Isolation and Characterization of Hydrocarbon-utilizing Bacteria from Petroleum Sludge Samples obtained from Crude Oil Processing Facility in Nigeria

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ABSTRACT: The isolation and characterization of hydrocarbon-utilizing bacteria from petroleum oily sludge collected from crude oil processing plant in Rivers State, Nigeria was carried out. Microbiological analysis of the sludge sample showed that the microbial load consisted average of 2.5 x 10^6 cfu/g total heterotrophic bacterial (THB) count, while the hydrocarbon utilizing bacterial (HUB) count was average of 2.0 x 10^3 cfu/g. Phenotypic characteristics and phylogenetic analysis of the 16S rRNA gene sequence of the isolates revealed that they were related to members of the species Klebsiella pneumonia, Klebsiella oxytoca and Alcaligenes faecalis. The 16S rRNA of the isolates showed percentage similarities to the type strains (99% sequence similarities) of the following; Klebsiella pneumonia strain B21 (gi: 922317936), Klebsiella pneumonia strain ICB-C183 (gi: 908478837), Klebsiella oxytoca strain BCNAI (gi: 846993354), Klebsiella oxytoca strain BC4 (gi: 884060138), Alcaligenes faecalis strain IOU PMR (gi: 686028963) and Alcaligenes faecalis strain AQ-I (gi: 816845513). Of the six hydrocarbon utilizing bacteria identified, 4(67%) were Klebsiella species while 2(33%) were identified as Alcaligenes sp of different strains. The result suggest that these isolated organisms from the petroleum sludge can be explored and used to promote environmentally friendly technology clean-up of petroleum hydrocarbon polluted sites in the Niger Delta region of Nigeria.© JASEM https://dx.doi.org/10.4314/jasem.v21i2.17

Keywords: Hydrocarbon-utilizing bacteria, Petroleum sludge, 16S rRNA, Phylogenetic, Gene sequence.

One of the most encountered pollutants in the crude oil exploration and production companies is the crude oil that is entrapped with the effluents during treatment, conditioning and storage of crude oil. The oily waste (sludge) is semi-emulsified mixture of oxidized crude oil, water and miscellaneous solids (corrosion product, sand, silt and clay). These petroleum wastes and crude oil enter into the environment through the activities of petroleum extraction, refining, transportation, storage, bad practices (human error), accidents, illegal dumping of crude oil wastes, and leakages due to corrosion of tanks and pipelines as well as vandalization. The release of hydrocarbon pollutants into the environment whether accidentally or due to human error is the main cause of water and soil pollution. Fortunately, microorganisms are highly efficient and versatile in their ability to degrade hydrocarbons; (Odokuma and Dickson 2003; Head et al., 2006; Odokuma and Smith 2007; Adebuseye et al., 2007). Bacteria that biodegrade polynuclear aromatic hydrocarbons (PAHs), monoaromatic hydrocarbons and aliphatic hydrocarbons are readily isolated from the environment that have been contaminated with petroleum hydrocarbon (Chamkha et al., 2011; Manchola and Dussan, 2014). A reviewed list of bacterial genera has listed 79 genera that can utilize hydrocarbon as the sole source of carbon and energy (Head et al., 2006). The effect of hydrocarbon contaminants on human health and the environment can not be over emphasized. It is therefore important to reduce or eliminate the total petroleum hydrocarbon (TPH) content in petroleum sludge by applying desirable property of hydrocarbon-utilizing bacteria isolated or found in the contaminated environments. This study was therefore aimed at isolating and characterizing hydrocarbon-utilizing bacteria from petroleum sludge sample collected from crude oil processing plant.

MATERIALS AND METHODS

Sample collection: The petroleum oily sludge sample used in the study was collected from crude oil processing plant located at Obegi community in Ogbia-Egbama Ndoni Local Government Area of Rivers State, Niger Delta, Nigeria. Fresh sludge samples were collected directly from different sludge outlet valves in the plant and mixed together into a 500ml glass jar and covered. The sludge sample was transported in ice pack to the microbiology research...
laboratory of the University of the Port Harcourt for analysis. The sludge sample was preserved in the refrigerator at 4°C.

Reagents: The entire reagents used in this study were of analytical grade and were obtained from BDH chemicals limited, Poole England. Nutrient agar (NA), and MacConkey’s agar were obtained from International diagnostic group; England. Filter Paper (What man No.1) was obtained from WER Bauston Limited. The Bonny light crude oil used for hydrocarbon utilizing bacterial (HUB) test was obtained from Port Harcourt Refinery Company, Eleme, Rivers State Nigeria.

Enumeration of bacterial populations: The total heterotrophic bacterial (THB) and hydrocarbon utilizing bacterial (HUB) counts of the petroleum oil sludge were carried out by ten-fold serial dilution with normal saline. One gram of petroleum sludge was weighed into test tubes containing 10ml normal saline. Then 1ml was transferred from the stock into another test tube containing 9ml normal saline giving 10⁻¹ dilution. This was repeated up to 10⁻³. Aliquots of dilution (0.1ml) were inoculated onto nutrient agar (NA) plates in triplicates using spread plate method (APHA, 1998). The plates were incubated at 37°C for 24h. The HUB count was carried out in triplicates on mineral salt agar (MSA) of Mills (1978 as modified by Okpokwasili and Odukuma (1990). MSA was composed of 0.29g of KCl, 10g of NaCl, 0.42g of MgSO₄•7H₂O, 0.42g of NH₄NO₃, 1.25g of KH₂PO₄, 0.83g of K₂HPO₄ and 15g of agar all dissolved in one litre of distilled water and autoclaved at 121°C for 15mins. Aliquots of 0.1ml of dilution was inoculated onto MSA plates in triplicates by spread plate method (APHA, 1998). Sterile filter papers (What man No.1) saturated with Bonny light crude oil were aseptically placed on the inside lid of each plate and kept in an inverted position and incubated at 37°C for 48h. Plates were enumerated after incubation period. The filter paper saturated with crude oil served as a sole source of carbon and energy.

Isolation and identification of HUB: Culturable bacteria isolates from the HUB plates were sub-cultured onto NA plates and incubated at 28°C for 24h. Distinct colonies were further sub-cultured onto slant NA in Bijou bottles and incubated at 28°C for 24h. The NA slant cultures were stored in a refrigerator at 4°C and served as pure stock culture for subsequent characterization and identification of the isolates. Standard characterization tests were performed as described by Cheesbrough, (2006) which included Grams staining, motility, methyl-red, Vogues- Proskauer, indole, citrate utilization and sugar fermentation. The pure isolates were identified on the basis of their cultural, morphological and physiological characteristics (Cheesbrough, 2006).

Molecular Identification of Isolates: DNA extraction: DNA extraction was carried out by using a ZR fungal/ bacterial DNA Miniprep extraction kit obtained from Inquba, South Africa. The DNA extraction protocol was based on manufacturer’s instruction. Heavy growth of the pure six isolates subcultured on MacConkey’s agar plates were suspended in 200µl of isotonic buffer into ZR bashing bead lysing tubes, 750µl of Lysing solution was added. They were processed at maximum speed for 5mins. The ZR Bashing bead lysing tubes were centrifuged at 10,000 xg for 1min. Four hundred (400) µl of the supernatants were transferred into ZYMO-spin IV spin filter in collection tubes and centrifuged at 7000xg for 1min. The amount of 1,200µl of fungal/bacterial DNA binding buffer was added to each filtrate in the collection tubes bringing the final volume to 1,600µl, 800µl was then transferred into ZYMO-spin11C column in a collection tube and centrifuged at 10,000xg for 1min, the flow through were discarded from the collection tubes. The remaining volumes were transferred to the same ZYMO-spin and spun at 10,000xg for 1min. Two hundred (200) µl of the DNA pre-wash buffer were added to the ZYMO-spin 11C in new collection tubes and spun at 10,000xg for 1min followed by addition of 500µl of fungal/bacterial DNA wash buffer and centrifuged at 10,000xg for 1min. The ZYMO-spin 11C column were transferred to clean 1.5µl centrifuge tubes and 100µl of DNA elution buffer were added to the column matrix and centrifuged at 10,000xg for 30 seconds to elute the DNA. The ultra-pure DNA of each isolate properly labeled were then stored at -20°C for use. After extraction, the DNA samples were quantified using NANODROP (ND1000).

PCR amplification of 16S rRNA: The 16S of the rRNA genes of the isolates were amplified using the 27F and 1492R primers on a PCR System (9700 Applied Bio system thermal cycler) at a final volume of 25µl for 40 cycles. The PCR mix include: the x2 Dream tag master mix supplied by Inquba, South Africa (tag polymerase, DNTPs, MgCl₂, the primers at a concentration of 0.4M and extracted DNA as a template. The PCR conditions were as follows: initial denaturation, 95°C for 4mins; denaturation 95°C for 30s; Annealing, 52°C for 3mins. Then holding temperature at 4°C. The products were resolved on a 1% agarose gel at 120v for 15min and visualized on a
UV transilluminator (Cebron et al., 2008). The generated PCR products were dried and sent to Inquaba (South Africa) for purification and sequencing using an AB13500 genetic analyzer adopting the Bigdye Termination technique by Inquaba South Africa.

**Phylogenetic analysis:** The sequences were edited using the bio informatics algorithm Bioedit. Similar sequences were downloaded from the National Biotechnology Information Centre (NCBI) data base using Blast N, these sequences were aligned using cluster IX. The evolutionary history of the isolates was inferred using Neighbor- Joining method in MEGA 6.0 (Thompson et al., 1994). The bootstrap consensus tree inferred from 500 replicates was taken to represent the evolutionary of the taxa analyzed (Felsenstein, 1985)

**RESULTS AND DISCUSSION**

<table>
<thead>
<tr>
<th>Type of count</th>
<th>Value</th>
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<tbody>
<tr>
<td>THB</td>
<td>(2.5 \times 10^6) cfu/g</td>
</tr>
<tr>
<td>HUB</td>
<td>(2.0 \times 10^3) cfu/g</td>
</tr>
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</table>

**Fig. 1:** Agarose electrophoresis of the 16S gene bands of the isolates:

L: represents the 1kb ladder, 1-6 represents 16 S gene bands of the isolates.

**IDENTIFICATION OF ISOLATES WITH ACCESSION NUMBERS**

<table>
<thead>
<tr>
<th>HUB isolates identified using 16S rRNA gene nucleotide sequence(s)</th>
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<tbody>
<tr>
<td><em>Klebsiella pneumonia</em> strain B21                                           - SUB1917764B1  KX817218</td>
</tr>
<tr>
<td><em>Klebsiella pneumonia</em> strain ICB-C183                                       - SUB1917764B2  KX817219</td>
</tr>
<tr>
<td><em>Klebsiella oxytoca</em> strain BCNA1                                              - SUB1917764B3  KX817220</td>
</tr>
<tr>
<td><em>Klebsiella oxytoca</em> strain BC4                                               - SUB1917764B4  KX817221</td>
</tr>
<tr>
<td><em>Alcaligenes faecalis</em> strain IOU PMR                                          - SUB1917764B5  KX817222</td>
</tr>
<tr>
<td><em>Alcaligenes faecalis</em> strain AQ-1                                             - SUB1917764B6  KX817223</td>
</tr>
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</table>

The bacterial populations of the sludge sample are presented in Table 1. The enumeration of high population of THB \((2.5 \times 10^6)\) cfu/g and HUB counts of \(2.0 \times 10^3\) cfu/g respectively is an indication that the sludge sample could be degraded by the indigenous hydrocarbon-utilizing bacteria isolated from the oily sludge sample (Ojo, 2006; Manchola and Dussan, 2014). The enumeration of these bacterial populations from the oily petroleum sludge implies that the bacterial genera are able to utilize crude petroleum oil hydrocarbons as the sole source of carbon and energy. Manchola and Dussan (2014) in a study to evaluate the ability of seven strains belonging to *Lysinibacillus sphericus* and *Geobacillus* species isolated from an off-shore “sercina” oil field, to biodegrade TPH in the presence...
of toxic metals, their potential to produce biosurfactants, and their ability to improve the biodegradation rate, reported that the bacterial strains were able to utilize crude petroleum oil hydrocarbons as the sole source of carbon and energy. In addition, their ability to degrade crude oil was not affected by the presence of toxic metals such as chromium and arsenic. At the same time, the strains were able to reduce toxic metals concentration through biosorption processes. (Manchola and Dussan, 2014).

The cultural and colonial characteristics showed that the colonies were of various shapes and forms such as large, raised, moist and mucoid, while others were non-slime (non mucoid) on NA plates.

Morphological characterization of the isolates all showed Gram-negative, non-sporing, non-motile short rods. Biochemical reactions showed catalase positive, oxidase negative, indole negative, Vogus-Proskauer positive, citrate positive and various reactions to other biochemical reactions. The results suggested Klebsiella and Alcaligenes species.

The phylogenetic analysis of the 16S rRNA gene sequence results of the isolates produced an exact match during mega blast search for highly similar sequences from the NCBI non-redundant nucleotide (nr/nt) database. The 16S rRNA of the isolates showed percentage similarities to other species at 99% being most closely related to Klebsiella pneumonia, Klebsiella oxytoca and Alcaligenes faecalis. The evolutionary distances computed using Jukes-cantor methods were in agreement with the phylogenetic placement of the 16S rRNA of the isolates within the type strains of the following: Klebsiella pneumonia strain B2 (gi: 922317936), Klebsiella pneumonia strain IC-B18 (gi: 908478837), Alcaligenes faecalis strain IOU PMR (gi: 686028963), Alcaligenes faecalis strain AQ-1 (gi: 816845513), Kebisiella oxytoca strain BCNAI (gi: 846993354) and Kebisiella oxytoca strain BC4 (gi: 884060138) respectively.

Previous studies have reported the involvement of many genera of bacteria in hydrocarbon degradation and use them as a source of carbon and energy (Head, 2006; Bhattacharyya, 2003), but there are few reports that suggest that enteric bacteria especially those of genera Klebsiella, Enterobacter, Escherichia coli and Hafnia have the ability to degrade hydrocarbons (Diaz, et al., 2001; Serma et al., 2004). Klebsiella pneumonia is ubiquitous as it is found in mammals and ecological environment. It is an important cause of human infections. Infections or diseases are usually nosocomial or hospital-acquired (Podschum and Ullmann, 1998). The results of isolation and identification of Klebsiella Pneumonia and K. oxytoca to the strain levels in this study, agree with previous studies that implicated the same organisms in petroleum hydrocarbon degradation. Chamkha et al., (2011) reported the isolation and characterization of Klebsiella oxytoca strain degrading crude oil from Tunisian off-shore oil fields, similarly, Rodrigues et al., (2009) had earlier reported diversity of hydrocarbon-degrading Klebsiella strains isolated from hydrocarbon-contaminated estuaries. The oil-degrading Klebsiella isolates obtained from the estuary were closely related to Klebsiella pneumonia and Klebsiella orinthinmolytica. The isolates demonstrated a substantial degree of catabolic plasticity for hydrocarbon degradation. (Diaz, 2001; Sarma et al., 2004). Alcaligenes faecalis isolated from the oily sludge sample, in this study, is an aerobic gram-negative rod-shaped motile organism that occur or inhabit soil and water (Austin et al., 1981). Bharali et al., (2001) explored the use of Alcaligenes faecalis to promote biodegradation of petroleum hydrocarbons. A. faecalis produces biosurfactant compounds that increase the hydrophobicity of the cell surface during growth on hydrocarbons that enhances the contact with the hydrocarbons and as a result increases hydrocarbon degradation. In the study by Bharali et al., (2001), the capacity to produce biosurfactant was demonstrated by growing A. faecalis in salt media with a variety of hydrophobic substances (fuel, kerosene, crude oil) as the carbon source. Under different substrate concentrations surface tension and the rate of biosurfactant production were measured. It was found that biosurfactant produced by A. faecalis possessed high surface activity, decreasing surface tension adequately to allow for degradation by the microorganisms (Bharali et al., 2001).

In conclusion, from the findings of this study, it suggest that the organisms isolated and identified as Klebsiella pneumonia, K. oxytoca and Alcaligenes faecalis of different strains have the potential to degrade hydrocarbon pollutants when applied in the environmentally friendly technology clean-up (bioremediation) of hydrocarbon contaminated sites in the Niger Delta of Nigeria.

REFERENCES


