline.

The identification of the selected multidrug resistant bacteria isolates to determine the strain was carried out using molecular techniques. Table 4 below presents both the quantity and purity of genomic DNA that was extracted from the selected bacterial isolates for genomic identification. The PCR reaction of the extracted genomic DNA samples revealed positive amplification of the 16S rRNA gene in all the multidrug resistant isolates when ran on 1.5% agarose gel at 120V for 20 min as showed in Figure 1. Clear bands observed in lane 1, 3-13 shows positive amplification of the extracted DNA samples with Lane N representing a 1000 base pair molecular ladder.

The obtained 16s rRNA sequence from the isolate produced an exact match during the megablast search for highly similar sequences from the NCBI non-redundant nucleotide (nr/nt) database. The 16S rRNA of the isolate B8 showed a percentage similarity to other species at 99%. The evolutionary distances computed using the Jukes-Cantor method were in agreement with the phylogenetic placement of the 16S rRNA of the isolate within the *Salmonella* sp revealed a closely relatedness to *Salmonella bongori* strain KC153129.116(MG663494) than other *Salmonella* sp respectively. The other organisms identified are; *Vibrio* sp strain 201707CJKOP-Y162(MG593726), *Bacillus subtilis* clone N55(JQ622582), *Chryseobacterium* sp strain CB2915-325-DE_0621(MH512534), *Enterobacter* sp strain AB55(MF407145), *Klebsiella pneumonia* strain M792-16(MH680830), *Shigella sonnei* strain AR_0030(CP032523), *Escherichia coli* strain WECHEC025943(CP027205), according to the phylogenetic tree on Figure 1.

Table 1

Mean microbial count from streams and wells water samples in Ikwerre, Emohua, and Etche LGA's of Rivers State

| Water sources | Microbial load in Ikwerre LGA (CFU/ml) | | | Microbial load in Etche LGA (CFU/ml) | | | Microbial load in Emohua LGA (CFU/ml) | | |
|---------------------|--|-----------------|--------------------|--------------------------------------|-----------------|--------------------|---------------------------------------|-----------------|--------------------|
| | THBC | TCC (cfu/100ml) | TFC | THBC | TCC (cfu/100ml) | TFC | THBC | TCC (cfu/100ml) | TFC |
| Well ₁ | 2.68×10^2 | 43 | 2.58×10^2 | 3.57×10^2 | 25 | 1.52×10^2 | 3.10×10^3 | 20 | 1.25×10^2 |
| Well ₂ | 1.96×10^2 | 50 | 2.17×10^3 | 1.93×10^2 | 20 | 2.17×10^2 | 1.71×10^4 | 28 | 1.82×10^3 |
| Well ₃ | 1.34×10^3 | 52 | 3.48×10^3 | 2.33×10^2 | 22 | 2.67×10^2 | 2.53×10^3 | 37 | 2.25×10^4 |
| Stream ₁ | 3.30×10^2 | 11 | 2.63×10^2 | 2.64×10^2 | 6 | 2.96×10^2 | 3.69×10^3 | 1 | 2.55×10^4 |
| Stream ₂ | 3.88×10^2 | 3 | 4.20×10^3 | 2.25×10^2 | 3 | 2.48×10^2 | 2.75×10^2 | 9 | 1.38×10^3 |
| Stream ₃ | 2.90×10^3 | 8 | 1.89×10^3 | 3.85×10^2 | 2 | 1.80×10^2 | 2.35×10^4 | 6 | 2.76×10^3 |

Table 2

Biochemical characterization and identification of bacterial isolates from streams and well water samples in Ikwerre, Emohua, and Etche LGA's of Rivers State

| Number of isolates | Gram's reaction | Cell morphology and arrangement | Catalase | Motility | Oxidase | Coagulase | Indole | Citrate | H ₂ S | MR | VP | Nitrate | Urease | Glucose | Lactose | Mannose | Sucrose | Mannitol | Maltose | Ribose | Probable organism |
|--------------------|-----------------|---------------------------------|----------|----------|---------|-----------|--------|---------|------------------|----|----|---------|--------|---------|---------|---------|---------|----------|---------|--------|-------------------------|
| | | | | | | | | | | | | | | | | | | | | | |
| 7 | - | Rods in clusters | + | + | - | - | - | + | - | - | + | + | - | + | - | + | + | + | + | + | <i>Enterobacter</i> sp |
| 9 | - | Short rods in pairs and | + | - | - | - | ± | - | - | + | - | + | - | ± | - | + | - | + | ± | | <i>Shigella</i> sp |
| 12 | - | Rods in pairs | + | + | - | - | - | + | + | - | + | - | + | - | + | - | + | + | + | | <i>Salmonella</i> sp |
| 21 | - | Rods in pairs | + | + | - | - | + | - | - | + | - | + | - | + | + | - | ± | + | - | - | <i>Escherichia coli</i> |
| 8 | - | Rods in singles | + | - | - | - | - | + | - | - | + | + | + | + | + | + | + | + | + | - | <i>Klebsiella</i> sp |
| 13 | - | Curved Rods | + | + | + | - | + | + | - | - | ± | - | - | + | ± | + | + | + | + | - | <i>Vibrio</i> sp |
| 10 | + | Long Rods in singles | + | + | ± | - | - | + | ± | - | + | + | - | + | ± | + | + | + | + | + | <i>Bacillus</i> sp |

UNDER PEER REVIEW

Table 3
Antibiogram of *E. coli* isolates in water samples from Emohua Local Government Area, Rivers State.

| Isolate No. | Zone of Inhibition (mm) | | | | | | | | | | | |
|-------------------------|-------------------------|-----|-----|-----|-----|-----|-----|-----|----------|----------|----------|-------|
| | AUG | TET | AMX | COT | OFL | GEN | NAL | NIT | R (%) | I (%) | S (%) | MARI |
| <i>Enterobacter</i> sp | 15I | 13R | 13R | 12I | 21S | 14I | 30S | 20S | 2(25) | 3(37.5) | 3(37.5) | 0.25 |
| <i>Shigella</i> sp | 16I | 17I | 11R | 21S | 19S | 13I | 29S | 18S | 1(12.5) | 3(37.5) | 3(37.5) | 0.125 |
| <i>Salmonella</i> sp | 16I | 18I | 11R | 20S | 23S | 11R | 16I | 19S | 2(25) | 3(37.5) | 3(37.5) | 0.25 |
| <i>Escherichia coli</i> | 12R | 16I | 12R | OR | 20S | 15S | 22S | 12R | 3 (37.5) | 1 (12.5) | 1 (12.5) | 0.375 |
| <i>Klebsiella</i> sp | 17I | 16I | 13R | 14I | 18S | 16S | 20S | 18S | 1 (12.5) | 3 (37.5) | 4 (50) | 0.125 |
| <i>Vibrio</i> sp 1 | 11R | 18I | OR | 15I | 22S | 13R | 10R | 19S | 4 (50) | 2 (25) | 2 (25) | 0.5 |
| <i>Enterobacter</i> sp | 12R | 13R | OR | 16I | 19S | 17S | 12R | 10R | 5 (62.5) | 1 (12.5) | 2 (25) | 0.625 |
| <i>Shigella</i> sp | 13R | 15I | 11R | 18S | 18S | 22S | 13R | 24S | 3 (37.5) | 1 (12.5) | 4 (50) | 0.375 |
| <i>Salmonella</i> sp | 11R | 11R | 8R | 19S | 20S | 14I | 10R | 18S | 4 (50) | 1 (12.5) | 3 (37.5) | 0.5 |
| <i>Escherichia coli</i> | 12R | 13R | 15I | 17S | 16I | 11R | 15I | 11R | 4 (50) | 3 (37.5) | 1 (12.5) | 0.5 |
| <i>Klebsiella</i> sp | 15I | 13R | 13R | 10R | 17S | 21S | OR | 13R | 5 (62.5) | 1 (12.5) | 3 (37.5) | 0.625 |
| <i>Bacillus</i> sp | 12R | 15I | 10R | 12R | 15I | 19S | 12R | 16I | 4 (50) | 3 (37.5) | 1 (12.5) | 0.5 |
| <i>Vibrio</i> sp 2 | 12R | 16I | 15I | 16I | 21S | 21S | 18S | OR | 2 (25) | 3 (37.5) | 3 (37.5) | 0.25 |

Key:

OFL- Ofloxacin Gen-Gentamicin Nal-Nalidixic Acid NIT-Nitrofurantoin
Cot-Cotrimazole Aug- Augumentin Amx Amoxicillin TET – Tetracycline
R = Resistant, I= Immediate, S = Sensitive
WV: *Vibrio* species in well (no.1,2,5& 6) water samples
SV: *Vibrio* species in stream (no. 7 & 8) water samples

Table 4
Nanodrop quantified bacterial DNA

| S/N | Isolate code | Quantity (ng/μl) of bacterial DNA |
|-----|--------------|-----------------------------------|
| 1 | B1 | 12.7 |
| 2 | B2 | 21.0 |
| 3 | B3 | 8.2 |
| 4 | B4 | 15.9 |
| 5 | B5 | 5.6 |
| 6 | B6 | 13.9 |
| 7 | B7 | 6.4 |
| 8 | B8 | 18.6 |

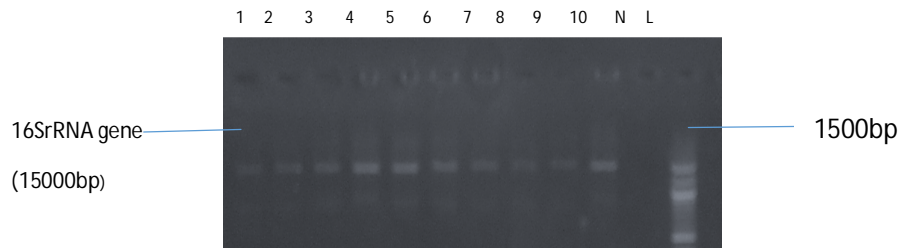


Plate 1: Agarose gel electrophoresis of the 16S rRNA gene of some selected bacterial isolates. Lanes 1, 3-12 represent the 16SrRNA gene bands (1500bp), Lane 2, failed amplification, Lane N represents the negative control, lane L represents the 1000bp molecular ladder.

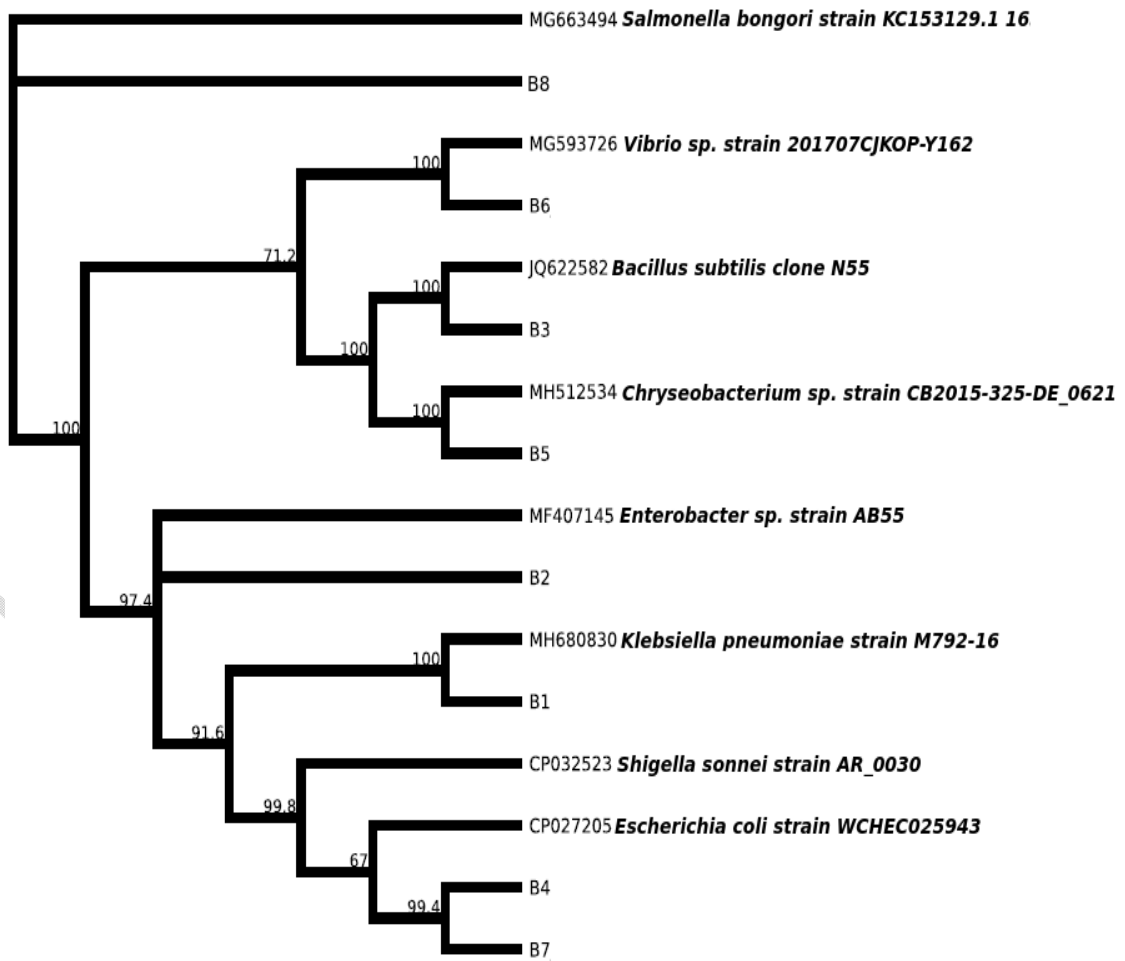


FIG.1: Phylogenetic tree showing the evolutionary distance between the bacterial isolates.

Table 5
Identification of the isolates by 16S rRNA genes amplification showing percentage similarities

| S/N | Gene Bank isolates | Accession number | Bacteriocin positive isolates | % Similarity |
|-----|---|------------------|-------------------------------|--------------|
| 1 | <i>Klebsiella pneumonia</i> strain M792-16 | MH680830 | B1 | 100 |
| 2 | <i>Enterobacter</i> sp. strain AB55 | MF407145 | B2 | 97.4 |
| 3 | <i>Bacillus subtilis</i> clone N55 | JQ622582 | B3 | 100 |
| 4 | <i>Escherichia coli</i> strain WCHEC025943 | CP027205 | B4 | 99.4 |
| 5 | <i>Chryseobacterium</i> sp. strain CB2015-325-DE_0621 | MH512534 | B5 | 100 |
| 6 | <i>Vibrio</i> sp. strain 201707CJKOP-Y162 | JQ622582 | B6 | 100 |
| 7 | <i>Shigella sonnei</i> strain AR_0030 | CP032523 | B7 | 99.8 |
| 8 | <i>Salmonella bongori</i> strain KC153129.1 16 | MG663494 | B8 | 100 |

Discussion

The results obtained were compared with both national and international standards. The examination of microbiological quality of drinking water sources is intended to prevent the development of waterborne illnesses or outbreaks among the rural dwellers as a result of consumption of water contaminated with harmful microorganisms [25]. Water or food items to be consumed or water that is designated for drinking should not harbour pathogenic microorganisms or any bacteria indicative of faecal contamination. The results of the mean total aerobic heterotrophic bacterial counts of the wells and stream water samples from the three Local Government Areas were found to be lower than the one reported by Anyanwu and Okoli [26] in a study of well water samples from different locations in Nsukka, Southeast Nigeria. However, Olatunji *et al.* [27] and Agwaranze *et al.* [25] in their assessment of the water quality of Asa river and well water sources in Wukari, Taraba State, Nigeria obtained a lower heterotrophic bacterial counts when compared to the present study. Although, the heterotrophic bacterial count observed in the well waters examined were high, the values were all lower than the one obtained by Shittu *et al.* [28] in their study of the Rivers and well waters in Kuta Town, Ogun State, Nigeria. Results from the total coliform bacterial count of the wells and stream water sources in this study revealed the presence of coliform bacteria in many of the wells and streams across the three LGA's. The result of this study is in line with the findings of Yahaya *et al.* [29] who reported that the contamination of water sources by coliform can be high during raining season than the dry season. This is evident when the water sources are not shielded from surface run-off during the wet season. Moreso, the grazing of domestic animals which are seen freely roaming about in the various communities of the three LGA could have also added to the pollution of the water sources.

The result of the biochemical test indicated the presence of *E. coli*, *Klebsiella* sp, *Enterobacter aerogenes*, *Staphylococcus aureus*, *Pseudomonas aeruginosa*, *Salmonella* sp, *Vibrio* sp, *Shigella* sp, and *Bacillus* sp. Romulus *et al.* [30] also isolated *Salmonella*, *Escherichia coli*, *Vibrio*, *Enterobacter*, *Klebsiella* and *Pseudomonas* from the shallow wells in Kitui Town, Kenya. Okunye and Odeleye [31] isolated and

identified *Vibrio* sp in their bacteriological investigation of well water samples from selected market locations in Ibadan, Nigeria.

In addition to the infections caused by these organisms, another major problem associated with these pathogens is the development of drug resistance. Saif *et al.* [32] stated that antibiotic resistant bacteria can infect human via contaminated food and drinking water, or directly from the environment. According to Samantha and Janeas [33], these superbugs result in infections that are only responsive to treatment with a few if any currently available antimicrobial agent. What is worrisome, is that these resistant organisms cause waterborne infections that leads to high rate of morbidity and mortality among residents of affected communities. There is therefore, a huge healthcare cost usually expended on waterborne diseases caused by resistance pathogens.

The contamination of water sources used for household chores and drinking purposes with harmful microorganism account for a major health problem, especially among the low income group and the rural dwellers in developing countries. Microbiological analysis of the well and stream water sources for the various communities in the three local government areas investigated showed a slightly high values for bacteria and fungi count. The biochemical characterization and identification of the isolates revealed the presence of potential waterborne pathogens, such as *E. coli*, *Salmonella* sp, *Enterobacter* sp., *Shigella* sp., *Vibrio* sp *Bacillus* sp. and *Klebsiella* sp that are hazardous to human health when consumed or ingested.

CONCLUSION

Access to good quality or potable drinking water and efficient sanitary practices are fundamental to human health and economic development. The occurrence of multidrug resistant pathogenic bacterial isolates in natural water sources requires serious routine evaluation in order to forestall the outbreak of waterborne disease epidemics. Although, some progress has been made by both the local, state and federal authorities towards the provision of potable water to prevent the occurrence of waterborne disease, a lot still remains to be done to totally eradicate this deadly menace.

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