Molecular characterization of crude oil degrading bacterial isolates from polluted soils and cow dung

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Abstract—Crude oil contamination of agricultural soil is frequent in the Niger-Delta Region of Nigeria and can devastate the soil thereby, negatively affecting the socio-economic lives of the people. This study aimed to characterize twelve bacterial isolates with potential for crude oil degradation using conventional and molecular tools. Isolates with potential for crude oil degradation were selected from among the crude oil degrading bacteria obtained from petroleum contaminated agricultural soils and Cow dung all in Ondo State, Nigeria. The identities of the isolates were confirmed via morphological and biochemical characterization and thereafter the CTAB method was used to prepare the DNA. PCR amplification of 16S rRNA gene of isolates was carried out using universal primers for bacteria. The PCR products were then purified using ethanol precipitation and thereafter sequenced with automated DNA sequencing machine. The sequence data were compared with gene sequences in GenBank database (NCBI) using a BLAST search to find closely related sequences. Phylogenetic analyses of 16S rRNA gene sequences were examined in order to determine the evolutionary relatedness of the isolates. Results revealed eight (8) gram positive bacteria consisting of Staphylococcus hominis, Geobacillus sp., Lactobacillus plantarum and four (4) different species of Bacillus, while the gram negative bacterial isolates were Brevundimonas diminuta, Klebsiella oxytoca, Escherichia coli and Enterobacter tabaci with 83% to 100% ribosomal RNA homology. Crude oil degrading bacteria characterized in this study can be developed as inoculums with high survival and activity to bioaugment the degradation of crude oil polluted agricultural soil.

Keywords—Bioremediation, Cow dung, Crude oil degrading bacteria, Polluted soils, 16S rRNA gene.

I. INTRODUCTION

Pollution of the environment due to discharge of petroleum or its derivatives into the environment is a concern to all and sundry including government, environmental researchers and residents of oil producing areas, particularly, the Niger Delta region of Nigeria. Crude oil spill, no matter its source, quantity and size (minor, medium, major or disaster) is potentially harmful to the environment and all forms of biomass (including indigenous micro flora and micro fauna [1, 2]. Among remediation techniques, bioremediation which relies on the use of microorganisms with desired metabolic capabilities to detoxify many hydrocarbon pollutants is most preferred. This method of biological mineralization of petroleum hydrocarbon involving primarily bacteria, yeast and mold is applicable over large area [3], ecosystem friendly, cost effective and non-invasive relative to the physical and chemical methods [4]. Microbial degradation of hydrocarbons by natural population of microorganisms is the major and ultimate natural mechanism by which petroleum hydrocarbon pollutants can be cleaned up from the environment [5]. These microorganisms use petroleum hydrocarbon as carbon and energy source for cellular activities. Microbial remediation of a hydrocarbon-contaminated site is accomplished with the activities of diverse group of microorganisms, particularly the indigenous bacteria present in the soil [6]. Hydrocarbon degrading bacteria and fungi are widely distributed in
marine, freshwater and soil environments. Typical bacteria groups already known for their potentials to degrade hydrocarbons are *Pseudomonas* sp., *Marinobacter* sp., *Micrococcus* sp., *Acinetobacter* sp., *Alcaligenes* sp., *Bacillus* sp., *Enterobacter* sp. [6,7]. Molds belonging to the genera *Aspergillus* sp., *Penicillium* sp., *Fusarium* sp., *Amphotecota* sp., *Paecilomyces* sp., *Talaromyces* sp. and the yeasts *Candida* sp., *Yarrowia* sp., and *Pichia* sp. have been implicated in hydrocarbon degradation [8]. The process of hydrocarbon biodegradation in soil is however limited by microorganism type and population among other factors [2, 9]. The removal of contaminated by natural attenuation is slow because degrading microbes in soil is only about 10% of the total heterotrophic population [9]. Many different microorganisms such as *Pseudomonas* species and the white rot fungus, *Phanerochaete chrysosporum* are being marketed for use in bioremediation [2]. *Klebsiella pneumoniae*, *Pseudomonas aeruginosa*, *Escherichia coli*, *Proteus mirabilis*, *Alcaligenes faecalis*, *Bacillus species*, *Brevundimonas diminuta*, *Pseudomonas pseudomallei*, *Staphylococcus hominis*, *Enterobacter* sp., *Aspergillus* sp., *Scopulariopsis brevicaulis*, *Gliocladium* sp., *Trichoderma* sp. *Pseudolichenes* variotii, *Trichophyton mentagrophytes*, *Candida parapsilosis*, *Kodamaea ohmeri* were crude oil degrading bacteria and fungi associated with cow dung and some crude oil contaminated and uncontaminated soils [10]. The studies however do not focus on the molecular identification and genetic relationship among these isolates. Therefore, as a step towards developing inoculum with high survival and activity to bioaugment crude oil biodegradation, it is important to identify the organisms to reasonable level of precision using molecular tools. In this study, bacterial isolates from crude oil polluted soil and Cow dung with potential for crude oil degradation were identified using conventional and molecular tools.

II. MATERIALS AND METHODS

2. 1 Sample collection

Twelve (12) crude oil degrading bacteria previously with potential for crude oil degradation were selected from among the crude oil degrading bacteria obtained from petroleum contaminated agricultural soils of Awoye, Orioke-Iwamimo, Igodan-Lisa, Oba-Ile and Ido-Ani and Cow dung from Shasha, all in Ondo State, Nigeria. These environmental sources had varying amount of total petroleum hydrocarbon (TPH) as evidence of hydrocarbon pollution [10] and their crude oil biodegradation potentials were also reported [11].

2. 2 Sterilization of materials

All media, distilled water and diluents were sterilized by autoclaving at 121°C at 15 psi for 15 minutes. Glass wares were sterilized in a hot air oven at 160°C for 2 hours.

2. 3 Purification and Characterization of the crude oil degrading bacterial isolates from soil and cow dung

The isolates were purified by repeated streaking on nutrient agar (NA) and Bushnell-Hass (MSM) incorporated with 1.5% agar and sterile crude oil (2%) and incubated at 28 ± 2°C for 48 hours and 14 days (for growth in oil) respectively. Pure and distinct colonies were then inoculated into their respective nutrient agar slant, labeled and incubated at 28 ± 2°C for 48 hours after which they were maintained on nutrient agar (NA) at 4°C for further use. The cultural, morphological and biochemical characteristics were also confirmed. These biochemical tests include gram staining, catalase, oxidase, nitrate reduction, citrate utilization, sugar (glucose, maltose, mannitol, sucrose and lactose) fermentation using [12, 13] as standard references.

2. 4 Molecular characterization

2. 4. 1 DNA extraction of bacterial isolates using CTAB method

DNA isolation is a routine procedure to collect DNA for subsequent molecular or forensic analysis. In this research, Cetyl Trimethyl Ammonium Bromide (CTAB) method of DNA extraction from microbes was used. Twelve bacterial isolates with potential for crude oil degradation were selected from among previously screened isolates from agricultural soils and Cow dung. Overnight grown broth cultures of isolates were respectively transferred to endopod tubes and spun down for 2 minutes at 14,000 rpm. The supernatant was discarded and the DNA was extracted using the Cetyl Trimethyl Ammonium Bromide (CTAB) method [14]. The DNA pellets were re-suspended in 100 µl of sterile distilled water. DNA concentration of samples were measured on spectrophotometer at 260 nm and 280 nm and the genomic purity were determined. Agarose gel electrophoresis was used to determine the quality and integrity of the DNA by size fractionation on 1.0% agarose gel and visualized on UV light source.

2. 4. 2 Polymerase Chain Reaction (PCR) analysis

Polymerase chain reaction (PCR) analysis was performed with MJ Research Thermal Cycler (PTC-200 model) using 16S (forward and reverse) universal primer for bacteria. The sequence for the 27F (5'-AGAGTTTGTGATCAGGCTAG-3') and 1492R (5'-CGGTACCTTGTTACGACTT-3') [15]. The PCR mix comprises of 1µl of 10X buffer, 0.4 µl of 50mM MgCl₂, 0.5µl of 2.5mM dNTPs, 0.5µl 5mM forward primer, 0.5µl
of 5mM reverse primer, 0.05µl of 5units/µlTaq with 2µl of template DNA and 5.05µl of distilled water to make-up 10µl reaction mix. The PCR profile used was initial denaturation temperature of 94°C for 3minutes, followed by 30 cycles of 94°C for 60 seconds, 56°C for 60 seconds, 72°C for 120 seconds and the final extension temperature of 72°C for 5 minutes and the 10°C hold.

2. 4. 3 Gel Electrophoresis of 16S rDNA products
The PCR products were analyzed on a 1% TAE agarose gel and stained with ethidium bromide at 80 V for 45-60 min. The bands were thereafter, visualized using ultraviolet (UV) light trans - illumination and photographed with a digital imaging system (Kodak UV illumination System).

2. 4. 4 Purification of PCR products
The amplicon was further purified before the sequencing using 2M sodium acetate washing techniques. To about 10 µl of the PCR product was added 1µl 2M sodium acetate pH 5.2, followed by 20 µl absolute ethanol. This was kept at -20°C for 1hour, spun at 10,000rpm for 10 minutes, and thereafter washed with 70% ethanol and air- dried. This was then re-suspended in 5µl sterile distilled water and kept at 4°C for sequencing.

2. 4. 5 DNA Sequencing analysis
The 16S rRNA gene products (forward and reverse primers) of isolates were sequenced. The clean PCR products were subjected to cycle sequencing in both direction using universal primers. The PCR mix used for the sequencing include 0.5µl of BigDye Terminator Mix, 1µl of 5X sequencing buffer, 1µl of 16S forward primer with 6.5µl distilled water and 1µl of the PCR product making a total of 10µl. The PCR profile for sequencing is a rapid profile. The initial rapid thermal ramp to 96°C for 1minute followed by 25 cycles of rapid thermal ramp to 96°C for 10 seconds, Rapid thermal ramp to 50°C for 5 seconds and Rapid thermal ramp to 60°C for 4 minutes, then followed by Rapid thermal ramp to 4°C and hold forever. This was then re-suspended in 5µl sterile distilled water and kept at 4°C for sequencing running. The cocktail mix is a combination of 9µl of Hi Di Formamide with 1µl of purified sequence making a total of 10 µl. The samples were loaded on the machine (ABI 3100) and the data in form A, C, T, and G was released.

2. 4. 6 Construction of Phylogenetic Tree
The nucleotides sequences obtained were compared with other nucleotides sequences using BLASTn tools of the National Centre for Biotechnology Information (NCBI). The UPGMA method was used to infer the evolutionary history [16]. The software, MEGA7 was used to align all the sequences obtained in this study while all positions containing gaps and missing data were eliminated. The Maximum Composite Likelihood method was used to compute the evolutionary distances [17]. Thereafter, the evolutionary analyses were conducted in MEGA7 [18].

III. RESULTS AND DISCUSSION
3. 1 Characterization of crude oil degrading bacterial isolates from soils and cow dung
In the quest to proffer solution to cleaning up the environment polluted with crude oil using microorganisms, twelve potential crude oil degrading bacteria that could be employed in bioremediation techniques were selected from polluted soils and cow dung samples for characterization. Tables 1 and 2 show the detailed morphological and biochemical characteristics of the twelve crude oil degrading bacteria from polluted soils and cow dung. Results indicated that the isolates showed different morphology and responses to biochemical tests isolates SS1A, SS1B, SS2A, SS2E, SS3C, SS4A, SS5A and SS5C were tentatively identified as Brevundimonas diminuta, Lactobacillus plantarum, Klebsiella sp., Bacillus megaterium, Bacillus subtilis, Bacillus sp., Bacillus sp. and Geobacillus sp respectively for the soil samples (Table 1). Cow dung bacterial isolates (Table 2) CD6A, CD6E, CD6J, and CD6K were similarly identified as Escherichia coli, Enterobacter tabaci, Staphylococcus hominis and Geobacillus sp. Interestingly, only Staphylococcus hominis obtained from cow dung was coccii shaped. Thus, the conventional techniques of identification of bacteria revealed four Bacillus species, Brevundimonas sp. and Lactobacillus sp. Klebsiella sp. and Geobacillus sp from polluted soils (table 1) and Escherichia coli, Enterobacter sp., Staphylococcus sp. and Geobacillus sp from cow dung samples (Tables 2). The results obtained agree with the characterization pattern of [12, 13].
Table 1: Characteristics of crude oil degrading bacteria isolated from soil samples

| Characteristics | Soil isolates |
|-----------------|---------------|
|                 | SS1A | SS1B | SS2B | SS2E | SS3C | SS4A | SS5A | SS5C |
| **Morphological** |      |      |      |      |      |      |      |      |
| Colony          | Circular | Spherical | Spherical | Cylinder | Elliptical | Undulated | Elliptical |      |
| Colour          | None | Whitish | None | Yellow | None | None | Brown | None |
| Edge            | Entire | Lobate | Regular | Entire | Lobate | Entire | Entire | Entire |
| Surface         | Smooth | Rough | Smooth | Smooth | Smooth | Smooth | Smooth | Smooth |
| Cell shape      | Rod | Rod | Rod | Rod | Rod | Rod | Rod | Rod |
| Gram stain      | - | + | - | + | + | + | + | + |
| Motility        | + | - | - | + | + | + | + | + |
| **Biochemical** |      |      |      |      |      |      |      |      |
| Catalase        | + | - | + | - | + | + | + | + |
| Oxidase         | + | - | - | - | - | - | - | - |
| Spore staining  | - | - | - | + | + | + | + | + |
| Citrate reaction| - | + | + | + | - | - | + | + |
| MR              | + | - | - | + | - | - | + | + |
| VP              | - | - | + | - | + | + | - | - |
| Nitrate reduction| - | - | + | + | + | + | + | + |
| **Fermentation** |      |      |      |      |      |      |      |      |
| Glucose         | NC | A | AG | A | A | A | A | A |
| Maltose         | NC | A | AG | A | A | NC | A | NC |
| Mannitol        | NC | A | AG | A | A | A | A | A |
| Sucrose         | NC | A | AG | A | A | NC | NC | NC |
| Lactose         | NC | A | NC | A | G | NC | A | NC |
| **Possible Organism** | Brev. diminuta | Lactobacillus plantarum | Klebsiella oxytoca | Bacillus egaterium | Bacillus subtilis | Bacillus aryabhattai | Bacillus sp. | Geobacillus sp. |

Legend: A= Acid production only, AG = Acid & Gas production, G = Gas production, NC = No change, + = positive, - = negative. SS1A and SS1B are isolates from Awoye soil sample, SS2B and SS2E are isolates from Orioke-Iwamimo soil, SS3C is isolate from Igodan-Lisa soil, SS4A is isolate from Oba-Ile soil, SS5A and SS5C are isolates from Ido-Ani soil.

The bacterial isolates in this work belong to both the gram positive and gram negative groups. These findings corroborate the report that both gram positive and positive bacteria have been implicated in the mineralization of hydrocarbon pollutants [19]. This study also revealed that gram positive crude oil degrading bacteria are ubiquitously distributed, diverse and dominant in all the soils samples and cow dung. This finding deviates from the reports that gram positive bacteria if detected in bioremediation are never diverse and dominant [20]. Also, the ability to isolate high number of certain crude oil degrading microorganisms from these environments is commonly taken as evidence that those microorganisms are the most active degraders in that environment and can be used in the bioremediation of petroleum oil contaminated sites [21, 22]. This probably suggests that these organisms are able to adapt to different hydrocarbons and varying environmental parameters since samples were collected from different locations. B.
*diminuta* belongs to the phylum Proteobacteria, class Alphaproteobacteria and order Caulobacter. *B. diminuta* has been reported to have considerable ability for bioremediation following a bioremediation study of seawater contaminated with diesel in China [23]. *Staphylococcus hominis*, a gram negative and coagulase negative bacterium, was also isolated from oil contaminated soil [24]. It was observed from this work that the dominant bacterial species were *Bacillus*, belonging to the phylum Firmicutes. The abundance of *Bacillus* sp. in the crude oil polluted soils shows that they are indigenous to the soil. The presence of *Bacillus* species have been reported by different researchers in crude oil contaminated soils with the ability to degrade oil [25], thereby, using crude oil as sole source of carbon and energy [22, 25]

| Characteristics | Cow dung isolates |
|-----------------|-------------------|
| **Morphological** |                  |
| Colony          | Circular          |
| Colour          | Greyish           |
| Edge            | Entire            |
| Surface         | Smooth            |
| Cell shape      | Rod               |
| Gram stain      | -                 |
| Motility        | +                 |
| **Biochemical**  |                  |
| Catalase        | +                 |
| Oxidase         | -                 |
| Spore staining  | -                 |
| Citrate reaction| -                 |
| MR              | +                 |
| VP              | -                 |
| Nitrate reduction| +               |
| **Sugar Fermentation** |            |
| Glucose         | AG                |
| Maltose         | AG                |
| Mannitol        | AG                |
| Sucrose         | NC                |
| Lactose         | AG                |
| **Possible Organism** |          |
| *Escherichia coli* | Entrobactersp     |
| *Staph.hominis* | Geobacillus      |

**LEGEND:** A= Acid Production only, AG = Acid & Gas Production, G = Gas Production, NC = No Change, + = Positive, - = Negative , CD6A, CD6E, CD6J and CD6K are isolates from cow dung
3.2 Molecular identity of the bacterial isolates

Plate 1 shows the agarose gel electrophoresis of twelve bacterial isolates subjected to molecular identification using 16S universal primer. Fig. 1 shows the phylogenetic tree of the bacterial isolates while table 3 shows the nearest relative, accession numbers and the percentage homology of the isolates. The blasting of the sequence also revealed that there are eight types of bacteria genera present in the samples. These include four *Bacillus* species, *Brevundimonas diminuta*, *Escherichia coli*, *Enterobacter* sp. *Geobacillus*, *Staphylococcus hominis*, *Lactobacillus plantarum* and *Klebsiella oxytoca*. All the bacteria obtained in this work had between 83% and 100% homology with the bacterial isolates deposited in the NCBI data.

Plate 1: Agarose gel electrophoresis of DNA of the bacterial isolates using 16S universal primer (100 basepair ladder)

Legend: M = Molecular Weight Marker/Ladder (1kb)

**Fig. 1: Phylogenetic tree based on partial 16S rRNA gene sequences analysis indicating the relationships between crude oil-degrading bacteria from polluted soils and cow dung and the related strains from the Gene Bank.**
Table 3: Complete nucleotides blast of 16S rRNA genes from crude oil degrading bacteria

| Isolate Code | Source       | Organisms                            | Accession Number | (%) Identity |
|--------------|--------------|--------------------------------------|------------------|--------------|
| SS1A         | Soil         | *Brevundimonas diminuta* strain VBE65 | MG027643.1       | 100          |
| SS1B         | Soil         | *Lactobacillus plantarum* strain CAU9791 | MF425505.1       | 83           |
| SS2B         | Soil         | *Klebsiella oxytoca* strain U760     | K4572971.1       | 99           |
| SS2E         | Soil         | *Bacillus megaterium* strain Edgemont-007A | MF965192.1       | 99           |
| SS3C         | Soil         | *Bacillus subtilis* strain 3-2       | JX051359.1       | 99           |
| SS4A         | Soil         | *Bacillus aryabhattai* strain CN13-5 | MH762878.1       | 100          |
| SS5A         | Soil         | *Bacillus* sp. strain AHBR19         | KF241532.1       | 100          |
| SS5C         | Soil         | *Geobacillus stearothermophilus* strain N23-E | KF768847.1       | 100          |
| CD6A         | Cow dung     | *Escherichia coli* strain 42         | MH671423.1       | 100          |
| CD6E         | Cow dung     | *Enterobacter tabaci* UPM18          | MH794127.1       | 99           |
| CD6J         | Cow dung     | *Staphylococcus hominis* strain FC1738 | MH665980.1       | 100          |
| CD6K         | Cow dung     | *Geobacillus* sp. strain BD63        | MF767892.1       | 94           |

Legend: SS1A and SS1B are isolates from Awoye soil sample, SS2B and SS2E are isolates from Orioke-Iwamimo soil, SS3C is isolate from Igodan-Lisa soil, SS4A is isolate from Oba-Ile soil, SS5A SS5C are isolates from Ido-Ani soil, CD6A, CD6E, CD6j and CD6K.

The molecular techniques used in this study to identify bacteria were based on the conserved sequence of the 16S rRNA genes that were amplified by PCR [26]. The molecular weights of the PCR amplification fragments obtained were about 1.6kb each (Plate 1). The results from the conserved sequence of the 16S rRNA coupled with the nucleotide sequences revealed that the twelve bacteria isolates were closely related to *Brevundimonas diminuta* strain VBE65, *Lactobacillus plantarum* strain CAU9791, *Klebsiella oxytoca* strain U760, *Bacillus megaterium* strain Edgemont-007A, *Bacillus subtilis* strain 3-2, *Bacillus aryabhattai* strain CN13-5, *Bacillus* sp. strain AHBR19, *Geobacillus stearothermophilus* strain N23-E, *Escherichia coli* strain 42, *Enterobacter tabaci* UPM18, *Staphylococcus hominis* strain FC1738 and *Geobacillus* sp. strain BD63. The high percentage (99 - 100%) similarities observed between the 16S rRNA gene, partial sequences of ten of the bacterial isolates and previously identified bacteria in the GeneBank, indicates that they are homologous to each other (Table 3). The Phylogenetic tree, based on partial bacterial 16S

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rRNA gene sequence analysis, indicates the relationships between the isolated microbes and the related genera of the phylum Proteobacteria and Firmicutes. The evolutionary history was inferred by using the statistical method of UPGMA. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) is shown next to the branches. The higher the bootstrap value, the more reliable the phylogenetic analysis. The code indicates the microorganisms used in this study. The strains of reference species are indicated (Fig. 2).

IV. CONCLUSION

This study has been able to characterize crude oil degrading bacteria from environmental sources (soil and cow dung) using both conventional and molecular techniques. The study revealed the molecular identity of crude oil degrading bacteria from environmental sources that can be harvested as biomass and marketed for use as inoculums to enhance bioremediation of polluted sites in the future.

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