

1 **Comparative study on physicochemical parameters study of oil polluted sites and**
2 **hydrocarbon degradation potentials of heterotrophic bacteria in southern Nigeria**
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5 **ABSTRACT**

6 In this study, Comparative study on physicochemical parameters study of oil polluted sites
7 and hydrocarbon degradation potentials of heterotrophic bacteria in southern Nigeria

8 hydrocarbon degradation potentials of heterotrophic bacteria isolated from oil-polluted soil
9 were examined; Samples were collected from Sakpenwa, an oil producing community in Tai
10 LGA of Rivers State, The gravimetric analysis showed the bacteria were capable of utilizing
11 96.9-99.7% the oil samples, The amount of hydrocarbon in the soil samples were determined
12 using Gas Chromatography-Flame Ionization Detector, GC- FID. Statistical analysis was
13 carried out using Statistical Package for Social Sciences (SPSS, version 20.0). Analysis of
14 variance (ANOVA) carried out at 95% level of confidence software that the degree of
15 hydrocarbon degradation varied amongst isolates, *Pseudomonas aeruginosa* and *Alcaligenes*
16 sp. showed highest degrading activities while *Bacillus subtilis* sp. showed least activity. This
17 study revealed that indigenous bacterial species possess the requisite gene necessary for
18 hydrocarbon biodegradation. This concluded that degradation is most often the primary
19 mechanism for contaminant destruction including petroleum contaminants.
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21 **KEYWORDS:** physicochemical parameters, hydrocarbon degradation Gas Chromatography,
22 Analysis of variance,
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One major substance causing environmental pollution in the South-South of Nigeria is Crude oil. Annually, a large amount of petroleum hydrocarbon is being released into the marine and estuarine environments, causing great harm to many organisms, while some (microbial degraders) use them as energy source (McKew *et al.*, 2007). A wide range of pollutants (e.g. PAHs) can be degraded by microorganisms found within the polluted environment (Barathi and Vanudevan, 2001). However, environmental factors like soil pH, moisture, temperature and nutrients affect the rate of pollutants' degradation by microorganisms in the soil. Other factors that influence pollutants' degradation are toxicity of pollutant (Kenawy *et al.*, 2007), Bacteria are primary producer in the marine food web, where their role is to recycle nutrients and breaking down of hydrocarbon (Bordenave *et al.*, 2004, Edmund and Jansson, 2006; Head *et al.*, 2006). Various bacterial species isolated from soil have been proved to degrade PAHs (Aislabie *et al.*, 2004). Other hydrocarbon degraders include *Alcaligenes* sp., *Acinetobacter* sp., *Flavobacter* sp., *Cyanobacterium* sp., *Moraxella* sp. and *Bacillus* sp (Bhattacharya *et al.*, 2002).

Hydrocarbon utilizing bacterial genera include: *Pseudomonas*, *Arthrobacter*, and *Micrococcus* (Atlas, 1981). Previous studies have found the evolution of some obligate hydrocarbon degraders (also named as obligate hydrocarbonoclastic bacteria) of indigenous marine bacterial genera. (Brooijmans *et al.*, 2009). In summary, only few number of this organism can degrade hydrocarbon *in situ* (Head *et al.*, 2006). *Alcanivorax* sp. is a good hydrocarbon degrader because it has been proved in many parts of the world including the United States to potential hydrocarbon degrader (Yakimov *et al.*, 2007). Microorganisms are active degraders of hydrocarbon in an environment if high numbers of oil degrading microbes are present in the environment (Okerentugba and Ezeronye, 2003). Bacteria are the prominent degraders of petroleum in oil contaminated environment (Rahman,*et al.*, 2003). A lot of bacteria utilize hydrocarbons as their energy source (Yakimov, *et al.*, 2007). Years ago, bacteria were used as the easiest agent for removing petroleum contaminants off the environment (Leahy and Colwell, 1990). Utilization of n-alkanes ranging from C10–C40 by *Acinetobacter* sp. have been discovered (Throne-Holst, *et al.*, 2007). Isolation of bacterial species from the genus, *Mycobacterium* capable of hydrocarbon degradation in the soil has been ascertained (Chaillan, *et al.*, 2004). Hydrocarbon biodegradation usually requires the consortium of species. One of the most persistent groups of organic pollutant in the ecosystem is petroleum hydrocarbons (Huang *et al.*, 2005). They can disrupt the food chain, leading to ecological cycle destruction (Khan, 2005).

MATERIALS AND METHODS

2.1 Study area and sample collection.

The study site was located at the oil polluted sites in Sakpenwa community in Ogoni land, Tai Local Government Area, Rivers State. Soil samples were collected 500m and 1000m away from the major spill sites. Fifty grams (50g) of the oil-polluted soil samples were collected from each of the sampling points using a soil sampler. The collected soil samples were transported in plastic nylon bags from the polluted sites to the Department of Microbiology, University of Port Harcourt laboratory for analysis within 24 hours (Iheanacho *et al.*, 2014).

2.2 Samples preparation

The soil samples collected were passed through a mesh sieve (2mm pore size) to remove large particles and were thoroughly mixed. Thereafter, 5g of each soil sample was suspended in 45 ml of distilled water. The suspended samples were mixed properly in a rotary shaker at 100 rpm at room temperature ($28 \pm 2^{\circ}\text{C}$) for 1hour, 30 minutes to liberate the organisms into the liquid medium (Iheanacho *et al.*, 2014). The pH of the samples was also taken.

89 **2.3 Isolation and enumeration of total heterotrophic bacteria**

90 The total culturable heterotrophic bacterial count for each degradation set-up was enumerated
91 using the streak plate method (Odokuma and Okpokwasili, 1992). Serial dilutions of the
92 samples were made and 0.1ml aliquot of the 10^{-1} to 10^{-4} dilutions of each sample were
93 transferred onto well dried, sterile nutrient agar plates (in triplicate) and incubated at 37°C for
94 24. After incubation, the bacterial colonies that grew on the plates were counted and sub-
95 cultured onto fresh nutrient agar plates using the streak-plate method in order to obtain pure
96 cultures of each colony. Discrete colonies on the plates were then transferred into nutrient
97 agar slants, properly labelled and stored at 4°C as a stock culture for preservation and
98 identification (Odokuma and Ibor, 2002).

99 **2.4 Biodegradation studies**

100 The method proposed by Ekpo and Ekpo (2006) was used. The biodegradation study of
101 hydrocarbons in the polluted soil was carried out using the Bushnell-Haas broth. This
102 medium consist $MgSO_4$ 0.02g; $CaCl$ 0.2g; K_2HPO_4 100g; $KHPO_4$ 1g; NH_4NO_3 1g; F_2Cl 0.05g
103 was autoclaved in 2 litres conical flasks. 99ml of the liquid medium (Bushnell-Haas broth)
104 was dispensed into five (5) conical flasks into which 1 ml of sterile crude oil was added (Ekpo
105 and Ekpo, 2006). Precisely, 5ml of each of the bacterial isolates (in liquid broth) were
106 inoculated into five (5) different conical flasks containing the liquid medium. The
107 concentration of day zero was use as control to the other subsequent days. The bacterial
108 cultures were incubated at ambient temperature (4°C) in an electric shaker of 100 strokes per
109 minute for 30 minutes each day. Sampling period was set for every 5 days for 30 days (Okoh,
110 2003). Bacterial utilization of hydrocarbon was monitored using their optical density at
111 600nm wavelength (Ekpo and Ekpo, 2006). The total petroleum hydrocarbon was measured

112 **2.5 Determination of Crude oil Degradation rate in Soil**

113 The crude oil degradation rate in the soil was determined by the solvent extraction method
114 (Chithra and Shenpagam, 2014). Five grams (5g) of soil sample was mixed with 100ml of
115 normal hexane in a flask and corked. The mixture was shaken using a mechanical shaker for
116 1hr, and then allowed to settle. With the use of a sterile syringe, an aliquot of the oil extract in
117 the solvent solution (20ml) was withdrawn and put in a previously weight evaporation dish.
118 The dish and its content were evaporated to dryness in a rotary evaporator and the dish was
119 reweighted to obtain the difference.

120 The percentage (%) of the degradation was shown as follows:

121 Weight of residue crude oil = weight of beaker containing extracted crude – weight of empty
122 beaker

123 Amount of crude oil degraded= weight of crude oil added to the media – weight of residual
124 crude oil.

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$$\% \text{ Degradation} = \frac{\text{amount of crude oil degraded}}{\text{amount of crude oil added to the media}} \times 100$$

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127 **2.6 Sample preparation for Total Petroleum Hydrocarbon (TPH) analysis (AOAC**
128 **1990).**

129 Precisely, 2ml of sample was extracted with 20ml of dichloromethane. Separating funnel
130 used to separate the sample and the dichloromethane layer was concentrated in a rotator
131 evaporator. 1ml of acetonitrile was added into the concentration and transferred into vial
132 ready for analysis.

133 **Fixed setting:** Generally, adjusting of gas flows to the columns was done, the inlets, the
134 detectors and the split ratio. In addition, the injector and the detector temperature were also
135 set. The detector was held at a high end of the oven temperature range to minimize the risk of
136 analyte precipitation. All of these parameters were set to the correct values but double
137 checking of all the instrument was done: Buck 550 gas chromatograph equipped with an on-
138 column, automatic injector, flame ionization detectors, HP 88 capillary
139 column(100mm×0.25µm film thickness,) CA,USA

140 Detector temperature A: 250° C
141 Injector temperature 22° C
142 Integrator chart speed: 2cm/min
143 The oven was set at a temperature of 180° C which warms up the Gas Chromatogram, while
144 its warming set was:

146 Table 1: Temperature condition for warming the oven.

Initial temp.	Hold time	Ramp time	Final temp
70° C	5mins	10mins	220° C
22° C	2mins	5mins	280° C

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148 Light will turn off and you begin your run your experiment, when the instrument is ready,
149 then Inject 1 microliter sample onto column A, using proper injection technique.

150 **2.7 Determination of changes in Total Petroleum Hydrocarbon content of soil.**

151 The extract was analyzed using the Buck 550 gas chromatography equipped with a Field
152 Ionization Detector (FID), High Performance (HP) 88 capillary column (100mm × 0.25 μm)
153 with a nominal film thickness of 0.25μm, while the volume of the injection was 0.8μL at
154 22°C. The carrier gas was Helium at (2cm/min) because hydrocarbon is a volatile compound.
155 The holding capacity of the column is 35°C for 1.50 min. The temperature increases gradually
156 from 22°C min⁻¹, to 280°C min⁻¹ and held for 5mins. The total sum of the components
157 present is equal to the sum of Total Petroleum Hydrocarbon (TPH) present in the GC
158 capillary column between of 5 to 35 min retention time. (Saari *et al.*, 2007).

159 **2.8 Statistical analysis**

160 Statistical analysis was carried out using Statistical Package for Social Sciences (SPSS,
161 Version 20.0). Analysis of variance (ANOVA), P- values test of significance, was carried out
162 at 95% level of confidence, P - values was use to determine the significance levels between
163 various treatments and data obtained during the study.

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RESULTS AND DISCUSSION

The bacterial diversity present in the Control soil, Site A (500m) and Site B (1000m) of this study was as represented in Table 2.

Table 2: Culturable bacterial diversity presents in the various study sites

Control	Polluted site (500m) away	Polluted site (1000m) away
<i>Acinetobacter</i> sp.	<i>Pseudomonas</i> sp.	<i>Alcaligenes</i> sp.
<i>Alcaligenes</i> sp.	<i>Bacillus</i> sp.	<i>Citrobacter</i> sp.
<i>Pseudomonas</i> sp.	<i>Acinetobacter</i> sp.	<i>Bacillus</i> sp.
<i>Serratia</i> sp.		<i>Acinetobacter</i> sp.
<i>Bacillus</i> sp.		

Physicochemical Properties of the test soil

It is an established fact that pollutants (e.g. hydrocarbons) affect the physicochemical condition of perturbed soil ((Roling *et al.*, 2004; Margesin *et al.*, 2007). Table 3 showed the level of changes in the physicochemical parameters of soil caused by the pollutant and shows how the contamination level increase, the amount of nitrate and phosphorous in the soil reduce. The control soil has a high and normal quantity of nitrate and phosphorous of 58.30mg/kg, and 10.20mg/kg respectively, after 500m away, from the polluted site the quantity of the nitrate and phosphorous were 13.90mg/kg and 1.50mg/kg and to further confirm the soil was polluted, a sample was taken from 100meter away from the polluted site for further study and this shows a value of 42.70mg/kg and 3.2mg/kg of nitrate and phosphorous respectively. Essential nutrients in the soil were reduced as soil is contaminated, the decrease in nitrate and phosphate level is attributed to the fact that they were been used in the metabolism of organism in building biomass. There is a positive correlation in the utilization of both nitrate and phosphate and this indicate their importance in cell metabolism. It was establish that the availability of nitrogen and phosphorus limit the microbial degradation of hydrocarbon (Abu and Ogiji, 1996; Zhu *et al.*, 2001) and the pH of the soil varies from 7.24 from the control soil sample to 5.08 and 6.47, 500 and 1000meters away from the polluted site respectively, which shows that the soil sample was acidic. This concord with the work of (Amund and Adebisi, 1991; Okpokwasili and James, 1995) which proofs that microbial utilization of hydrocarbons often leads to production of organic acids. Thus, the acids probably produced account for the reduction in pH levels.

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Table 3: Physicochemical parameters of the soil samples

S/N	Parameters	Control Soil	Polluted Soil 500m Away	Polluted Soil 1000m Away
1	pH	7.24	5.08	6.47
2	Conductivity ($\mu\text{s}/\text{cm}$)	126.00	750.00	1097.00
3	Sulphate (So_4) (mg/kg)	10.00	95.00	90.00
4	Moisture content (%)	0.20	2.20	0.04
5	Ammonia, (No_3) (mg/kg)	0.00	0.08	0.13
6	AmmoniaNitrogen (NH_3N) (mg/kg)	0.00	0.07	0.11
7	Nitrate (NO_3) (mg/kg)	56.30	13.90	42.70
8	Nitrate-Nitrogen ($\text{NO}_3\text{-N}$) (mg/kg)	13.20	3.10	9.70
9	Calcium (Ca) (mg/kg)	0.20	0.24	0.00
10	Magnesium (Mg) (mg/kg)	0.14	0.28	0.33
11	Sodium (Na) (mg/kg)	5.00	34.00	22.00
12	Potassium (K) (mg/kg)	1.30	8.00	5.08
13	Nickel (Ni) (mg/kg)	0.00	0.00	0.00
14	Mercury (Hg) (mg/kg)	0.00	0.00	0.00
15	Lead (Pb) (mg/kg)	0.31	0.21	0.29
16	Copper (Cu) (mg/kg)	0.05	0.00	0.00
17	Iron (Fe) (mg/kg)	4.13	3.93	4.33
18	Zinc (Zn) (mg/kg)	5.24	0.49	0.01
19	CEC (mg/kg)	0.56	1.70	1.17
20	Total Organic Carbon (%)	31.60	33.00	34.70
21	Phosphate (P_2O_5) (mg/kg)	31.40	4.60	9.70
22	Phosphorous (P) (mg/kg)	10.20	1.50	3.20
23	Phosphate (PO_4^{3-}) (mg/kg)	23.40	3.50	7.20
24	Ash content (%)	85.00	85.00	90.82
25	TPH, /kg	16.00	12984.00	184.80
26	Total Organic Matter	63.20	66.00	69.40

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231 Figure 1 to Figure 4 below showed the Chromatographic profile of hydrocarbon degradation
232 of the various hydrocarbon utilizing bacteria at the initial day, day (0) and the final day for
233 degradation study, day 30 respectively. Thus showing the result of how the hydrocarbon was
234 reduced by each bacterium

235 The hydrocarbon-utilizing bacterial genera isolated from the oil contaminated soil were
236 *Pseudomonas*, *Bacillus*, *Acinetobacter*, *Alcaligenes* and *Citrobacter*. Okpokwasili and Okorie
237 (1990) isolated similar hydrocarbon utilizing bacteria from Niger Delta aquatic systems.
238 Chikere and Okpokwasili (2004) also made similar findings on petroleum effluents. It has

239 also been observed that some microorganisms are more abundant in areas of high
240 concentration of hydrocarbons. These micro floras are actively oxidizing the hydrocarbons
241 and this is considered as another source of carbon for use in the ecosystem.
242 Results from the Gas Chromatography analyses from figure 1 to 4 shows that during the first
243 (5) days, Total Petroleum Hydrocarbon (TPH) reduction was high due to evaporation which
244 shows that some components of the hydrocarbon were volatile. The reductions in both the
245 number and sizes of the peaks and height from the profiles corroborate with the report by
246 Okpokwasili and Okorie (1988) which states that they are hydrocarbon degraders which
247 utilize the breakdown products of hydrocarbon. *Aciligenes* growth increased simultaneously
248 with decrease in TPH throughout the 30days period monitored leading to nutrient limitation
249 in the soil.

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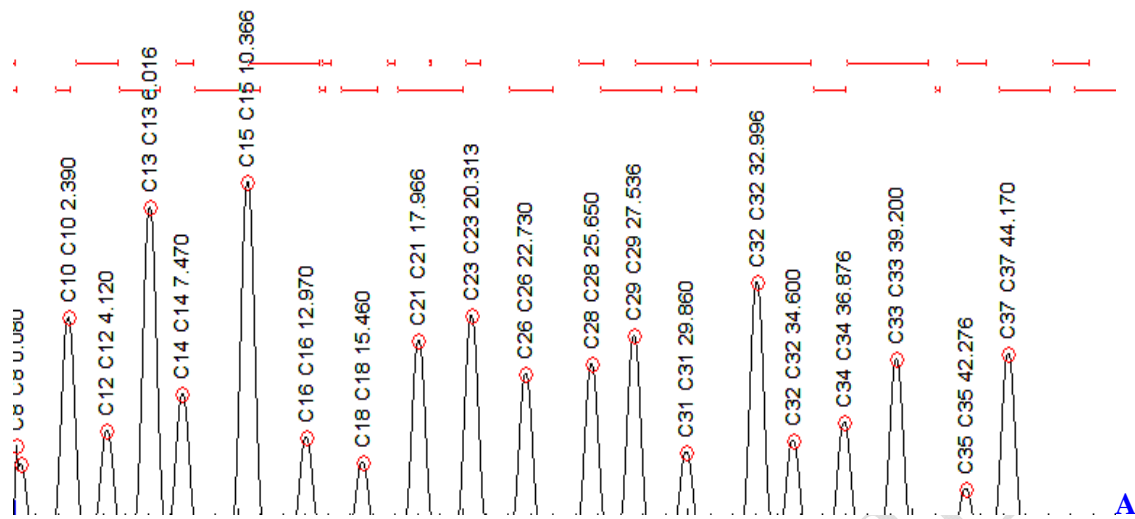
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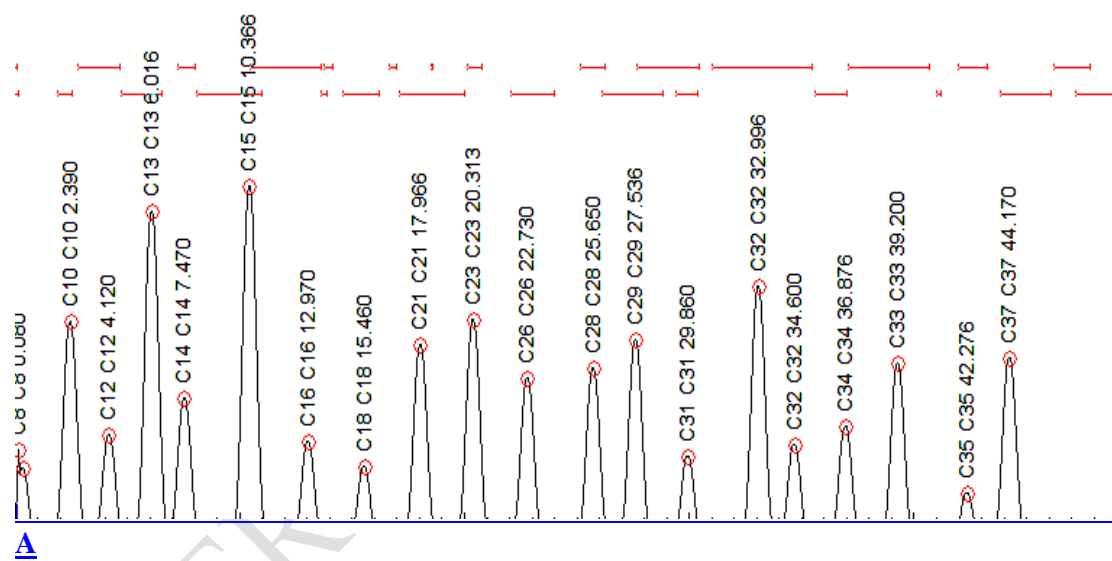
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UNDER PEER REVIEW

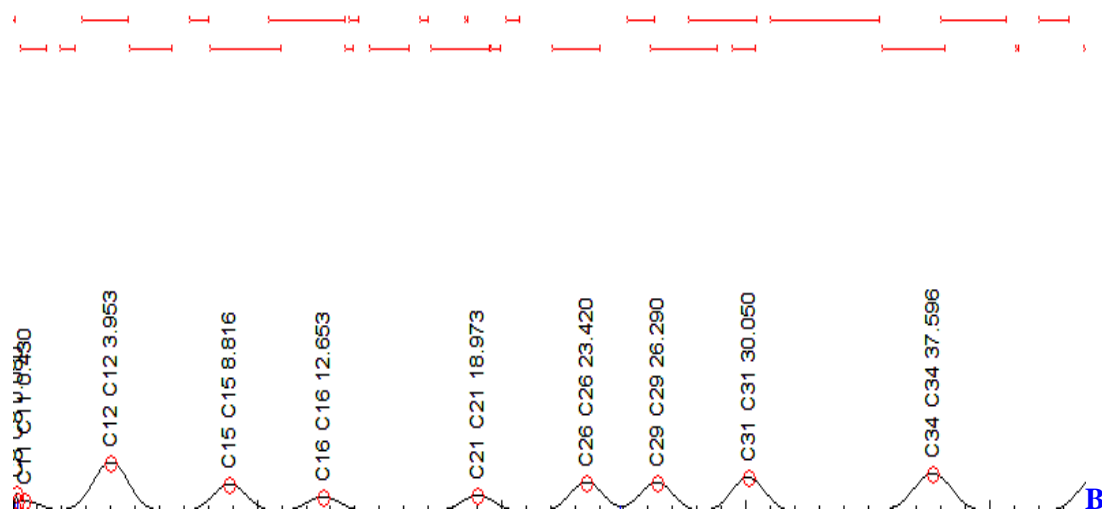
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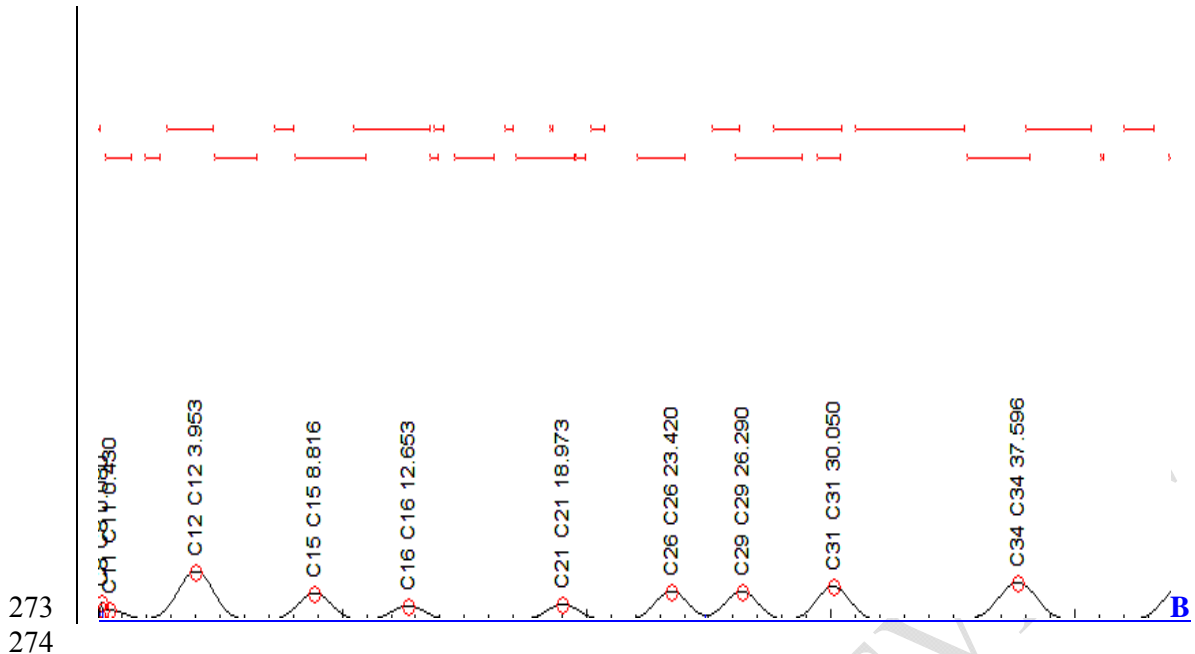


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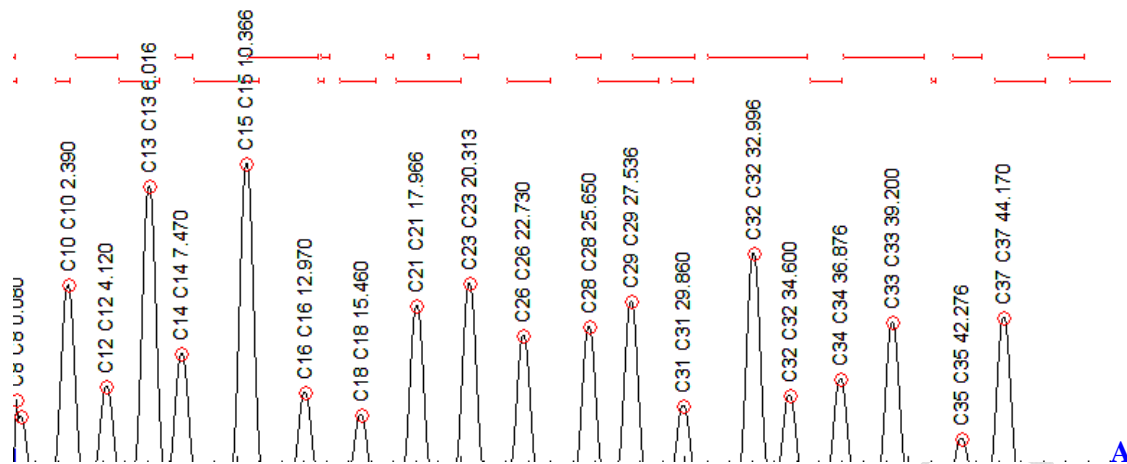


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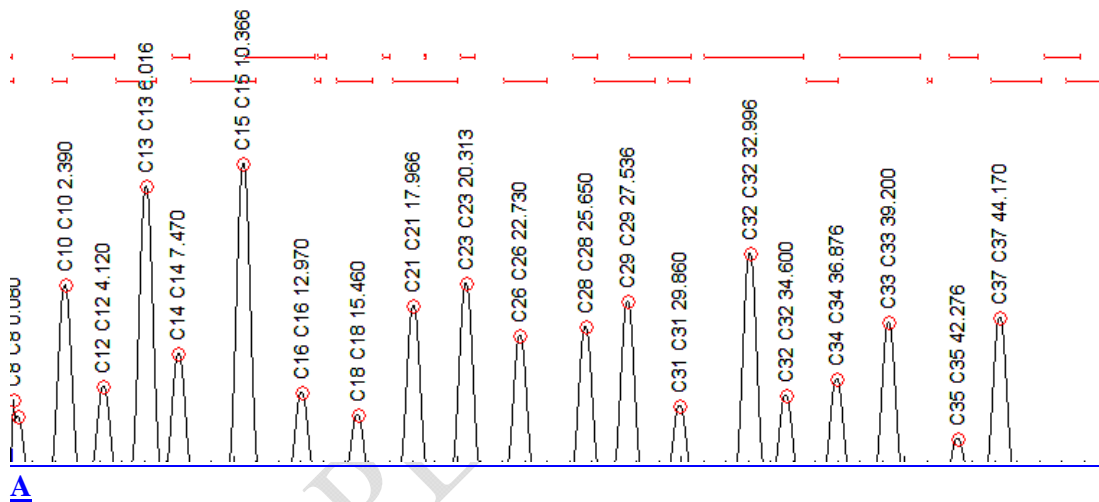
Figure 1: Profile of hydrocarbon degradation by *Alcaligenes* sp on day zero (A, top) and after 30 days (B, below).

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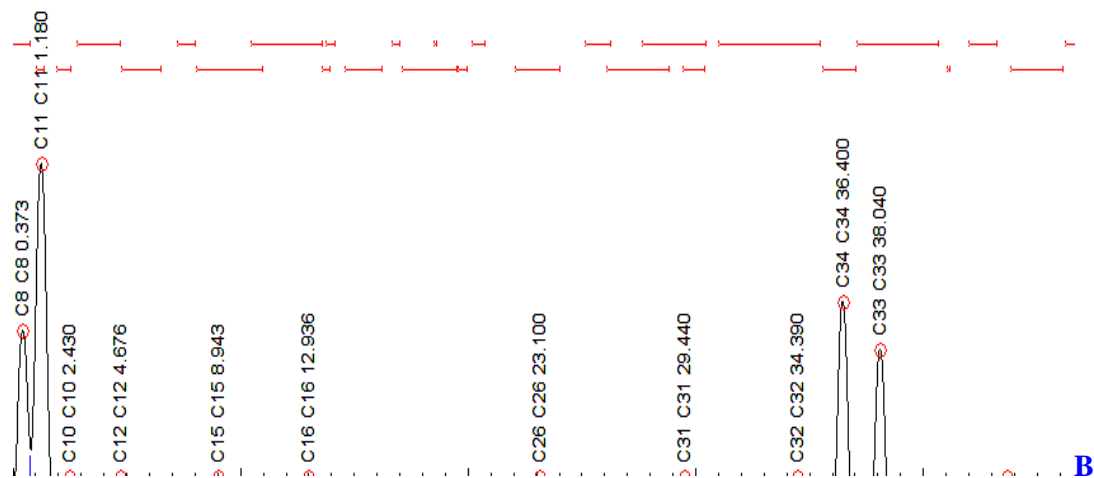
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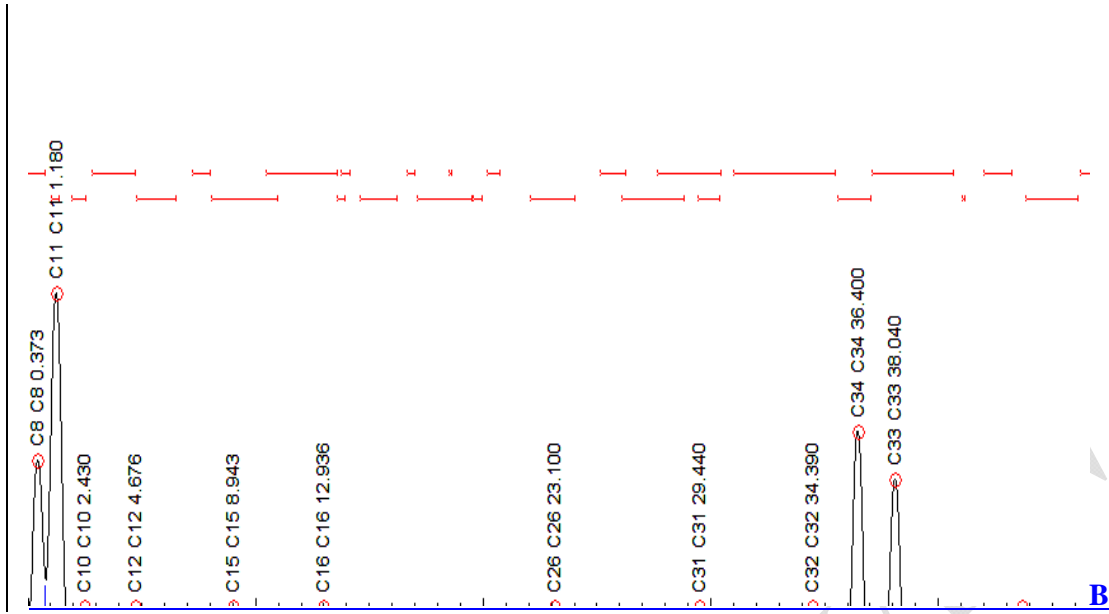
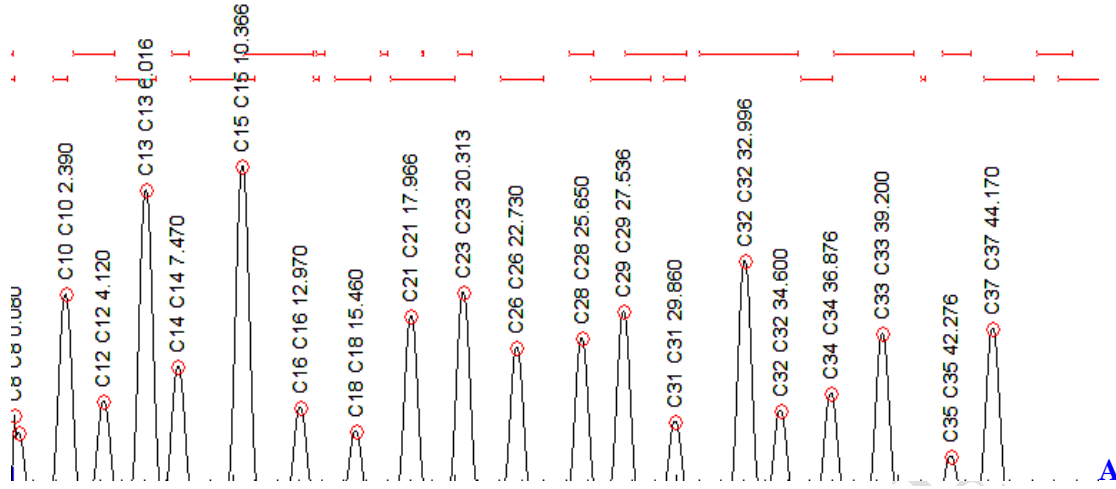


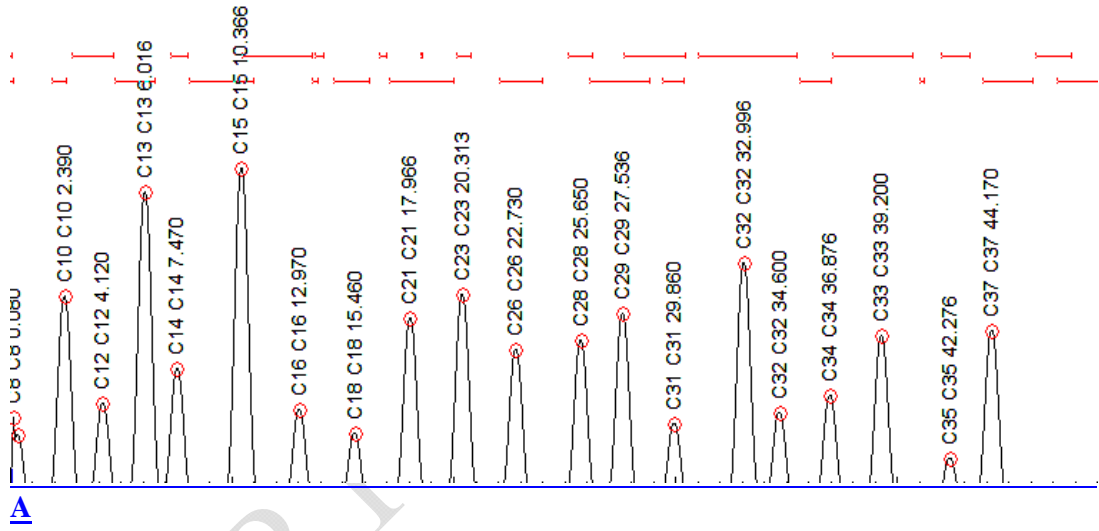
Figure. 2: Profile of hydrocarbon degradation by *Acinetobacter* sp on day zero (A, top) and after 30 days (B, below).

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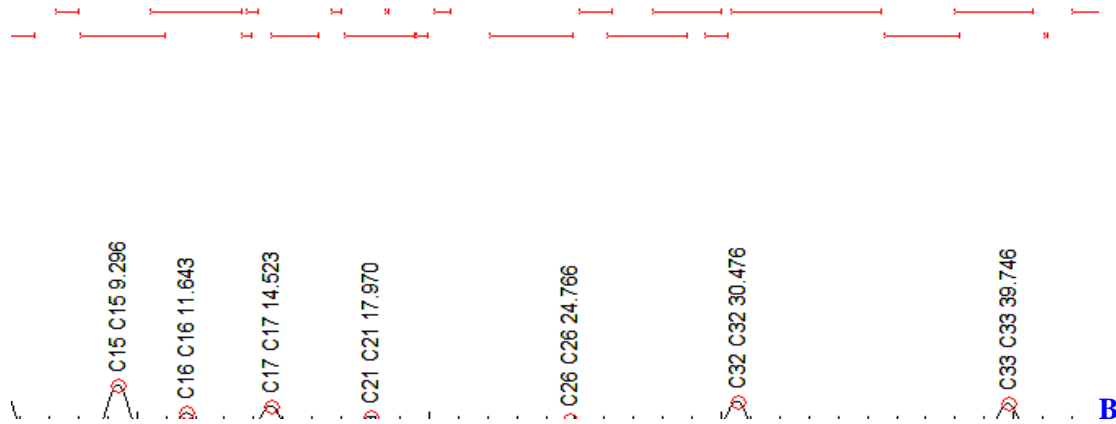
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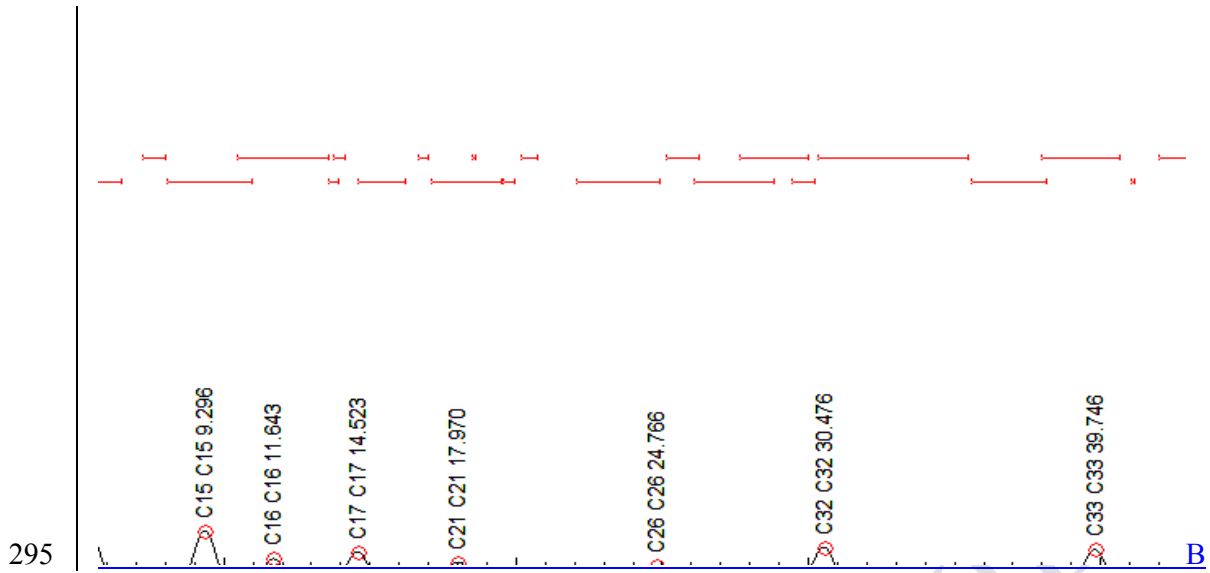


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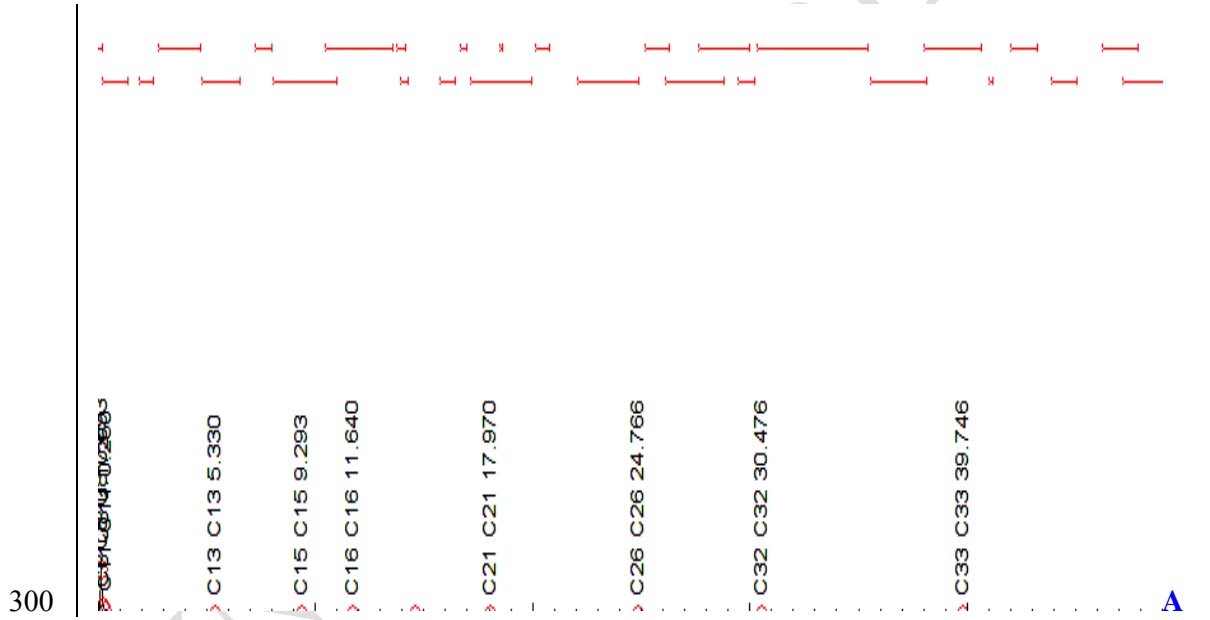
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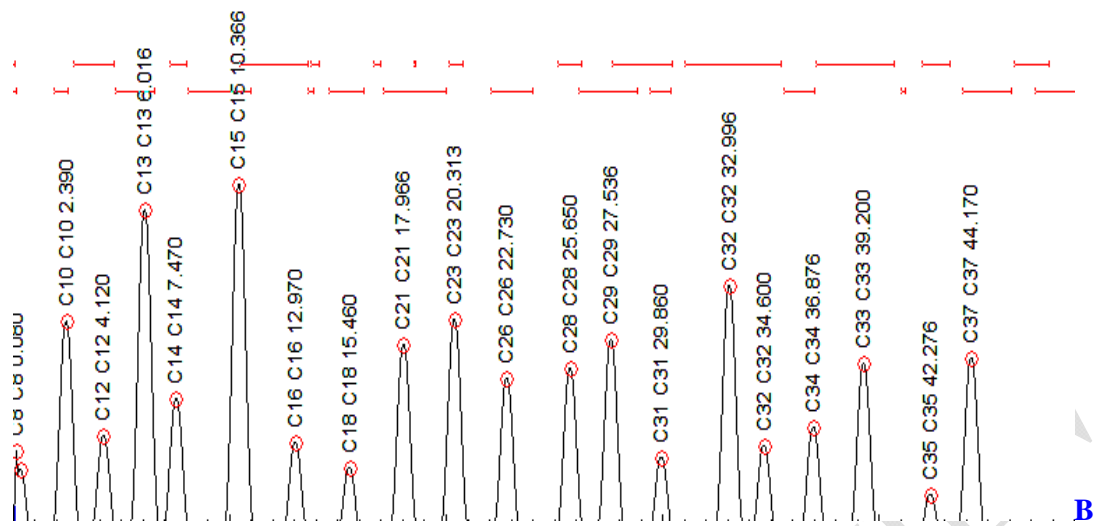
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Figure.3: Profile of hydrocarbon degradation by *Bacillus subtilis* on day zero (A, top) and after 30 days (B, below)

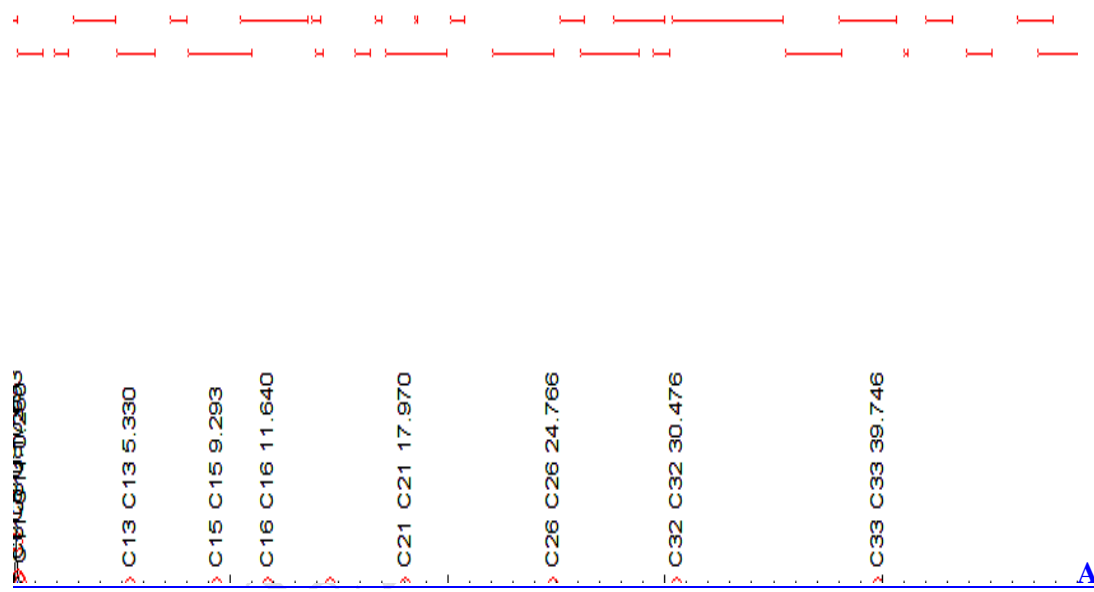


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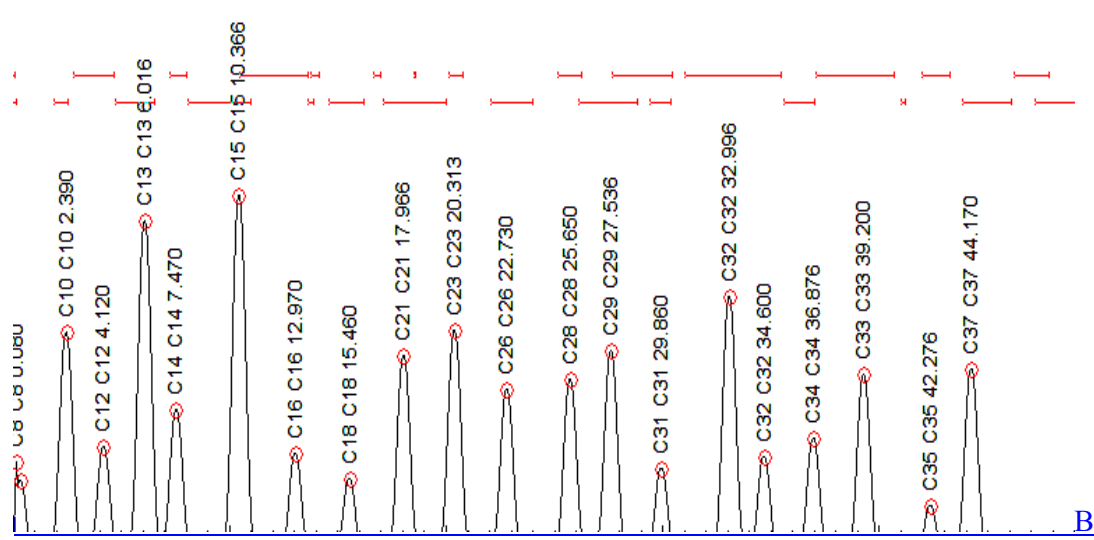
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Figure.4: Profile of hydrocarbon degradation by *Pseudomonas aeruginosa* on day zero (A, top) and after 30 days (B, below)

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The TPH present were C-8,C-10, C-11, C-12, C-13, C-14, C-15, C-16, C-18, C-21, C-23, C-26, C- 28, C-29, C-31, C-32, C-33, C-34, C-35 and C-37 for the treatment on day 0-30 days as showed in table 3.. The total amount of TPH after contamination of soil with crude oil showed that the treatment had TPH of (13729.70mg/ml) for the first day and reduced significantly after 30 days by each of the bacterium as follows: *Bacillus* sp (426.15mg/ml) which was able to degrade the Hydrocarbon by 96.90%, *Pseudomonas* sp. (58.68mg/ml), which was able to degrade the Hydrocarbon by 99.60% *Alcaligenes* sp.(111.07mg/ml) which was able to degrade the Hydrocarbon by 99.20% and *Acinetobacter* sp. (38.37mg/ml) which was able to degrade the Hydrocarbon by 99.70% This concord with the work of Margesin *et al.* (2003) and Quatrini *et al.* (2008) which demonstrated that *Actinobacter* sp. play important role during petroleum hydrocarbon degradation. C-8, C-11, C-15, and C-26, were recalcitrant to degradation while C-10, C-14, C-18, C-23, C-28, C-29, and C-35 were all degraded in the degradation study. Statistical analysis of the result shows that there is significant difference between various hydrocarbon and soil samples.

Table 3: The extent of degradation of the hydrocarbons by selected bacterial degraders after 30days

Hydrocarbons	<i>Bacillus</i> sp.	<i>Pseudomonas</i> sp.	<i>Alcaligenes</i> sp.	<i>Acinetobacter</i> sp.
C ₈	✓	✓	✓	✓
C ₁₀	-	-	-	-
C ₁₁	✓	✓	✓	✓
C ₁₂	-	-	✓	✓
C ₁₃	✓	✓	-	-
C ₁₄	-	-	-	-
C ₁₅	✓	✓	✓	✓
C ₁₆	-	✓	✓	✓
C ₁₈	-	-	-	-
C ₂₁	✓	✓	-	-
C ₂₃	-	-	-	-
C ₂₆	✓	✓	✓	✓
C ₂₈	-	-	-	-
C ₂₉	-	-	-	-
C ₃₁	-	-	✓	✓
C ₃₂	✓	✓	-	✓
C ₃₃	✓	✓	-	-

C ₃₄	-	-	✓	✓
C ₃₅	-	-	-	-
C ₃₇	-	-	✓	-
Hydrocarbon remaining	426.15mg/ml	58.68mg/ml	111.07mg/ml	38.37mg/ml
Hydrocarbon degraded	96.90%	99.60%	99.20%	99.70%

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Key: Present (remaining) = ✓ ; Complete removal/degradation = -

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345 CONCLUSION

346 This study revealed that the location Sakpenwa Community, Tai L.G.A, Rivers State, is one
 347 of the oil exploration zone of southern Nigeria, harboring a large number of microbial
 348 hydrocarbon degraders which are able of utilizing Bonny light crude oil as carbon source
 349 when monitored. The increasing values obtained indicate that both test organisms: *Bacillus*
 350 sp. and *Pseudomonas* sp. can grow and utilizing Bonny light as carbon source but *Aciligenes*
 351 sp grow at a slower rate as the days of incubation increase further. The results of the
 352 chromatographic analyses of the total petroleum hydrocarbons (TPHs) also showed that
 353 *Acinetobacter* sp. can degrade Bonny light crude oil more efficient and this evidence is
 354 shown from the reductions of both the number, sizes of the peaks and in the percentage of
 355 degradation. Therefore, it appeared that *Acinetobacter* sp. a bacterium could be more useful
 356 in the degradation study contaminated with Bonny light crude oil and is recommended for
 357 controlling oil polluted site. This study revealed that indigenous bacterial species possess the
 358 requisite hydrocarbon degradation gene necessary for hydrocarbon degradation we concluded
 359 from the findings that this study revealed that indigenous bacterial species possess the
 360 requisite gene necessary for hydrocarbon biodegradation. This concluded that biodegradation
 361 is most often the primary mechanism for contaminant destruction including petroleum
 362 contaminants.

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