

# Microbiological and Physicochemical Analysis of some Selected Municipal dumpsites in Calabar Municipality, Cross River State

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**Abstract:** *The study was aimed at investigating the microbiological and physicochemical analysis of some selected municipal dumpsites. Municipal wastes its leachate, soil and air samples within the dumpsites were randomly sampled (during wet and dry seasons) within Calabar metropolis. The research was undertaken within a period of six months. Standard microbiological methods were used to isolate, characterized and identify both bacteria and fungi isolates from the collected samples. a leachate collection system constructed using four-sided polyethylene water container attached to bottle with perforated holes on one side, and pushed under (perforated sides up) the waste hip at various locations was used for leachate collection. The result of the dump site samples collected showed that the total bacterial count of the dump site sample was generally higher in the wet season than dry season. In the dry season, there were no significant differences ( $p < 0.05$ ) between the waste soil bacterial count, waste soil fungal count, waste fungal count and dump site air fungal count. For the wet season, there was no significant differences in the waste-soil bacterial count and waste bacteria count but a significant difference in the waste fungal count and waste soil fungal count was observed. Bacteria isolates from the waste samples were identified as *Bacillus* spp, and *Klebsiella* sp, *Escherichia coli*, *Enterobacter* sp, *Salmonella* sp, *Staphylococcus* sp, *Shihella* sp and *Pseudomonas* sp, while the fungi isolates were identified as *Aspergillus* sp, *Mucor* sp, *Nocardia* sp, and also the filamentous bacteria-*Actinomyces* sp was identified. The result of the physicochemical properties of the waste samples revealed the presence of trace and major elements such as potassium (878.511mg/L) Iron (50.0mg/L), Magnesium (23.25mg/L), Calcium (21.851mg/L), Manganese (4.45mg/L), Cobalt (1.30mg/L), Copper (0.50mg/L), with sodium (28600mg/L) having the highest content and cadmium (0.157mg/L). Apart from the temperature and BOD, all other values obtained from the physicochemical analysis of the waste samples were lower than the WHO permissible limits. However, from this research study, the garden street dump site has been proven to be a*

*considerable source of microbiological contamination and infection in Calabar municipality, as the examined waste samples from the aforementioned site contained considerable quantities of potentially pathogenic microorganisms. Therefore, its extended influences on the air, as well as the soil, plant and surrounding ground water should be given due considerations by government and other environmental agencies in the state, by providing a well constructed and operated land fill sites for municipal waste disposal, so as to safe guard both the public and environmental health from future potential hazards.*

## INTRODUCTION

Solid waste management has remained an undisputable environmental problem in the developing countries of the world and it stands out amongst the arrays of global environmental hazards besieging metropolitan cities (1). This problem has become increasingly complex due to the increase in human population, industrial and technological revolutions, in addition to the fact that the processes that control the fate of wastes in the receiving media are complex (2). Solid waste are any non-fluidic/non-flowing substance which has been identified to be of no use or has no immediate economic demand at a particular point or source either as a raw material, end product, expired products, containers or after use remnants and which must be disposed of (3), (4). They are generated from various human activities such as domestic, hospital, industrial and agricultural activities. According to (5),(6), it may be categorized according to its origin (domestic, industrial, commercial, construction or institutional), according to its contents (organic material glass, metal, plastic paper etc) or according to hazard potential (toxic, non-toxin, flammable, radioactive, infectious etc). landfills and open dumping remains the major method of disposing solid waste in Nigerian cities, and solid waste disposed in landfills is usually subjected to series of complex biochemical and physical processes, which leads to the production of both leachate and gaseous emissions (7) while municipal

waste dumping sites as an alternative, are designate places where waste from various sources are deposited (8). These sites are not properly constructed nor designed, and consequently wastes dumped there over the years biodegrade and generate leachates that ultimately become point source of pollution into soil and ground water (9) when precipitation occurs, percolating water (leachates) dissolves many organic and inorganic salts with may be transported to nearby aquifers resulting in the alternation of the water quality (10). The rate of production and characteristics of leachate produced depends on a number of factors which include but not limited to solid waste composition, particle size, degree of compaction, hydrology of site, age of land fill, moisture, temperature conditions and availability of oxygen (11). The implication of the dumpsite on groundwater hydrology is that leachates from the dumpsite infiltrates into the ground and also move in the direction of groundwater flow thereby contaminating the groundwater along its path (12), consumption leads to several health challenging and it has been observed that bad quality water issue such as typhoid, kidney disease, liver )and environmental health related problems. However, various searches have shown that the indiscriminate handling and disposal of wastes, leads to environmental degradation, destruction of ecosystem and noses great risks to public health (13). It is on this basis that this research study is focused on analyzing the microbiological and physiochemical properties of some selected municipal dumpsites in Calabar Municipality, Cross River State.

## MATERIALS AND METHODS

### The study area,

The study area was Calabar Municipality (fig. 1), is located between latitude 4<sup>0</sup>13' and 5<sup>0</sup>15' and longitudes 8<sup>0</sup>15' and 8<sup>0</sup>25' in Nigeria. The area is characterized by the wet and dry seasons with high annual rainfall in the range of 350-400mm and run-off estimated to reach 90% (CRBDA, 1982).

### Materials used

The materials used for this work includes; petri-dishes, test tubes, conical flasks, pipettes, slides, cover slips, filter paper, masking tape, aluminum foil, polyethylene bottles, and McCartney bottles.

### Culture media

The following culture media were used for the isolation, identification and characterization of microorganisms from the samples, nutrient agar (NA), MacConkey ager (MCA), Sabouraud pledextrose agar (SAD), Triple Sugar Iron (TSI) agar and Citrate agar. All the media were products

of Diagnostics laboratory, USA, and they were all prepared in accordance to the manufacture's instruction.

### Chemicals and reagents

For the identification and characterization of the isolates, chemicals used include; crystal violet, 95% ethanol, Lugol's iodine, Safranin, Lactophenol in cotton blue, Kovac's reagent, hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), Methyl red (MR) vogues proskauer reagents d-Napthnol solution, potassium hydroxide (KOH) and methylated spirit.

### Sample Collection

Leachate collection system was constructed using four- sided polyethylene water containers. Holes were perforated on one side of the bottle. These bottles were pushed (perforated sides up) under the waste heap at various locations with the stopper cover sticking out for easy identification. Each bottle was taken as leachate collection chamber. Sterile conical flasks well stoppered with cotton wool and aluminum foil were used for collection of leachate and borehole water samples. the samples were labeled and stored under refrigeration temperature until use.

### Soil Samples

A garden rake was used to remove the waste so as to expose the soil under it. Soil samples were then collected with hand trowel into aluminum foil and labeled.

### Other samples

The decomposing waste, and plant leaves were each aseptically collected into sterile disposable petri-dishes sealed with masking tape and properly labeled. All samples were stored in the refrigerator before use.

### Isolation of bacteria and fungi isolates

#### Soil at dumpsite (sd)

Serial dilutions 1.0g of soil samples was aseptically carried out in sterile distilled water. Using a 10-fold serial dilutions 0.1ml of dilutions, 10<sup>-3</sup>, 10<sup>-6</sup> and 10<sup>-9</sup> were plated in triplicates on nutrient agar, MacConkey agar and Sabouraud dextrose agar by spread plate techniques. The Nutrient agar and MacConkey agar plates were then incubated at 37<sup>0</sup>C for 18-24hours, while the Sabouraud dextrose agar plate was incubated at 35<sup>0</sup>C for 48hours.

#### Waste at dumpsite (Wd)

Mixed decomposing wastes were aseptically collected 10g of the waste was soaked in 100ml sterile distilled water in a conical flask and well shaken to dislodge the organisms using a 10<sup>-</sup> fold serial dilution, 0.1ml of dilutions of 10<sup>-3</sup>, 10<sup>-6</sup> and 10<sup>-9</sup> were each plated on nutrient agar, MacConkey agar and Sabouraud dextrose agar by spread plate technique. The Nutrient agar and MacConkey agar plates were then incubated at 37<sup>0</sup>C for 18-24hours,

while the Sabouraud dextrose agar plate was incubated at 35°C for 48 hours.

Plant growing at dumpsite (Pd)

10g of plant leaves covered with dust and dirt at the dumpsite were introduced, into 90mls of nutrient broth, to concentrate the organisms and was then incubated at 37°C for 18-24 hours. The broth cultures were then plated by streaking on nutrient agar, MacConkey agar and Sabouraud dextrose agar and incubated at 37°C for 24 hours (Nutrient and MacConkey agar plates) and at 48-72hrs (Sabouraud dextrose agar plates).

Air at dumpsite/100 meters distance from dumpsite (Ad and A100).

Plates of nutrient agar, MacConkey agar and Sabouraud dextrose agar were exposed in triplicates at the dumpsites and at 100m away from dumpsite for 60 minutes during the busy hours of the day. The plates were then covered and incubated at 37°C for 24hours (Nutrient and MacConkey agar plates) and 35°C for 48-72 hours (Sabouraud dextrose agar plates)

#### Total bacterial and fungal counts

For each sample, the Nutrient agar and Sabouraud dextrose agar plates were used to calculate the total number of bacteria and fungi in the sample. The average colony counts were multiplied by their dilution factor to obtain the total bacterial and fungal count while for the air sampling, the average number of colonies per location was recorded as the total bacterial and fungal count of the air in that environment.

#### Purification and maintenance of culture

Purification of culture was by streak plate technique, pure cultures of isolates were maintained on Nutrient agar slants (for bacteria) and Sabouraud dextrose agar slants (for fungi) and stored in the refrigerator.

#### Identification of isolates

Bergey's manual of determinative bacteriology (Holt *et al.*, 1994; Cheesbrough, 2000) was used in the identification of the isolates, and standard tests as specified in microbiological techniques of Steane (1999) and Kaiser (1998) were carried out.

#### Physicochemical analysis of sample

The trace elements in the leachate were determined using atomic absorption spectrophotometer (UNICAM 919). 50mls of the sample was weighed and concentrated by evaporation and the residue homogenized. The homogenized sample was weighed, a shed and leached with 5ml standard flask. The volume was made up to mark (20ml capacity). The reagent blank was prepared without sample, the readings were obtained and results recorded. The temperature, pH, turbidity, conductivity, dissolved O<sub>2</sub>, biological oxygen demand (BOD), chloride,

sulphate, nitrite, nitrate and aluminum content of the leachate were measured according to the methods of Ekpo and Ibok (1998) and Ekpo and Ulrich (2000).

#### Statistical analysis

Data of the bacterial and fungal counts of the wastes sample was subjected to analysis and significant means determined using chi-square.

#### Bore-hole water analysis

##### Total bacterial count

1ml of water samples collected from bore-hole close to the study site (waste dumpsite) was inoculated by pour plate technique into Nutrient agar and incubated at 37°C for 48 hours. Result was recorded as colony forming units per milliliter of sample (cfu/ml).

##### Membrane filtration

Using a sterile 10ml pipette, 1ml of the sample was transferred into at 99ml of sterile water contained in a bottle and shaken. This dilution was then filtered through a membrane filter and the membrane filter was placed on a plate of sterile MacConkey agar and incubated at 37°C for 48hours. Coliforms appeared pink and were counted and reported as total Coliforms/100ml.

## RESULTS

### Average total bacterial and fungal counts of the two dumpsites.

Table 1 present the average total bacterial counts of the two dumpsites during the month of December, 2015. It showed that the garden street dumpsite had higher counts of  $25.2 \pm 0.17 \times 10^7$  CFU/g from the waste sample against  $15.2 \pm 0.11 \times 10^7$  Cfu/g and  $14.4 \pm 0.23 \times 10^7$  Cfu/g from the Eastern, highway waste soil and waste samples respectively. The Eastern highway air sample had higher count of  $2.7 \pm 0.18 \times 10^7$  Cfu/g against  $2.4 \pm 0.15 \times 10^7$  Cfu/g from garden street. The average total fungal count of the two dumpsites during the month of December, 2015 is as presented in table 2. It showed that the fungal count for waste soil, waste and air samples while  $11.2 \pm 0.11 \times 10^7$  Cfu/g,  $18 \pm 0.38 \times 10^2$  Cfu/g,  $0.7 \pm 0.01 \times 10^1$  Cfu/g were recorded for the Eastern highway dumpsite samples respectively.

The average total bacterial counts of the two dumpsites during the month of January 2016 are recorded in Table 3. The Garden street dumpsites had higher counts of  $21.2 \pm 2.33 \times 10^7$  Cfu/g,  $23.2 \pm 1.44 \times 10^7$  Cfu/g and  $3.6 \pm 0.88 \times 10^2$  Cfu/g, as against  $14.8 \pm 2.43 \times 10^7$  Cfu/g,  $14.8 \pm 1.88 \times 10^7$  Cfu/g and  $2.2 \pm 0.03 \times 10^2$  Cfu/g for the eastern highway dumpsites from the soil, waste and air samples respectively.

Table 4 presents the average total fungal count for the month of January 2016, it showed that the count  $15.0 \pm 1.83 \times 10^2$  Cfu/g,  $14 \pm 2.33 \times 10^2$  Cfu/g and  $0.80 \pm 0.10 \times 10^1$  Cfu/g were obtain from the soil,

waste and air samples from Garden street dumpsites while Eastern highway dumpsites samples had  $10 \pm 1.33 \times 10^2$  Cfu/g,  $11.8 \pm 0.87 \times 10^2$  Cfu/g and  $0.50 \pm 0.01 \times 10^1$  Cfu/g respectively from soil, waste and air samples.

Table 5 shows the total average bacterial counts for the two dumpsites in the month of February 2016. Garden street dumpsites had  $18 \pm 1.38 \times 10^7$  Cfu/g,  $16.4 \pm 1.45 \times 10^7$  Cfu/g and  $2.5 \pm 0.44 \times 10^2$  Cfu/g from soil, waste and air samples respectively, while the Eastern highway dumpsite samples had  $14 \pm 1.58 \times 10^7$  Cfu/g,  $14.8 \pm 1.35 \times 10^7$  Cfu/g and  $2.0 \pm 0.33 \times 10^2$  Cfu/g from soil, waste and air respectively. The average total fungal counts for the month of February 2016 from both dumpsites are presented in Table 6. The eastern highway dumpsite recorded higher fungal counts of  $17 \pm 1.65 \times 10^2$  Cfu/g and  $17 \pm 0.85 \times 10^2$  Cfu/g for the soil and waste samples respectively as against  $14 \pm 0.85 \times 10^2$  Cfu/g and  $11 \pm 0.55 \times 10^2$  Cfu/g obtained for the garden street samples respectively. The fungal counts in air was  $0.7 \pm 0.10 \times 10^1$  Cfu/g for Garden street and  $0.3 \pm 0.01 \times 10^1$  Cfu/g for Eastern highway dumpsite.

Table 7 presents the results of the average total bacterial count of samples from the two dumpsites during the month of April 2016 for the soil samples from Garden street site, a bacterial count  $25.8 \pm 1.37 \times 10^7$  Cfu/g was obtained while  $15.6 \pm 1.26 \times 10^7$  Cfu/g was obtained for samples from the Eastern highway site soil, for the waste samples, an average of  $29.4 \pm 1.33 \times 10^7$  Cfu/g was obtained for Garden street dumpsite while for the Eastern highway dumpsite the count of  $15.4 \pm 1.28 \times 10^7$  Cfu/g was recorded. For the air samples,  $3.1 \pm 0.87 \times 10^1$  Cfu/g and  $2.80 \pm 0.55 \times 10^1$  Cfu/g were obtained for air at dumpsite (Ad) and air at a 100m distance (Ad<sub>100</sub>) away from the dumpsite respectively for the Garden street dumpsite, while  $1.90 \pm 0.06 \times 10^1$  Cfu/g and  $2.0 \pm 0.03 \times 10^1$  Cfu/g were obtained for the two samples (Ad and Ad<sub>100</sub>) respectively from the Eastern highway dumpsite. Bacterial counts for the plant samples (Pd) had an average of  $11.0 \pm 1.48 \times 10^4$  Cfu/g and  $7.3 \pm 1.33 \times 10^4$  Cfu/g for the Garden street and Eastern highway dumpsites respectively. The leachate analysis revealed an average total bacterial count of  $19.6 \pm 1.88 \times 10^7$  Cfu/g for the Garden street dumpsite.

Table 8 shows the average total fungal count of samples, from the two dumpsites during the month of April, 2016. The soil samples (Sd) gave  $1.8 \pm 0.06 \times 10^3$  Cfu/g for the garden street dumpsite and  $0.7 \pm 0.01 \times 10^3$  Cfu/g for the Eastern highway dumpsite. The waste samples (Wd) gave  $1.7 \pm 0.03 \times 10^3$  Cfu/g and  $0.9 \pm 0.01 \times 10^3$  Cfu/g for the Garden street and eastern highway dumpsites respectively. From the two air samples, air at

dumpsite (Ad) and air 100m away from the dumpsite (Ad<sub>100</sub>) gave  $0.5 \pm 0.1 \times 10^3$  Cfu/g and  $0.5 \pm 0.01 \times 10^3$  Cfu/g were recorded respectively for both dumpsites. The plant at dumpsite sample gave  $0.4 \pm 0.10 \times 10^3$  Cfu/g and  $0.5 \pm 0.11 \times 10^3$  Cfu/g. Total fungal count in the Garden street leachate was  $1.4 \pm 0.11 \times 10^3$  Cfu/g. The average total bacterial counts of the two dumpsites obtained for the month of May, 2016 are recorded in Table 9. For the soil samples (Sd), counts of  $26.4 \pm 1.88 \times 10^7$  Cfu/g and  $16.1 \pm 0.87 \times 10^7$  Cfu/g were obtained from garden street and eastern highway dumpsites respectively. The waste samples (Wd) from Garden street and Eastern highway dumpsite had a total bacterial count of  $26 \pm 2.87 \times 10^7$  Cfu/g and  $18.8 \pm 1.97 \times 10^7$  Cfu/g respectively. A total of  $2.6 \pm 0.86 \times 10^1$  Cfu/g and  $2.5 \pm 0.79 \times 10^7$  Cfu/g was recorded for the two air samples (Air at dumpsite (Ad) and Air 100m from dumpsite (Ad<sub>100</sub>)) respectively for the garden street dumpsite while  $2.1 \pm 0.18 \times 10^1$  Cfu/g and  $2.1 \pm 0.16 \times 10^1$  Cfu/g were recorded from the two air sample (Ad and Ad<sub>100</sub>) respectively for the Eastern highway dumpsite. The plant samples gave  $9.0 \pm 1.84 \times 10^4$  Cfu/g and  $5.2 \pm 0.98 \times 10^4$  Cfu/g for the garden street and eastern highway samples respectively while the leachate from Garden street gave  $2.4 \pm 0.88 \times 10^7$  Cfu/g.

Table 10 presents the average total fungal counts of samples from the two dumpsites during the month of May, 2016. The soil samples (Sd) had an average count of  $1.4 \pm 0.03 \times 10^3$  Cfu/g and  $0.9 \pm 0.01 \times 10^3$  Cfu/g. For the Garden street dumpsite and Eastern highway dumpsite. Air from dumpsite (Ad) and air 100m away from the dumpsite (Ad<sub>100</sub>) recorded  $0.4 \pm 0.01 \times 10^1$  Cfu/g and  $0.5 \pm 0.4 \times 10^{10}$  Cfu/g and  $0.5 \pm 0.02 \times 10^1$  Cfu/g respectively for the eastern highway dumpsite. The plant sample (Pd) and the leachate showed a count of  $0.4 \pm 0.02 \times 10^3$  Cfu/g and  $1.2 \pm 0.0 \times 10^1$  Cfu/ml respectively for the garden street dumpsite while the eastern highway plant sample had  $0.5 \pm 0.01 \times 10^3$  Cfu/g.

Table 11 presents the result of the average total bacterial count of the two dumpsites for the month of June, 2016. For the Garden street dumpsites, soil sample (Sd) and waste samples (Wd) recorded a count of  $25.6 \pm 2.86 \times 10^2$  Cfu/g and  $26. \pm 2.33 \times 10^7$  Cfu/g respectively, whereas Eastern highway dumpsite recorded  $17.6 \pm 1.77 \times 10^7$  Cfu/g and  $14.4 \pm 1.87 \times 10^7$  Cfu/g for the same samples respectively. For the air samples at dumpsite (Ad) and 100 meters away from dumpsite (Ad<sub>100</sub>), Garden street recorded  $2.4 \pm 0.85 \times 10^1$  Cfu/g and  $2.4 \pm 0.57 \times 10^1$  Cfu/g respectively while Eastern highway recorded  $19.0 \pm 0.86 \times 10^1$  Cfu/g and  $2.3 \pm 0.88 \times 10^1$  Cfu/g respectively. Plant leaf surfaces from the Garden street dumpsite had  $9.5 \pm 1.33 \times 10^4$  Cfu/g while plant from the Eastern highway dumpsite and  $5.0 \pm 1.26 \times 10^4$  Cfu/g. for the leachate

sample (L), the garden street dumpsite had  $23.6 \pm 2.89 \times 10^7$  cfu/ml.

Table 12 presents the average total fungal count for the month  $1.83 \pm 0.31 \times 10^3$  cfu/g was obtained in soil samples (Sd) from Garden street and Eastern highway dumpsites respectively. The waste samples from the Garden street dumpsite had an average of  $0.7 \pm 0.11 \times 10^3$  cfu/g while that of the Eastern highway dumpsite had  $1.7 \pm 0.23 \times 10^2$  cfu/g. the two air samples, Ad and Ad<sub>100</sub>, both had  $0.5 \pm 0.01 \times 10^1$  cfu/g for the Garden street dumpsite and  $0.4 \pm 0.11 \times 10^1$  cfu/g for the eastern highway dumpsite. The garden street leachate showed a count of  $1.2 \pm 0.18 \times 10^3$  cfu/ml.

Fig. 2 to 6 shows a comparative analysis of the two dumpsite based on the monthly counts of the microbial load of the samples. Fig. 2 and 3 showed that the air counts were higher in the dry season of the year, that is from December to February, than the wet season. In fig. 4, it was observed that the Garden street plant leaves sample had more bacterial counts than that of Eastern highway. Fig. 5 shows an increase in the waste soil bacteria count in the wet season of the year and a decrease in the dry season from the two dumpsite samples. In fig. 6, it was observed that apart from the high counts observed in February and June, the soil fungal counts from the eastern highway samples were lower than that of garden street.

#### Characterization and identification of isolates from the waste samples.

The characterization of cellular morphology and biochemical characterization and identification of bacterial isolates in the wet season are presented in table 13 and 14, while table 15 presents the biochemical characterization and identification of bacterial isolates obtained in the dry season. From these results, 17 bacterial isolates were obtained in the wet season while 8 isolates were obtained in the dry season. The bacterial isolates identified from the waste samples during the wet season were *Staphylococcus spp*, *Enterococcus spp*, *Proteus spp*, *E. Coli*, *Klebsiella spp*, *Shigella spp*, *Salmonella spp*, *Bacillus spp*, *Micrococcus spp* and *Pseudomonas spp*, while those identified during the dry season were; *E.coli*, *Kebsiella spp*, *Shigella spp*, *Salmonella spp*, *Bacillus spp*, *Micrococcus spp* and *Pseudomonas spp*.

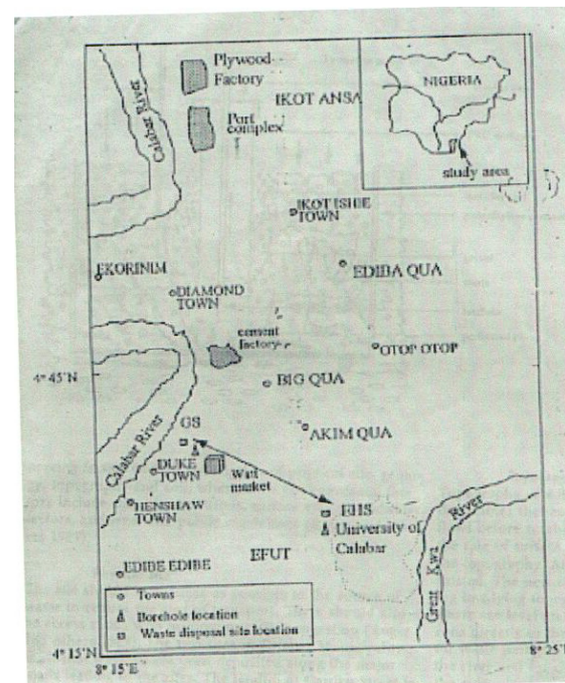
From the Sabouraud dextrose agar plates, nine isolates were obtained from the various samples as summarized in table 16. They were

identified based on their colony appearance on the agar medium and the morphological characteristics as *Aspergillus spp*, *Mucor spp*, *Nocardia spp* and *Actinomyces spp*.

#### Physicochemical analysis of the collected waste samples

The values for the trace and major elements of the leachate sample collected from garden street dumpsite is presented in table 17 while that of other physicochemical properties is present in table 18. It showed that the highest element was sodium (28600mg/l) followed by potassium (878.57mg/l), iron (50mg/l), magnesium (23.25mg/l), calcium (21.85mg/l), Manganese (4.447MG/L), Cobalt (1.30mg/l), Copper (0.500mg/l) and Cadmium (0.157mg/l) as the least. Apart from the temperature and BOD, the values obtained for the physicochemical properties were lower than the WHO permissible limits.

Microbiological analysis of bore-hole water sample located near the garden street dumpsite. Table 19 presents the result of microbiological analysis of the bore-hole water sample from a location near the garden street dumpsite. It showed that the average bacterial and coliform count for both the wet and dry season ( $8.0 \pm 1.35 \times 10^1$  and  $19.6$  cfu/ml) and ( $9.7 \pm 1.45 \times 10^1$  and  $22.33$  cfu/ml) respectively, were high and exceeded the accepted WHO limits of not more than 0-3/100ml of drinking water.



**Fig 1: Map of Calabar Municipality showing sample location**

**Table 1: Total heterotrophic bacterial counts of the Eastern highway and Garden street dumpsites during the month of December 2005**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	25.2±0.22x10 <sup>7</sup> cfu/g	15.2±0.11x10 <sup>7</sup> cfu/g
W <sub>d</sub>	22.4±22.4x10 <sup>7</sup> cfu/g	14.4±0.23x10 <sup>7</sup> cfu/g
A <sub>d</sub>	2.4±0.15x10 <sup>2</sup> cfu/g	2.7±0.17x10 <sup>2</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table 2: Total fungal count of the Garden street and Eastern highway dumpsites during the month of December 2015**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	14 ±0.33x10 <sup>2</sup> cfu/g	11 ±1.22 x10 <sup>2</sup> cfu/g
W <sub>d</sub>	12±1.32x10 <sup>2</sup> cfu/g	18±0.38x10 <sup>2</sup> cfu/g
A <sub>d</sub>	1.1±0.11x10 <sup>1</sup> cfu/g	0.7±0.01x10 <sup>1</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table 3: Total bacterial counts of the Garden street and Eastern highway dumpsites during the month of January 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	21.2 ±2.33x10 <sup>7</sup> cfu/g	14.8±2.4 x10 <sup>7</sup> cfu/g
W <sub>d</sub>	23.3±1.44x10 <sup>7</sup> cfu/g	14.8±1.88 x10 <sup>7</sup> cfu/g
A <sub>d</sub>	3.6 ±0.88 x10 <sup>2</sup> cfu/g	2.2±0.03x10 <sup>2</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table 4: Total fungal count of the Garden street and Eastern highway dumpsites during the month of January 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	15.0 ±1.83x10 <sup>2</sup> cfu/g	10±1.33 x10 <sup>2</sup> cfu/g
W <sub>d</sub>	14±2.33x10 <sup>2</sup> cfu/g	11±0.87 x10 <sup>2</sup> cfu/g
A <sub>d</sub>	0.80±0.01x10 <sup>1</sup> cfu/g	0.50±0.01x10 <sup>1</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table .5: Total heterotrophic bacteria counts of the Garden street and Eastern highway dumpsites during the month of February 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	18±1.38x10 <sup>7</sup> cfu/g	14±1.58 x10 <sup>7</sup> cfu/g
W <sub>d</sub>	16.4±1.45x10 <sup>7</sup> cfu/g	14.8±0.1.35 x10 <sup>7</sup> cfu/g
A <sub>d</sub>	2.5±0.48 x10 <sup>2</sup> cfu/g	2.0± 0.033 x10 <sup>2</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table 6: Total fungal count of the Garden street and Eastern highway dumpsites during the month of February 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	14 ±0.8x10 <sup>2</sup> cfu/g	17±1.65 x10 <sup>2</sup> cfu/g
W <sub>d</sub>	11±0.55 x10 <sup>2</sup> cfu/g	17±0.0.85 x10 <sup>2</sup> cfu/g
A <sub>d</sub>	0.7±0.11x10 <sup>1</sup> cfu/g	0.3±0.01x10 <sup>1</sup> cfu/g

**Key:** S<sub>d</sub> - Soil dumpsite  
 W<sub>d</sub> - Waste at dumpsite  
 A<sub>I</sub> - Air at dumpsite

**Table 7: Total heterotrophic bacterial count of the Garden street and Eastern highway dumpsites during the month of April 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	25.8 ± 1.37 x 10 <sup>7</sup> cfu/g	15.6 ± 1.26 x 10 <sup>7</sup> cfu/g
W <sub>d</sub>	29.4 ± 1.33 x 10 <sup>7</sup> cfu/g	15.4 ± 1.28 x 10 <sup>7</sup> cfu/g
A <sub>d</sub>	3.1 ± 0.87 x 10 <sup>1</sup> cfu/g	1.90 ± 0.06 x 10 <sup>7</sup> cfu/g
Ad <sub>100</sub>	2.80 ± 0.55 x 10 <sup>1</sup> cfu/g	2.0 ± 0.03 x 10 <sup>1</sup> cfu/g
P <sub>d</sub>	11 ± 1.48 x 10 <sup>1</sup> cfu/g	7.3 ± 1.33 x 10 <sup>4</sup> cfu/g
L	19.6 ± 1.87 x 10 <sup>7</sup> cfu/g	

**Key:** S<sub>d</sub> - Soil dumpsite  
W<sub>d</sub> - Waste at dumpsite  
A<sub>d</sub> - Air at dumpsite  
Ad<sub>100</sub> - Air at a distance of 100meters from the dumpsite  
P<sub>d</sub> - Plant at dumpsite  
L - Leachate

**Table 8: Total fungal counts of the Garden street and Eastern highway dumpsite during the month of April, 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	1.8 ± 0.06 x 10 <sup>3</sup> cfu/g	0.7 ± 0.01 x 10 <sup>3</sup> cfu/g
W <sub>d</sub>	1.7 ± 0.03 x 10 <sup>3</sup> cfu/g	0.9 ± 0.01 x 10 <sup>3</sup> cfu/g
A <sub>d</sub>	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g
Ad <sub>100</sub>	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g
P <sub>d</sub>	0.4 ± 0.01 x 10 <sup>3</sup> cfu/g	0.5 ± 0.11 x 10 <sup>3</sup> cfu/g
L	1.4 ± 0.11 x 10 <sup>3</sup> cfu/g	-

**Key:** S<sub>d</sub> - Soil dumpsite  
W<sub>d</sub> - Waste at dumpsite  
A<sub>d</sub> - Air at dumpsite  
Ad<sub>100</sub> - Air at a distance of 100meters from the dumpsite  
P<sub>d</sub> - Plant at dumpsite  
L - Leachate

**Table 9: Total heterotrophic bacterial count of the Garden street and Eastern highway dumpsites during the month of April 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	26.4 ± 1.88 x 10 <sup>7</sup> cfu/g	16.1 ± 0.87 x 10 <sup>7</sup> cfu/g
W <sub>d</sub>	26 ± 2.87 x 10 <sup>7</sup> cfu/g	18.8 ± 1.197 x 10 <sup>7</sup> cfu/g
A <sub>d</sub>	2.6 ± 0.86 x 10 <sup>1</sup> cfu/g	2.1 ± 0.18 x 10 <sup>1</sup> cfu/g
Ad <sub>100</sub>	2.5 ± 0.70 x 10 <sup>1</sup> cfu/g	2.1 ± 0.16 x 10 <sup>1</sup> cfu/g
P <sub>d</sub>	9.0 ± 1.184 x 10 <sup>4</sup> cfu/g	5.2 ± 0.98 x 10 <sup>4</sup> cfu/g
L	2.4 ± 0.88 x 10 <sup>7</sup> cfu/g	-

**Key:** S<sub>d</sub> - Soil dumpsite  
W<sub>d</sub> - Waste at dumpsite  
A<sub>d</sub> - Air at dumpsite  
Ad<sub>100</sub> - Air at a distance of 100meters from the dumpsite  
P<sub>d</sub> - Plant at dumpsite  
L - Leachate

**Table 10: Total fungal count of the Garden street and Eastern highway dumpsites during the month of May 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	1.4 ± 0.03 x 10 <sup>3</sup> cfu/g	0.9 ± 0.01 x 10 <sup>3</sup> cfu/g
W <sub>d</sub>	1.2 ± 0.05 x 10 <sup>3</sup> cfu/g	0.7 ± 0.01 x 10 <sup>3</sup> cfu/g
A <sub>d</sub>	0.4 ± 0.01 x 10 <sup>1</sup> cfu/g	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g
Ad <sub>100</sub>	0.5 ± 0.01 x 10 <sup>1</sup> cfu/g	0.5 ± 0.04 x 10 <sup>1</sup> cfu/g
P <sub>d</sub>	0.4 ± 0.02 x 10 <sup>3</sup> cfu/g	0.5 ± 0.01 x 10 <sup>3</sup> cfu/g
L	1.2 ± 0.01 x 10 <sup>3</sup> cfu/g	

**Key:** S<sub>d</sub> - Soil dumpsite  
W<sub>d</sub> - Waste at dumpsite  
A<sub>d</sub> - Air at dumpsite

A <sub>d100</sub>	-	Air at a distance of 100meters from the dumpsite
P <sub>d</sub>	-	Plant at dumpsite
L	-	Leachate

**Table 11: Total heterotrophic bacterial count of the Garden street and Eastern highway dumpsites during the month of June 2016**

Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	25.6 ± 2.86 × 10 <sup>7</sup> cfu/g	7.6 ± 1.77 × 10 <sup>7</sup> cfu/g
W <sub>d</sub>	26 ± 2.33 × 10 <sup>7</sup> cfu/g	16.4 ± 1.87 × 10 <sup>7</sup> cfu/g
A <sub>d</sub>	2.4 ± 0.85 × 10 <sup>1</sup> cfu/g	1.9 ± 0.086 × 10 <sup>1</sup> cfu/g
A <sub>d100</sub>	2.4 ± 0.57 × 10 <sup>1</sup> cfu/g	2.3 ± 0.088 × 10 <sup>1</sup> cfu/g
P <sub>d</sub>	9.5 ± 1.33 × 10 <sup>4</sup> cfu/g	5.0 ± 1.26 × 10 <sup>4</sup> cfu/g
L	23.6 ± 2.89 × 10 <sup>7</sup> cfu/g	

**Key:**

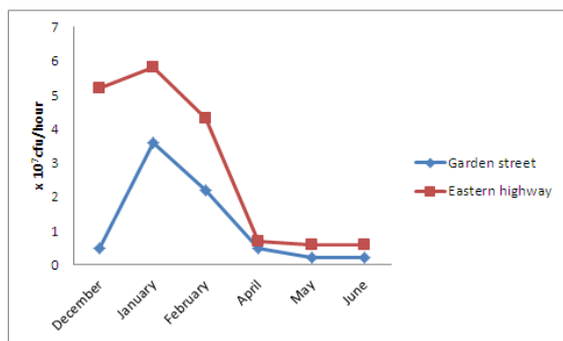
S <sub>d</sub>	-	Soil dumpsite
W <sub>d</sub>	-	Waste at dumpsite
A <sub>d</sub>	-	Air at dumpsite
A <sub>d100</sub>	-	Air at a distance of 100meters from the dumpsite
P <sub>d</sub>	-	Plant at dumpsite
L	-	Leachate

**Table 12: Total fungal counts of the Garden street and Eastern highway dumpsites during the month of June 2016**

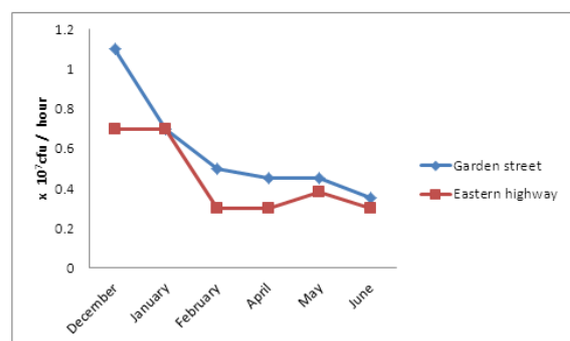
Sample	Garden street (cfu/g)	Eastern highway (cfu/g)
S <sub>d</sub>	1.1 ± 0.13 × 10 <sup>3</sup> cfu/g	1.83 ± 0.31 × 10 <sup>3</sup> cfu/g
W <sub>d</sub>	0.7 ± 0.11 × 10 <sup>3</sup> cfu/g	1.7 ± 0.23 × 10 <sup>3</sup> cfu/g
A <sub>d</sub>	0.5 ± 0.01 × 10 <sup>1</sup> cfu/g	0.4 ± 0.011 × 10 <sup>1</sup> cfu/g
A <sub>d100</sub>	0.5 ± 0.01 × 10 <sup>1</sup> cfu/g	0.5 ± 0.01 × 10 <sup>1</sup> cfu/g
P <sub>d</sub>	0.4 ± 0.11 × 10 <sup>4</sup> cfu/g	0.5 ± 0.12 × 10 <sup>4</sup> cfu/g
L	1.2 ± 0.18 × 10 <sup>3</sup> cfu/g	

**Key:**

S <sub>d</sub>	-	Soil dumpsite
W <sub>d</sub>	-	Waste at dumpsite
A <sub>d</sub>	-	Air at dumpsite
A <sub>d100</sub>	-	Air at a distance of 100meters from the dumpsite
P <sub>d</sub>	-	Plant at dumpsite
L	-	Leachate



**Fig. 2: Bacterial counts of the air samples for the two seasons of the year**



**Fig. 3: The fungal counts of the air samples for the two seasons of the year**

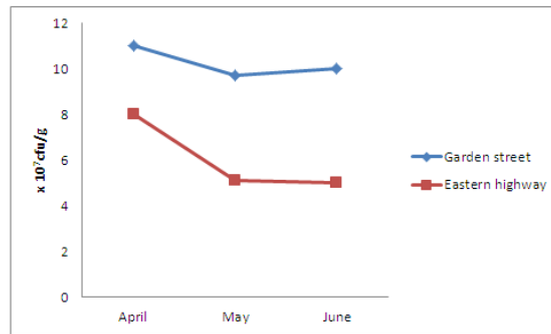


Fig.4: The bacterial counts of the plant samples for the wet season of the year

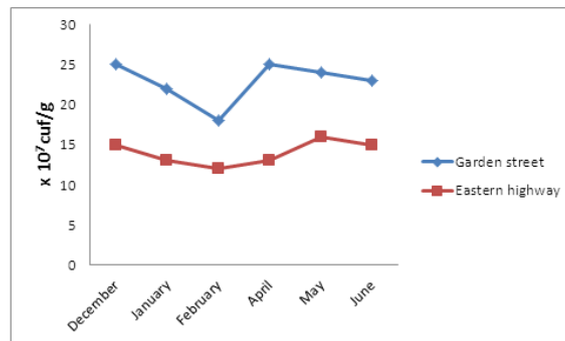


Fig. 5: The bacterial counts of the soil samples of the two season of the year

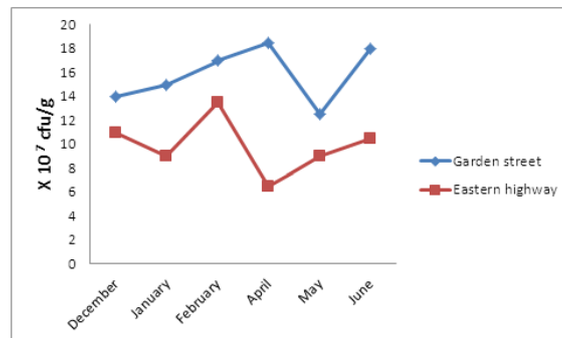


Fig. 6: The fungal counts of the soil samples for the two seasons of the year

**Table 13: Characterization of cellular morphology of bacterial isolate (April to June 2016)**

Colony	1	2	3	4	5	6	7	8	9	10	11	12	13	
Pigmentation	Cream	Straw yellow	Off white	Off white	Cream	Pin	Cream	Cream	Cream	Yellow	Cream	Cream	Pink	
Elevation	Flat entire	Raised entire	Flat lobate	Flat lobate	Raised entire	Flat entire	Convex entire	Flat entire	Raised entire	Raised entire	Flat rhizoid	Raised entire	Nat undulate	
Optical characteristics	Translucent	Translucent	Translucent	Opaque viscid	Translucent	Translucent	Transparent	Translucent	Opaque	Opaque	Opaque	Opaque	Translucent	
Consistency colony	-	butyrous	Butyrous	Membranous	Butyrous	Viscid	Viscid	Viscid	Viscid	Viscid	Butyrous	Dry	Butyrous	
Surface colony shape	Glossy circular	Glossy circular	Dull/dry circular	Dull/dry circular	Glossy circular	Dry	Glossy	Dull						
Gram's reaction	+ve	+ve	+ve	+ve	+ve	+ve	-ve	-ve	+ve	+ve	-ve	+ve	+ve	
Cell shape	Rods	Cocci	Rods	Rods	Rods	Cocci	Rods	Rods	Cocci	Cocci	Rods	Cocci	Rods	
motility	+ve	+ve	+ve	+ve	+ve	+ve	+ve	+ve	-ve	+ve	+ve	+ve	-ve	

**Table 14: Biochemical characterization/identification of bacterial isolates (April-June 2016)**

Isolate	1	2	3	4	5	6	7	8	9	10	11	12	13
Catalase	+	+	+	+	+	+	+	+	-	+	+	+	+
citrate	-	-	-	+	-	-	-	-	-	-	-	-	+
Hs											-	-	
Productio	-	-	-	-	-	-	-	+	-	-	+	-	-
n	-	-	-	-	-	-	-	-	-	-	-	-	-
Coagulase	-	-	-	-	-	-	-	-	-	-	-	-	-
Indole													
MR-VP	- +	+ -	- +	- -	+ +	- -	+ +	- +	+ +	+	+ +	+	+ +
Oxidase	+	+	+	+	-	+	-	+	+	-	-	-	-
Glucose	A	-	A	A	A	-	A	A	-	-	AG	A	A
Lactose	-	-	+	-	-	+	+	A	A	A	AG	-	A
Manitol	-	-	-	-	-	-	-	-	-	-	-	-	A
Sucrose	-	-	-	-	-	-	-	-	-	-	AG	-	A
Probable organism	<i>Bacillus spp</i>	<i>Micrococci</i>	<i>Bacillus spp</i>	<i>Bacillus spp</i>	<i>Bacillus sp</i>	<i>Micrococc</i>	<i>Bacillus spp</i>	<i>Bacillus spp</i>	<i>Enterococcus faecalis</i>	<i>Staphylococcus spp</i>	<i>Proteus spp</i>	<i>Staphylococcus spp</i>	<i>Klebsiella spp</i>

**Key:** + Positive  
 - Negative  
 A Acid  
 AG Acid and gas  
 MR Methyl red  
 VP Vogues proskaeur

**Table 14: Biochemical characterization/identification of bacterial isolates (April-June 2016)**

Isolate	1	2	3	4	5	6	7	8
Grams reaction	-	-	-	-	-	+	+	-
Cell shape	Rod	Rod	Rod	Rod	Rod	Rod	Cocci	Rod
Motility	+	-	+	-	+	+	-	+
Indole	+	-	-	-	-	-	-	-
Catalase	+	+	+	+	+	+	+	+
Citrate	-	+	+	-	+	+	-	-
MR	+	-	-	+	-	+	-	-
VP	-	+	+	-	+	+	-	-
Glucose	AG	AG	AG	-	AG	A	A	A
Lactose	A	A	A	-	-	A	A	-
Mannitol	A	A	A	-	A	A	-	-
Sucrose	A	A	A	-	A	A	A	A
Probable organism	<i>E.coli</i>	<i>Klebsiella spp</i>	<i>Enterobacter spp</i>	<i>Shigella spp</i>	<i>Salmonella spp</i>	<i>Bacillus spp</i>	<i>Micrococcus spp</i>	<i>Pseudomonas</i>

**Key:** + Positive  
 - Negative  
 A Acid  
 AG Acid and gas  
 MR Methyl red  
 VP Vogues proskaeur

**Table 16: Morphological characterization of isolates from the garden street dumpsite samples grown on sabouroud dextrose agar**

Isolates	Colony appearance	Nature of hyphae	Arrangement of spores	Columella	Probable organism
1.	Black	Aseptate	Spores not embedded	Absent	<i>Aspergillus sp</i>
2.	Grey	Aseptate	Spores in sporangia	Present	<i>Mucoer sp.</i>
3.	Greenish yellow	Aseptate	Spores not enclosed in sporangia	Absent	<i>Aspergillus sp</i>
4.	Cottony white	Aseptate	Spores in sporangia	Present	<i>Mucor sp.</i>
5.	Velvet green	Mass of tiny filaments	Not observed	Absent	<i>Nocardia.sp</i>
6.	Yellow	Aseptate	Spores not enclose	Absent	<i>Aspergillus sp</i>
7.	Dark green	Mass of tiny filaments	Not observed	Absent	<i>Nocardia sp</i>
8.	Black	Aseptate	Spores not enclose	Absent	<i>Aspergillus sp</i>
9.	White drops	Not observed	Granular spores	Absent	<i>Actinomyces sp</i>

**Filamentous bacteria****Table 17: Trace and major elements content of the lachate from the garden street dumpsite**

Metals	Sample mg/l	WHO (1984) std mg/l
Iron (Fe)	50.00	3.0
Copper (Cu)	0.500	1.0
Sodium (Na)	2800.0	200.00
Potassium (k)	878.571	-
Cobalt (Co)	1.30	-
Cadmium (Cd)	0.157	0.005
Manganese (Mn)	4.447	-
Calcium (Ca)	21.851	100.00
Magnesium (Mg)	23.25	-
Lead (Pb)	1.9946085	-

**Table 18: The physicochemical properties of the leachate from the garden street dumpsite**

Parameters	Values obtained	WHO (1984)
Temperature ( <sup>o</sup> C)	28.4	12-25
pH	8.5	6.5-9.5
Turbidity (FTU)	1305.00	-
Dissolved O <sub>2</sub> (mg/L)	3.80	4.0
BOD <sub>5</sub> (mg/l)	210.40	0.5
Chloride (mg/l)	125.8	250
Sulphate (mg/l)	50.45	500
Nitrate (mg/l)	0.04	0.1
Nitrite (mg/l)	0.43	45

**Table 19: Microbiological analysis of the bore-hold water sample from a location near the garden street dumpsites**

Sample	Bacterial count CFU/ml		Total coliforms/ml	
	Wet season	Dry season	Wet season	Dry season
Month 1	8.0 ± 0.55x10 <sup>1</sup>	9.8 ± 0.88x10 <sup>1</sup>	21	23
Month 2	7.5 ± 1.008x10 <sup>1</sup>	9.0 ± 0.87x10 <sup>1</sup>	18	22
Month 3	8.6 ± 1.33x10 <sup>1</sup>	1.02 ± 1.39x10 <sup>1</sup>	20	22
Average	8.0 ± 1.35x10 <sup>1</sup>	9.7 ± 1.88x10 <sup>1</sup>	19.6	22.33

## DISCUSSION

From this study, a higher total bacterial count was observed in the garden street dumpsite compared to its eastern highway dumpsite counterpart. This could be due to the fact that the garden street dumpsite is an unplanned, uncontrolled and open dumpsite compared to the eastern, highway dumpsite which was a bit closed. Also the higher bacteria count observed in the garden street dumpsite could have also been due to its close proximity to the market, as it receives fresh waste daily and in large quantities. A comparative analysis of the two dumpsites showed that there was no significant difference ( $P \geq 0.05$ ) in the bacterial and fungal counts of the two dumpsites during dry season (December to February) as in calculated  $\chi^2$  (2.818) was less than the tabulated bacterial count (5.991) was also lesser than the tabulated (5.991) while for the fungal count, the calculated  $\chi^2$  (0.8303) was also lesser than the tabulated (5.991). during the rainy season (April to June), there was no significant difference in the soil bacteria count between the two dumpsites as the calculated  $\chi^2$  (0.806) was less than the tabulated  $\chi^2$  (5.991), as the reverse was the case in the soil fungal count, as the calculated  $\chi^2$  (6.698) was greater than the tabulated  $\chi^2$  (5.991). Also during the dry season. For the dumpsite air samples during the dry season showed significant difference (Calculated  $\chi^2$ , 23.174 >  $\chi^2$  tabulated 5.991), while there was no significant difference in the fungal counts between the dumpsites (calculated  $\chi^2$ , 0.416 <  $\chi^2$  tabulated, 5.991). a comparative analysis of the dumpsite samples showed that there was a higher bacterial and fungal counts during the wet seasons (April to June) compared to the dry seasons (December to February). The rationale behind this observation could probably have been due to the increased moisture content in the waste dumpsite during the wet season, as research has shown that water is required for both metabolic activities and proliferation of microbial cells (4). The presence of pathogenic bacteria such *Bacillus spp*, *Proteus sp*, *Enterococcus sp*, *Micrococcus Pseudomonas sp*, *Staphylococcus sp* and *Coliforms* such as *Klebsiella sp*, *E.coli*, *Shigella sp*, and *Salmonella sp* in the waste dumpsite as observed in the study was not surprising, as it corroborates with that of (14), who reported to have identified the presence of *Coliforms*, *Facal Coliforms* and pathogens such as *Escherichia Coli*, *Streptococcus*, *Pseudomonas* and *Salmonella* from samples collected close to sewage sites. Also the observation was in line with that of who identified the presence of *Bacillus*, *Staphylococcus*, *Klebsiella* from a waste dumpsite located at eagle island, River State (15). The presence of these identified organisms in the study site is a thing of great concern as these bacteria

have been associated with a number of public health problems *Proteus* are human pathogens and has been shown to occur in manure, soil and polluted waters (16) they are human pathogens and are capable of causing urinary tract infections and also they serve as secondary inhalers that may cause septic lesions In burn patients. *Klebsiella* and other species are opportunistic pathogens that has been reported to occur in soil, water, vegetables and waste sites, they can cause bactericemia, Pneumonia, urinary tract and other human infections (17), they frequently cause infections in neonatal, intensive care and immune suppressed patients (17). *Enterobacter* has been reported to occur in soil, fresh water, sewage, plants, vegetables and are associated with urinary tract infections *E. coli* are capable of producing enterotoxins and other virulence factors including invasive and colonization factors, they can cause diarrheal disease, urinary tract infections and nosocomial infections including septicemia and meningitides (18). *Salmonella* and *Shigella* are human pathogens and are the causative agents of typhoid fever, enteric fevers, gastroenteritis, septicemia and dysentery respectively (18).

The presence of fungi such as *Aspergillus sp* and *Actinomyces sp*, is of public health importance, as actinomycetes are known to actinomycosis and nocardiosis in cattle, dogs and humans (19), while *Aspergillus spp* are important producers of mycotoxins (e.g aflatoxins of *A. Flavus*). In human, they cause Aspergillosis and they have been reported to colonize the mucosal surface and then lungs thereby causing immune-suppression (19). The spores of *Aspergillus* in the environment also causes allergic response, and they may also cause opportunistic infections in immunocompromised persons. The result of physicochemical analysis of the leachate of the waste sample shows that most of the trace and major elements content of the leachate values obtained were under the WHO recommended values except for sodium and cadmium for other physicochemical parameters, the BOD, (210.40mg/L) exceeded the WHO permissible value (0.5). This observation corroborates with that of (20) the high amount of cadmium is of great concern and public health importance, the metal is known to be carcinogenic, embryo toxic and could cause kidney damage (21) (22) (23). The high total Coliform count (196-23cfu/ml) obtained from the microbiological analysis of the bore-hole water sample from a location near the garden street dumpsite was an indication of gross contamination from the leachate of the dumpsite, as researches has shown that inadequate control of refuse dumpsite where leachate generated are allowed to escape the surrounding and underlying water body is a major

threat to bore-hole water. The high Coliform count obtained from the bore-hole water sample exceeds that of 0-3cfu/100ml for drinking water as stipulated by (26)(27). However, this bore-hole was thereby unfit for consumption and could be a potential source of water borne disease to the public at large.

## CONCLUSION

Solid waste disposed in open dumpsite is usually subjected to series of complex biochemical and physical processes, which lead to the production of both leachate and gaseous emissions, and when this leachate leaves the dumpsite and reaches the water tables, it results in borehole contamination. Besides, the aforementioned consequence, open dumpsite also serve as breeding sites for pathogenic organisms which are capable of causing both public health as well as environmental hazards. With all this negative approaches accrued to open dumpsite, we therefore call on government and environmental agencies to help provide well planned and closed dumpsite, as well as good waste management systems, so as to help reduce or curb further public health risk and environmental hazards that may result from the use of open and unplanned dumpsites for sewage deposition.

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