Determination of the Bacterial Community Structure in a Crude oil-inundated Tropical Soil using Next Generation Sequencing Technique

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8 ABSTRACT

9 Aim: To identify the community composition of a crude oil impacted soil in Gbarain kingdom of
 10 Bayelsa State, Nigeria
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Study design: A crude oil-impacted soil sample (0-10 cm depth) was collected from Etelebuo-Ogboloma, flow station in Yenagoa L.G.A. of Bayelsa State and taken to the laboratory for various microbiological analyses.

Place and duration of study: The study was carried out at the Environmental Microbiology
 Laboratory, University of Port Harcourt, Nigeria, for 35days.

17 Methodology: Cultural morphology of the isolates was studied based on their physical appearance 18 such as colour, shape, size, elevation and margin. While catalase test, oxidase test, indole test, 19 motility test, Methyl-Red Voges-Proskauer's (MRVP) test and citrate utilization test were used for the 20 biochemical identification of the isolates. Deoxyribonucleic acid (DNA) eextraction from crude oil 21 polluted soil sample was performed using ZYMO soil DNA extraction Kit (Model D 6001, Zymo 22 Research, USA) following the manufacturer's instructions. DNA sequencing was performed by the 23 Next Generation Sequencing Technique to determine the nucleotide sequence of all microorganisms 24 present in the soil sample using sequencing primer -16S: 27F: 5'-GAGTTTGATCCTGGCTCAG-3' and 25 518R: 5'- ATTACCGCGGCTGCTGG-3'.

Results: Five different bacterial genera were isolated and identified using the cultural techniques, and they include *Acetobacter* sp., *Pseudomonas* sp., *Arthrobacter* sp., *Bacillus* sp. and *Micrococcus* sp. However, the molecular characterization revealed that the soil was mainly dominated by the Alphaproteobacteria (54.64%), followed by the Actinobacteria (9.67%), Gammaproteobacteria (6.55%), Betaproteobacteria (2.27%) and Bacilli (0.95%) as well as Clostridia (0.34%); as the most dominant class of bacteria. The unknown group accounted for 25.28%. A total of one hundred and four (104) diverse bacterial species were identified, in the overall metagenomics.

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34 Conclusion: This study has shown the bacterial community composition of the crude oil polluted soil 35 obtained from Gbarain Kingdom. These findings are fundamental in understanding the biological fate 36 of crude oil in these oil rich regions. 37 Keywords: Metagenomics; bacterial composition; Gbarain Kingdom; crude oil.

38 1.0 INTRODUCTION

Crude oil spill has remained a major source of environmental pollution in the Niger Delta area of Nigeria. These spills impact on the environment at varying degrees, depending on the volume spilled and physicochemical properties of the oil. The deleterious effects of crude oil contamination on the flora and fauna of the impacted media usually result in biodiversity loss, as exposure to crude oil fractions reduces bacterial population in the affected media, leading to loss in species diversity [1].

44 Bioremediation has remained an effective impact mitigation strategy (alongside other physical and 45 chemical methods of site remediation). The process of bioremediation involves the interplay of 46 various factors meant to achieve an efficient breakdown of these highly complex crude oil products in 47 the environment, in which case the synergistic activities of microbial consortium are required; where 48 hydrocarbonoclastic and hydrocarbon degrading microorganisms work together [2]. Bacillus, 49 Pseudomonas, Rhodococcus, Arthrobacter and Corynebacterium spp. are a group or bacterial 50 consortium having the ability to degrade hydrocarbons [3]. However, for bioremediation to be very 51 effective, the involvement of well adapted species to the prevailing environmental/site conditions is 52 very necessary with the following hydrocarbonoclastic bacteria; Arthrobacter, Flavobacterium, 53 Sphingomonas (a novel Pseudomonas sp), and Pseudomonas spp., well known for their adaptation to 54 petroleum contaminated environments [4]. The use of well adapted species helps in facilitating a 55 better and faster rate of hydrocarbon biodegradation in the contaminated site. However, not all 56 microorganisms found in the environment can degrade a particular hydrocarbon due to genetic 57 factors. Various authors have reported the efficacy of these adapted species. For instance, [5] 58 conducted a plasmid curing experiment and established that Nitrosomonas and Nitrobacter degraded 59 crude oil by the aid of crude oil degrading plasmid. Also, Staphylococcus sp. has been shown to have 60 the ability to utilize drilling fluid base oil and is therefore, a good agent for the remediation oil from 61 drilling fluid [6]. Fungal species such as Candida sp. and Schizosaccharomyces pombe have been 62 also found to be involved in the biodegradation of hydrocarbon [7; 8].

Recent advancement in bioremediation studies relies on molecular methods. These methods are useful in characterizing microorganisms associated with biodegradation of crude oil contaminated sites or media. The advantages of these molecular techniques are owed to their ability to identify autochthonous bacterial group *in situ* and also provide a rapid as well as efficient non-culture dependent methods of studying the bacterial composition (structure and function) of a site undergoing bioremediation [9].

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Several molecular techniques are available to study microbial communities in nature, especially the non-culturable ones which have not been identified in the laboratory via cultural approach. Comparative studies and analysis involving culture dependent and molecular metagonomic approach have revealed that only about 1% of the total microorganisms are amenable to culture [10]. Currently, nucleic acids, lipids, and proteins are harnessed to provide primary information that aid in the identification of these uncultured bacterial groups. Whole genomes or selected genes analysis which

76 includes 16S for prokaryotes and 18S rRNA for eukaryotes are molecular techniques that are non-77 culture dependent. Analytical studies of these genomes help to group microorganisms into three (3) 78 primary categories: two prokaryotic groups such as bacteria and archaea as well as one eukaryotic 79 group (eukarya) [11]. The development of techniques to characterise or identify functional diversity as 80 well as phylogenic differences has been of enormous importance in microbiology. Approaches such 81 as partial community analysis and whole community analysis are methods used to probe into bacterial 82 genetic properties which help in identifying microbial population based on their structure as well as 83 functional attributes. Modern molecular techniques have made way for an increased understanding of 84 bacterial diversity as well as functionality during crude oil degradation.

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Sequence-based and function-based sequencing are two conventional paths metagenomic screening has followed. The analysis of genome sequence data that has been recovered from the environment is motivated by many objectives, which include the establishment of gene inventories and natural product discovery [12].

91 Various researchers have used different molecular techniques to characterise and identify functional 92 genes in hydrocarbon degradation. However, the study of microbial diversity is somewhat complex 93 and requires several approaches, used in combination. This will provide more useful information 94 regarding diversity of the organisms. Owing to the paucity of information regarding the bacterial 95 community composition of hydrocarbon impacted sites in Gbarain kingdom, Bayelsa State, Nigeria, 96 this study was therefore carried out to identify the community composition using metagenomics and 97 culture dependent techniques. Data from this investigation could provide useful information for 98 potential bioengineering of the species through targeting specific genes of interest for efficient 99 bioremediation of crude oil contaminated sites.

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101 2.0 MATERIALS AND METHODS

102 2.1 Soil Sample Source and Collection

A crude oil-impacted soil (0–10 cm depth) was collected from Etelebuo-Ogboloma, flow station in
 Yenagoa L.G.A. of Bayelsa State using an Elderman auger into a Ziploc bag. Thereafter, the samples
 were kept in an ice box and immediately transported to the laboratory for microbiological analyses.

2.2 Culture-dependent Isolation, characterization and identification of hydrocarbon utilizing bacteria.

- **2.2.1 Place and duration of study:** The study was carried out at the Environmental Microbiology
 Laboratory, University of Port Harcourt, Nigeria, for 35days.
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111 **2.2.2** Isolation of bacterial strains from soil sample

- 112 The soil sample was diluted using 10-fold serial dilution, which was carried out to a dilution of 10⁻⁵ by
- 113 weighing 1 g of the soil sample into a sterile test tube containing 9ml of sterile physiological saline.

116 The heterotrophic bacteria were isolated by spreading 0.1 ml of the diluted soil sample on Nutrient Agar (NA) plates and incubated at 35 °C for 24hours. In the same manner, the hydrocarbon utilizing 117 118 bacteria were isolated by culturing the diluted soil samples on Mineral Salts Agar (MSA). A 119 Whatman's filter paper was saturated with crude oil and placed on the lid of each glass Petri dish 120 using sterile forceps. The crude oil served as the sole source of hydrocarbon (that is carbon and 121 energy source for the hydrocarbon utilizers). The inoculated Mineral Salts Agar (MSA) plates were 122 inverted and placed over the lid containing the saturated filter paper, and incubated at room 123 temperature for seven (7) days.

124 **2.2.3** Morphological identification of isolates.

Representative colonies of the different morphological types that appeared on the plates after incubation were carefully picked with a sterile inoculating loop and sub-cultured to obtain pure cultures. This was done by streaking aseptically, onto freshly prepared nutrient agar plates. Cultural morphology of the isolates was studied based on their physical appearance such as colour, shape, size, elevation and margin

130 **2.2.4 Biochemical characterization of isolates**

Isolated colonies were further purified by sub-culturing and identified using standard biochemical test
 methods [13].

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134 2.3 Molecular Analyses

135 2.3.1 Deoxyribonucleic acid (DNA) extraction from soil sample

136 Deoxyribonucleic acid (DNA) extraction from crude oil polluted soil sample was performed using 137 ZYMO soil DNA extraction Kit (Model D 6001, Zymo Research, USA) following the manufacturer's 138 instructions. According to this method, genomic DNA was extracted by weighing out 0.25 grams of 139 soil sample using an analytical balance (Ohaus, Germany). The sample was then added into a ZR 140 Bashing Bead [™] lyses tube followed by the addition of 750 µl lyses solution to the tube. The content 141 of the 2 ml tube was disrupted by mixing in a vortex mixer at maximum speed for 5 minutes. The ZR 142 Bashing Bead TM lyses tube was centrifuged in a micro centrifuge at \leq 10,000 x g for 1 minute. After 143 this, 400 µl of the filtrate was added to a Zymo-Spin ™ IV spin filter in a collection tube and 144 centrifuged at 7,000 rpm for 1 minute. This was followed by the addition of 1,200 µl of DNA binding 145 buffer to the filtrate in the collection tub after which 800 µl of the mixture from above was added to a 146 Zymo-Spin ™ IIC column in a collection tube and centrifuged at 10,000 x g for 1 minute. Flow through 147 from the collection tube was discarded and this particular step was repeated with the remaining 148 filtrate. This was followed by the addition of 200 µl of DNA pre-wash buffer into the Zymo-Spin ™ IIC 149 Column in a new collection tube and centrifuged at 10,000 x g for 1 minute, and then 500 µl DNA 150 wash buffer was added to the Zymo-Spin ™ IIC column and centrifuge at 10,000 x g for 1 minute. The 151 Zymo-Spin ™ IIC column was transferred into a clean 1.5 ml micro centrifuge tube and 100 µl DNA 152 elution buffer was directly added to the column matrix. This was centrifuged at 10,000 x g for 30 153 seconds to elude the DNA. The eluded DNA was transferred into a filter unit of Zymo-Spin ™ IV-HRC 154 Spin Filter in a clean 1.5 ml micro centrifuge tube and centrifuged at exactly 8,000 x g for 1 minute. 155 The filtered DNA was then used for PCR and DNA sequencing.

156 2.3.2 DNA Sequencing

157 All sequencing analysis was carried out at Ingaba Biotechnical Pty Ltd, South Africa. DNA sequencing 158 was performed by Next Generation Sequencing Technique to determine the nucleotide sequence of 159 all microorganisms present in the soil sample using sequencing primer -16S: 27F: 5'-160 GAGTTTGATCCTGGCTCAG-3' and 518R: 5'- ATTACCGCGGCTGCTGG-3'. The sequencing was 161 carried out using an automated PCR cycle-Genome Sequencer™ MiSeq (Illumina). Analysis and 162 alignment was performed using Vecton NTI suit 9 (InforMax, Inc.). Overall bioinformatics analysis was 163 done using NCBI-BLAST-2.2.24 and CLC bio Genomics workbench v7.5.1, for every sample set: 164 every read was BLASTed and the result file saved. The top 5 hits for every BLAST result (that is, 165 species name) was counted and a record was kept of how many times each species appeared as a 166 hit. The number in the last column is the number of times a read hit/matched that species. The 167 frequency (i.e count/total number of reads) and absolute count of each species were reported and 168 used to name the specific organism.

Sequencing Codons were finally saved in fasta format and output of results showing the kingdom, phylum, class, order, family, genus, species present in the sample given. The percentage of each variable was also enumerated. The names of species of culturable and non culturable organisms present in the sample, the corresponding accession number in NCBI data base, the number of hits and the e score were also given.

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175 3.0 RESULT AND DISCUSSION

176 **3.1 Colonial Morphology of Isolates**

177 All the isolates were circular in shape. The margin of the isolates were irregular, regular and entire.

178 While the elevation was either flat or convex (Table 1).

S/N	lsolate code	Colour	Shape	Size (mm)	Elevation	Margin
1	А	Brown	Circular	3.0	Flat	Irregular
2	В	Cream	Circular	3.5	Flat	Entire
3	С	Cream	Circular	3.2	Flat	Entire
4	D	Yellow	Circular	1.0	Flat	Entire
5	Е	Cream	Circular	1.4	Convex	Entire
6	F	Brown	Circular	5.0	Flat	Regular
7	G	Yellow	Circular	1.0	Flat	Entire
8	Н	Cream	Circular	1.5	Convex	Entire
9	I	Yellow	Circular	1.0	Flat	Entire
10	J	Cream	Circular	3.5	Flat	Entire
11	К	Cream	Circular	1.2	Convex	Entire
12	L	Yellow	Circular	1.0	Flat	Entire
13	Μ	Brown	Circular	3.0	Flat	Irregular
14	Ν	Cream	Circular	1.2	Convex	Entire
15	0	Cream	Circular	4.0	Flat	Entire
16	Р	Brown	Circular	3.0	Flat	Irregular
17	Q	Brown	Circular	4.5	Flat	Regular
18	R	Cream	Circular	3.5	Flat	Entire
19	S	Cream	Circular	3.2	Flat	Entire
20	Т	Yellow	Circular	1.0	Flat	Entire
21	U	Cream	Circular	3.5	Flat	Entire
22	V	Yellow	Circular	1.0	Flat	Entire
23	W	Cream	Circular	1.2	Convex	Entire
24	Y	Cream	Circular	3.0	Flat	Entire

179 Table 1: Colonial Morphology of bacterial isolates obtained from crude oil contaminated soil.

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3.2 Culture based identification of Isolates

Five different bacterial genera (*Acetobacter* sp., *Pseudomonas* sp., *Arthrobacter* sp., *Bacillus* sp. and *Micrococcus* sp.) were isolated and identified (Table 2). Most of the organisms isolated were gramnegative rods except *Bacillus* sp. and *Micrococcus* sp. that were gram-positive rod and cocci, respectively.

- 198 All the organisms isolated were indole negative and also showed an inability to produce hydrogen
- 199 sulphide (H₂S) and gas as well as an inability to ferment lactose. Bacillus sp. was the only spore
- 200 former isolated. All the bacterial isolates were urease negative except Acetobacter sp which was
- 201 urease positive (Table 2).

Table 2: Morphological and biochemical characteristics of bacteria isolated from crude oil

203 contaminated soil

S/N	Isolate Code	Gram reaction	Cell morphology	Oxidase	Citrate utilization	Methyl Red	Voges Proskauer	Catalase	Indole	Urease	Motility	Endospore	H ₂ S production	Spore test	Gas production	Glucose	Lactose	Maltose	Sucrose	Mannitol	Probable organism
1	А	-	Rod	-	-	-	-	+	-	-	+	-	-	-	-	A	-	-	+	-	Arthrobacter sp.
2	В	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
3	С	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
4	D	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
5	Е	+	Cocci	+	+	+	-	+	-	-	-	-	-	-	-	А	-	-	+	-	Micrococcus sp.
6	F	-	Rod	-	+	-	-	-	-	+	+	-	-	-	-	AG	-	+	-	-	Acetobacter sp.
7	G	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
8	Н	+	Cocci	+	+	+	-	+	-	-	-	-	-	-	-	А	-	-	+	-	Micrococcus sp.
9	T	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
10	J	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
11	Κ	+	Cocci	+	+	+	-	+	-	-	-	-	-	-	-	А	-	-	+	-	<i>Micrococcus</i> sp.
12	L	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
13	Μ	-	Rod	-	-	-	-	+	-	-	+	-	-	-	-	А	-	-	+	-	Arthrobacter sp.
14	Ν	+	Cocci	+	+	+	-	+	-	-	-	-	-	-	-	А	-	-	+	-	<i>Micrococcus</i> sp.
15	0	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
16	Ρ	-	Rod	-	-	-	-	+	-	-	+	-	-	-	-	А	-	-	+	-	Arthrobacter sp.
17	Q	-	Rod	-	+	-	-	-	-	+	+	-	-	-	-	AG	-	+	-	-	Acetobacter sp.
18	R	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
19	S	-	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
20	Т	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
21	U	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp.
22	V	-	Rod	+	-	+	+	-	-	-	+	-	-	-	-	AG	-	-	-	-	Pseudomonas sp.
23	W	+	Cocci	+	+	+	-	+	-	-	-	-	-	-	-	А	-	-	+	-	Micrococcus sp.
2 4	Υ	+	Rod	-	-	-	+	+	-	-	+	+	-	+	-	AG	-	-	-	-	Bacillus sp

205 These organisms have been reported by several researchers to be involved in the bioremediation of 206 petroleum hydrocarbons [14; 15]. As previously reported by [16], in a bioremediation study of a crude 207 oil polluted mangrove soil in Port Harcourt, using NPK as source of limiting nutrients, Acetobacter sp., 208 Pseudomonas sp., Arthrobacter sp., Bacillus sp., Alcaligenes sp., Flavobacterium sp., Citrobacter sp., 209 Vibrio sp. and Corynebacterium sp. are members of hydrocarbon utilizing bacterial population. The 210 findings of [16] were in consonance with the hydrocarbon utilizing bacterial population isolated in this 211 study, from Etelebou- Ogboloma Community (Gbarain Kingdom), an oil rich area in the Niger Delta 212 region of Nigeria.

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214 3.2 Molecular identification of autochthonous bacterial group

Metagenomics approach was used for *in situ* identification of bacterial population in the hydrocarbon impacted soil sample. The 16s bacterial metagenomic report revealed the taxonomic classification as follows:

Kingdom Classification: The report revealed bacteria as the predominant group (99.87%) and
protozoa was the next group with 0.09%, followed by fungi and archaea group with 0.02%
each.

Phylum Classification: The result showed that the soil had Proteobacteria (63.5%) as the predominant group. This was followed by Actinobacteria (9.67%), Firmicutes (1.29%), Chlamydiae (0.07%) and Ciliophora (0.05%). However, 25.28% of the taxa that belonged to this phylum where unknown whereas others (Bacteroidetes, Chloroflexi, Tracheophyta, Fusobacteria, Cyanobacteria) made up 0.11% of this phylum classification. Also, groups not assigned had 0.04% (Fig. 1).

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Class Classification: The class taxonomy shows the prevalence of the different groups in the
following order: Alphaproteobacteria (54.64%) > Unknown (25.28%) > Actinobacteria (9.67%)
Gammaproteobacteria (6.55%) > Betaproteobacteria (2.27%) > Bacilli (0.95%) > Clostridia
(0.34%) while the rest of the groups (others) made up 0.31% of the population (Fig 2).

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234Order Classification: Rhizobiales (54.29%) were found to be the more dominant group. While235Actinomycetales, Pseudomonadales, Bacillales, Xanthomonadales, Hydrogenophilales as236well as other members of the group, were 9.61%, 1.33%, 0.88%, 0.85%, 0.62% and 1.8% of237the population, respectively, the unknown group had 30.63% (Fig. 3).

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239 Family Classification: The metagenomic analysis of the polluted soil showed that the family 240 level taxonomical groups were mostly of the Methylobacteriaceae family (45.93%). Those that 241 belonged to the Mycobacteriaceae family were8.92% in population whereas, the 242 Bradyrhizobiaceae family made uр 8.06% of the population. The family of 243 Pseudomonadaceae accounted for 1.33% of this taxonomical classification. 244 Hydrogenophilaceae and Xanthomonadaceae constituted0.88% and 0.62% of the population,

respectively while the unknown group was 30.67% and others made up 3.58% of the population (Fig. 4).

BLAST output result: Data from the basic local alignment search tool (BLAST) showed (in Table 3) the most dominant bacteria to be *Methylobacterium* sp. which constituted 45.90% of the population. The uncultured bacterium was 24.63% and *Mycobacterium* sp., 8.86%. Also, *Bradyrhizobium* sp. was 7.99% whereas, uncultured gamma, *Pseudomonas aeruginosa* and betaproteobacterium accounted for 3.96, 1.29 and 1.05% of the bacterial population respectively.

255 Phylogenetic relationships: A phylogenetic tree based on the sequence analysis obtained
256 from the soil metagenomics is shown in Figs. 5a and b.

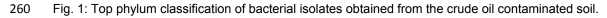
Ciliophora (0.05%) Chlamydiae(0.07%) Mot Assigned (0.04%) Others (0.11%) Actinobacteria (9.67%) Unknown (25.28%) Unknown (25.28%) Proteobacteria(63.5%)

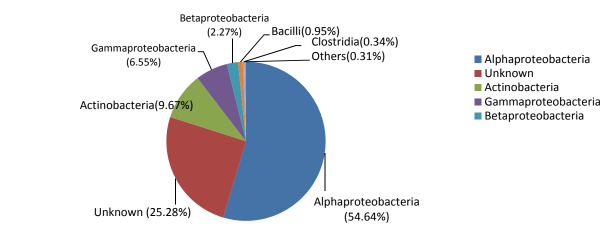
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263 Fig. 2: Top class classification of bacterial isolates obtained from the crude oil contaminated soil



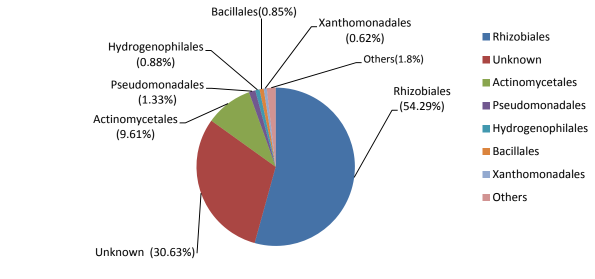


Fig. 3: Top order classification of bacterial isolates obtained from the crude oil contaminated soil.

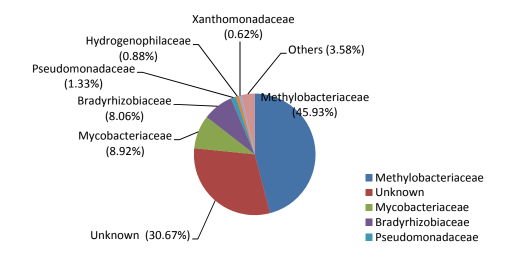


Fig. 4: Top family classification of bacterial isolates obtained from the crude oil contaminated soil.

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Table 3: BLAST output results

BLAST HIT	Read	Count %
1. Methylobacterium sp.	5857	45.90
2. uncultured bacterium	3143	24.63
3. Mycobacterium sp.	1130	8.86
4. Bradyrhizobium sp.	1020	7.99
5. uncultured gamma	505	3.96
6. Pseudomonas aeruginosa	165	1.29
7. beta proteobacterium	134	1.05
8. Thiobacillus prosperus	106	0.83
9. Stenotrophomonas maltophilia	79	0.62
10. Enterobacter sp.	69	0.54
11. Brachybacterium paraconglome	ratum 54	0.42
12. bacterium btn	46	0.36
13. Alicyclobacillus hesperidum	35	0.27
14. Ochrobactrum anthropi	32	0.25
15. unidentified eubacterium	26	0.20
16. Bacillus sp.	24	0.19
17. uncultured alpha	23	0.18
18. Staphylococcus aureus	22	0.17
19. Achromobacter sp.	11	0.09
20. uncultured beta	11	0.09
21. No hits	10	0.08
22. uncultured chlamydia	9	0.07
23. Geobacillus thermoleovorans	9	0.07
24. Anaerobic bacterium	9	0.07
25. Spirulina subsalsa	9	0.07
26. Bradyrhizobium japonicum	8	0.06
27. <i>Dietzia</i> sp.	8	0.06
28. Mycobacterium heidelbergense	8	0.06
29. uncultured streptococcus	7	0.05
30. Alicyclobacillus acidiphilus	7	0.05
31. uncultured eubacterium	7	0.05

306	32. <i>Clostridium</i> sp.	7	0.05
307	33. alpha proteobacterium	7	0.05
308	34. Providencia vermicola	6	0.05
309	35. <i>Thiobacillus</i> sp.	6	0.05
310	36. Chroococcidiopsis sp.	6	0.05
311	37. uncultured rothia	6	0.05
312	38. Acidovorax delafieldii	6	0.05
	0		
313	39. Bacterium 'smarlab	5	0.04
314	40. Ralstonia pickettii	5	0.04
315	41. Grimontella senegalensis	5	0.04
316	42. Staphylococcus epidermidis	5	0.04
317	43. Leuconostoc mesenteroides	5	0.04
318	44. Sphaerobacter thermophilus	4	0.03
319	45. Streptomyces sp.	4	0.03
320	46. Actinomycete species	4	0.03
321	47. Comamonas testosteroni	4	0.03
322		4	0.03
	48. <i>Paracoccus</i> sp.		
323	49. Sphingomonas sp.	3	0.02
324	50. Micrococcus lylae	3	0.02
325	51. Microcoleus vaginatus	3	0.02
326	52. Bacillaceae bacterium	3	0.02
327	53. uncultured brevundimonas	3	0.02
328	54. Aeromonas sp.	3	0.02
329	55. Pseudonocardia yunnanensis	3	0.02
330	56. uncultured hyphomicrobium	3	0.02
331		3	0.02
	57. Sulfuricurvum kujiense		
332	58. Bacillus soli	3	0.02
333	59. Micrococcus luteus	3	0.02
334	60. uncultured chloroflexi	2	0.02
335	61. Acidovorax sp.	2	0.02
	<u>^</u>	2	
336	62. uncultured soil		0.02
337	63. Oscillatoria amphigranulata	2	0.02
338	64. Bacteroidetes bacterium	2	0.02
339	65. uncultured fusobacterium	2	0.02
340	66. <i>Kocuria</i> sp.	$\overline{2}$	0.02
		2	
341	67. Alcaligenes sp.		0.02
342	68. Clostridium indolis	2	0.02
343	69. Acidocella sp.	2	0.02
344	70. uncultured pseudomonas	2	0.02
345	71. uncultured actinobacterium	1	0.01
346	72. Shewanella putrefaciens	1	0.01
347	73. Bacillus sphaericus	1	0.01
348	74. <i>Morganella</i> sp.	1	0.01
349	75. uncultured scenedesmus	1	0.01
350	76. agricultural soil	1	0.01
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351	77. uncultured chloroflexus	1	0.01
352	78. Pseudomonas stutzeri	1	0.01
353	79. Rhodopila globiformis	1	0.01
354	80. unidentified bacterium	1	0.01
355	81. uncultured archaeon	1	0.01
356	82. Bifidobacterium sp.	1	0.01
357	83. uncultured rubrobacteridae	1	0.01
358	84. Dysgonomonas sp.	1	0.01
359	85. Chondromyces crocatus	1	0.01
360		1	0.01
	86. Pseudomonas pertucinogena		
361	87. Agrobacterium tumefaciens	1	0.01
362	88. Saprospira sp.	1	0.01
363	89. <i>Delftia</i> sp.	1	0.01
364	90. uncultured candidate	1	0.01
365	91. <i>Rhizobium</i> sp.	1	0.01
366	92. uncultured gloeothece	1	0.01

367	93. uncultured delta	1	0.01
368	94. Marinospirillum alkaliphilum	1	0.01
369	95. uncultured rhodospirillaceae	1	0.01
370	96. microbulbifer sp.	1	0.01
371	97. uncultured sphingomonas	1	0.01
372	98. Clostridium subterminale	1	0.01
373	99. Cetobacterium somerae	1	0.01
374	100.Mobiluncus curtisii	1	0.01
375	101.uncultured methanogenic	1	0.01
376	102.uncultured syntrophorhabdaceae	1	0.01
377	103.Aquaspirillum peregrinum	1	0.01
378	104.Dellovibrio bacteriovorus	1	0.01

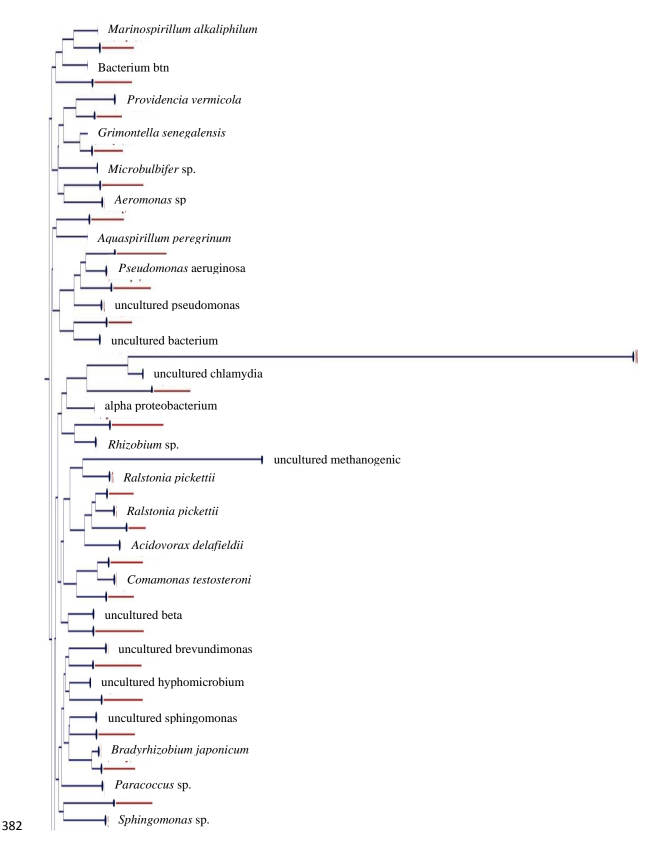


Fig. 5: Phylogenetic tree of 16S rRNA gene sequences of bacterial isolates obtained from crude oil contaminated soil sample.



Fig. 5 cont'd: Continuation of phylogenetic tree of 16S rRNA gene sequences of bacterial isolatesobtained from crude oil contaminated soil.

388 Metagenomic approach was used to identify the autochthonous bacterial population in the polluted 389 soil sample used in the investigation and results show that bacteria were predominant in the crude oil 390 impacted soil. Proteobacteria were the predominant bacterial population which may be indicative of 391 the major role played by the proteobacteria group in a crude oil polluted soil ecosystem. 392 Proteobacteria are Gram negative group of bacteria. This probably accounts for the predominance of 393 gram negative rods identified via culture-dependent approach described above. The data also 394 showed that the soil was mainly dominated by the Alphaproteobacteria (54.64%), followed by the 395 Actinobacteria (9.67%), Gammaproteobacteria (6.55%), Betaproteobacteria (2.27%) and Bacilli 396 (0.95%) as well as Clostridia (0.34%); as the most dominant classes of bacteria. 397

The order taxonomical classification showed that Rhizobiales (54.29%) were the most dominant group followed by the Actinomycetales, Pseudomonadales, Bacillales, Xanthomonadales and Hydrogenophilales.

402 The BLAST data indicated the dominant bacteria to be Methylobacterium sp. making up 45.90 % of 403 the population. The uncultured bacterium was 24.63% and Mycobacterium sp., 8.86%. Also, 404 Bradyrhizobium sp. was 7.99% and uncultured gamma, Pseudomonas aeruginosa and beta 405 proteobacterium accounted for 3.96, 1.29 and 1.05% of the bacterial population respectively. These 406 organisms have been identified by other researchers to be associated with biodegradation [17; 18]. 407 Molecular studies on microbial diversity have interestingly been related to degradation potentials of 408 these organisms by various researchers [17; 18; 19]. The research of [17] examined the degradation 409 of benzo[a]pyrene by a group of bacteria by using the DGGE technique, in order to determine their 410 dynamics during the degradation process, and identified Sphingomonas paucimobilis EPA505, 411 Mycobacterium str.PYR-1, and Alcaligenes denitrificans WW1 to be present in the consortium. These 412 organisms are well known to be associated with the degradation of aromatic compounds, as 413 Mycobacterium and Pseudomonas species have been also reported to be able to degrade PAH [18]. 414 In crude oil, carbazole is often present with its alkylated derivatives which often have 415 monomethyl, dimethyl, trimethyl, and tetramethyl side chains on different carbon positions of 416 carbazole (called C1-, C2-, and C3-, C4-carbazoles, respectively). Researchers have isolated strains 417 of Pseudomonas species, which could not only degrade carbazole efficiently, but also showed 418 nitrification and denitrification ability [19].

419

401

420 In the overall, one hundred and four (104) diverse bacterial species were identified as shown in table 421 3. The five (5) different bacterial genera (Acetobacter sp., Pseudomonas sp., Arthrobacter sp., 422 Bacillus sp. and Micrococcus sp.) isolated via cultural approach were similar to the bacterial species 423 identified using metagenomics. However, a comparison of the number of isolates obtained using both 424 methods clearly indicates that molecular techniques such as metagenomics increases the 425 understanding of bacterial diversity as well as functionality during bioremediation. Cultural techniques 426 only help to culture and identify minute population of microorganisms [20], representing about 1% of 427 the total microorganism found in a crude oil contaminated site [10].

430 The phylogenetic tree showed the predominant bacterial community in the hydrocarbon polluted soil 431 in relation to each other, using a scale of 0.2 and uncultured delta (0.01) at the root of the tree. The 432 phylogenetic sequence relationship shows that the most distant groups/species are the uncultured 433 scenedesmus (0.01%), uncultured alpha (0.18%) and Rhodpila globiformis (0.01%). These groups 434 however, belong to the same clade with Mobiluncus curtisii (0.01%). The no hits (0.08%) were also 435 seen to be distantly related to the uncultured Clamydia (0.07%) which is related to the alpha 436 proteobacteria (0.05%). The tree shows numerous clades for the 104 BLAST output result. This 437 shows the crude oil contaminated soil is composed of diverse groups of microorganisms and thus, 438 implies the hydrocarbon polluted soil is a reservoir for diverse bacterial groups.

439

450

440 Similar study by [21] involving the excision, amplification and sequencing of dominant DGGE bands in 441 biostimulated soils revealed the presence of distinct hydrocarbon degraders like Corynebacterium 442 spp., Dietzia spp., low G+C Gram positive bacteria and some uncultured bacterial clones. 443 Phylogenetic analysis of the 16S rRNA gene sequences of these dominant bacterial communities was 444 conducted using the neighbour joining method of PHYLIP. The researcher observed two distinct 445 clades appearing in the tree with members of the Actinobacteria and Firmicutes clustering separately 446 [21]. In another study, the analysis of 16S rRNA of the isolated isolates from a crude oil impacted soil 447 in the Niger Delta region of Nigeria showed the species belonged to eight bacterial genera namely: 448 Achromobacter, Alcaligenes, Azospirillus, Bacillus, Lysinibacillus, Ochrobactrum, Proteus, and 449 Pusillimonas, with Alcaligenes as the dominant genus [22].

451 The variations in the number of hits of bacterial species observed in this study conform to the fact that 452 the polluted soil has diverse bacterial groups with different levels of adaptation. The ability of these 453 bacteria to metabolize or adapt to crude oil hydrocarbons is relative to the possession of degradative 454 genes or resistance genes [23]. These studies have revealed the phylogeny of the polluted soil used 455 in this study, which shows the diverse species of bacteria. These bacterial species differ in their 456 structure and function. For example *Thiobacillus* sp. is known to be associated with iron oxidation and 457 inorganic sulphur reduction (Thiobacillus ferroxidans). Some others are associated with heavy metal 458 reduction. This implies the polluted soil ecosystem harbours diverse bacterial population having 459 different structures and function which aid the process of bioremediation. While some of these 460 organisms are hydrocarbonoclastic, others (like Pseudomonas sp.) are known for biosurfactant 461 production. Biodegradation of petroleum hydrocarbon pollutants is therefore feasible in such 462 ecosystem. However, the fate of these crude oil pollutants will partly depend on the ecological/site 463 characteristics of the impacted soil. Gbarain kingdom is an oil rich region in the Niger Delta area of 464 Nigeria. This area is replete with crude oil pipe lines traversing the land and water body. This may be 465 partly responsible for Thiobacillus sp. ranking among the top 8 in the BLAST output report. This 466 finding does not only make a case for bioremediation but also presents the critical need for pipeline 467 coating and biomonitoring.

468

469 4.0 CONCLUSION

470 The bacterial community composition of a crude oil polluted soil obtained from Gbarain Kingdom of 471 Bayelsa State, Nigeria suggests the presence of a rich and versatile group of hydrocarbon utilizing 472 bacteria adapted for crude oil degradation. These hydrocarbonoclastic bacterial species are involved 473 in the utilization of the petroleum hydrocarbons spilled at the site sampled and they have the 474 potentials to facilitate the bioremediation of the contaminated soil through natural attenuation or 475 enhanced bio-treatment protocols. Data obtained in this study make it obvious that only a very small 476 proportion of the bacterial population in the soil can be isolated and identified through cultural 477 techniques. Therefore, a detailed insight into the agents and dynamics of crude oil degradation in soil 478 thus, requires in-depth metagenomics which provide the structure and function of the diverse 479 microbial genera for the purpose of exploring and harnessing the potentials of these autochthonous 480 bacterial populations. The emergence of next generation gene sequencing (NGS) technique has 481 enabled the identification of a wide range of bacteria which were hitherto, unidentifiable via culture-482 based methods. The NGS method is performed independent of PCR products and boycotts the 483 limitations associated with such techniques and is suitable for the study of the bacterial community 484 structure of a crude oil impacted media. However, cultural methods are not to be sacrificed for 485 identification using molecular characterization as both methods are important in bioremediation 486 monitoring.

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