<u>Original Research Article</u> In Vitro Degradation and Reduction of Aromatic Hydrocarbons by Marine Bacteria Isolated from Contaminated Marine Environments of Niger Delta

7 **Aims:** To determine the *in vitro* degradation and reduction of aromatic hydrocarbons by 8 marine bacteria isolated from contaminated marine environments of Niger Delta. 9 Study Design: Nine treatments and nine controls designs were set up in triplicates 10 containing 100 mL of sterile modified mineral basal medium in 500 mL conical flasks 11 supplemented with 1 mg /L of xylene, anthracene and pyrene each; nine marine 12 hydrocarbon degraders and incubated at 24 °C for 24 days study. The nine treatments 13 and control set ups designated as ANT1, XYL2, PYR3, ANT4, PYR5, ANT6, XYL7, 14 15 XYL8, PYR9 and CTRL (Without hydrocarbons) were used to determine the aromatic 16 hydrocarbons degradability and transformation by the marine bacteria.

Place and Duration of Study: The studied sites were Abonema Wharf Water Front in
 Akuku-Toru Local Government Area, Nembe Water-side in Port Harcourt Local
 Government Area and Onne Light Flow Terminal Seaport located in Eleme Local
 Government Area of Rivers State, Nigeria between September, 2014 and March, 2017.

Methodology: A laboratory scale study was carried on six composite samples of the sediment and water samples from the three studied areas using enrichment, screening, phenotypical, degradation and TLC techniques.

Results: The results showed that the three studied areas harbour numerous promising aromatic degrading bacterial strains belonging to the genera: *Providencia, Alcaligenes, Brevundimonas, Myroides, Serratia,* and *Bacillus.* The bacterial strains especially *Serratia marcescens* XYL7 significantly (P = .05) had 99.50 ± 0.05 % and 60.00 ± 0.02 % degradations in weights of xylene and pyrene, respectively while *Alcaligenes faecalis* PYR5 significantly (P = .05) degraded 97.40 ± 0.01 % in weight of anthracene. TLC result revealed evidences of large spots size reductions or losses of test samples compare to control samples with minor spot sizes.

Conclusion: Thus, the outstanding degradative abilities of these strains could be exploited in bioremediation campaigns in Nigeria.

Keywords: Aromatic hydrocarbons, aquatic pollution, bioremediation, marine bacteria, Niger Delta.

41 1. INTRODUCTION

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43 The oil - rich Niger Delta region is the centre of exploration and production of petroleum in Nigeria. Over

44 80 -90 % of the Nigeria's crude oil originated from this important ecological area and its adjacent offshore

- 45 regions. Within the region, numerous causes of petroleum contamination may arise from transport
- 46 vehicles and mishap from tankers, oil fields and stations, pipelines and storage tanks [1]. As a result of
- 47 the activities mentioned above and from the petroleum companies, large scale contaminations of both the

terrestrial and aquatic ecosystems in the area have been reported and documented [2, 3, 4]. It is thus necessary to consider all remediation alternatives on the basis of their capacity to eliminate organic pollutants effectively as most of these waste products especially the benzene, toluene, and xylene (BTX) and polycyclic aromatic hydrocarbons (PAHs) are generally considered as toxicants [5]. The physical and chemical methods like volatilization, photooxidation, chemical oxidation, advanced oxidative processes and bioaccumulation are not safe and cost effective when compared to microbial bioremediation [6]. A healthier and economical way is to use enhanced biodegradation.

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Thus, much research has focused on the biological degradation of aromatics hydrocarbons (Ahs) through metabolism and co-metabolism [7]. Mao *et al.* [8] and Pathak and Bhatnagar [9], in their studies, recommended that supplementation and isolation of native aromatic hydrocarbon degraders is the most significant part of microbial degradation or bioremediation of aromatic hydrocarbons as the degraders have been well acclimatized leading to selection of better utilizers and diverse degraders of the hydrocarbons. Esedafe *et al.* [10] reported that an occurrence of 3/41 signifying 7.32 % of the isolates from refinery effluent were capable of utilizing phenanthrene and anthracene as sole carbon and energy

63 sources.

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65 Microbiological activity is affected by many environmental factors including energy source, donors and 66 acceptors of electrons, nutrients, pH, and temperature. Most microbes use respiration to transform 67 biological energy sources (organic and inorganic fuels, light) into ATP. Microorganisms especially 68 Prokaryotes employed a wide range of electron donors both organic carbon and inorganic compounds 69 such as H_2O , H_2 , as well as numerous alternative electron acceptors such as organic carbon, O_2 , CO_2 or 70 oxygen substitutes for respiration in the absence of molecular oxygen, which makes them exceptionally 71 adaptable with regard to energy [11]. Hydrocarbon degradation by microbial population in natural 72 ecosystem is influenced by physical, chemical and biological factors that contribute to the degradation of 73 petroleum and individual hydrocarbons [12]. Amongst the physical factors, temperature play a significant 74 role, firstly by directly changing the chemistry of pollutants and then secondly, by influencing the 75 physiology and diversity of the microbial environment [13]. Amongst the chemical factor, nutrients are

76 very essential requirements for effective biodegradation process, especially nitrogen, phosphorus and in 77 some cases iron [14]. Also, among the biological factor, the kind of microorganisms, their genomes and 78 their earlier contact to hydrocarbons affects the capacity with which the hydrocarbons could be degraded [15]. Several studies have reported that the growth rates of aromatic hydrocarbon degrading bacteria 79 80 were affected by the concentration of xylene compounds, temperature, pH of the medium and other 81 factors such as microbial number and nutrient availability [16]. The previous study reported that the growth rate of bacterial strains X1, X2, X3 and X4 under 1 % m-xylene decreased as the temperature 82 83 reduced from 30 - 25 °C, whereas at 45 °C, the growth rate was practically ceased. The optimum growth 84 rate of the strains was obtained at pH 6.5 but had fluctuating growth rate at pH 5.5 or 8.5 [16]. 85 86 The advancement in industrial and physical development as well as the enormous transportation activities 87 in the Niger Delta will definitely lead to increases in exploration, production, use, release and

bioaccumulation of aromatic hydrocarbons in the terrestrial and aquatic ecosystems with their concomitant health impacts/ risks on humans and animals population through food chain link. These facts prompted us to search for the presence of aromatic degrading microorganisms in crude oil - impacted Niger Delta aquatic ecosystem in particular for xylene, anthracene and pyrene hydrocarbon potent degraders as alternative to physical and chemical agents owing to greater advantages over them. In this work, we report the *in vitro* degradation and reduction of aromatic hydrocarbons by marine bacteria isolated from contaminated marine environments of Niger Delta, Nigeria.

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96 2. MATERIALS AND METHODS

97 **2.1 Description of the Sampling Sites**

The studied areas were chosen because high anthropologic and industrial acivities and are Abonema Wharf Water Front in Akuku-Toru Local Government Area, Nembe Water-side in Port Harcourt Local Government Area and Onne Light Flow Terminal Seaport located in Eleme Local Government Area of Rivers State. Abonema town is 53 Km and Abonema Wharf Water Front is 3 - 5 Km from Port Harcourt capital city; Nembe water side is located within Port Harcourt capital city of Rivers State, while Onne Light Flow Terminal is about 35 Km east from Port Harcourt capital city of Rivers State and 7 Km from Onne town. These sites were geo - referenced using Handheld Global Positioning System (GPS) GPSMAP 76
sc with the coordinates obtained from the sampling points or positions Abonema Wharf Water Front,
Nembe Water-side and Onne Light Flow Terminal Seaport were located between latitude 4°46'15.82"N to
latitude 4°46'38.01"N and longitude 7°0'0.54"E to longitude 7°0'34.82"E with average elevation of 4.1 m ,
latitude 4°45'8.72"N to latitude 4°45'26.42"N and longitude 7°1'11.37"E to longitude 7° 2'14.54"E with
average elevation of 2.7 m and latitude 4°41'32.58"N and 4°41'58.18"N and longitude 7°9'26.34"E and
7°10'48.82"E with average elevation of 2.3 m, respectively.

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112 **2.2 Sample of Collection**

113 Ten samples of marine sediment and water were collected randomly at ten (10) designated points in the 114 three sampling sites (Figures 1, 2 and 3). They were mixed together after which a total of six 115 representative sediment and water samples were taken for the analysis. The surface aerobic sediment 116 samples were obtained with a 95 % ethanol - sanitized plastic spatula at 5 cm depth inside 95 % ethanol -117 sanitized wide mouthed plastic containers. The water samples were collected at the air-water interface by 118 hand dipping the 95 % ethanol - sanitized cylindrical shaped 2 L plastic containers. The containers were wetted with the sediment and water samples before collecting the samples. All the composite or 119 120 representative sediment and water samples containers were placed into a sterile polythene bag in ice 121 packed coolers to keep them under a temperature not more than 4.00 °C. They were transported to the laboratory for microbiological analyses and stored at 4 °C in refrigerator for further processing [3, 17, 18]. 122

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124 **2.3 Isolation of Aromatic Hydrocarbon Degrading Bacterial Strains**

The hydrocarbon degraders were isolated from sediment and water samples of the three sampling sites using modified mineral basal agar (MBA) (4 g K₂HPO₄, 1.0 g (NH₄)₂SO₄, 0.1 g MgSO₄, 1.8 g KH₂PO₄, 0.1 g FeSO₄, 0.1 g NaCl, 0.2 g CaCl₂, 15 g Agar agar and distilled water 1,000 mL at pH 7.00 \pm 0.20) supplemented with xylene, anthracene and pyrene as sole carbon and energy sources. The medium was sterilized by autoclaving at 121 °C and 15 psi for 15 minutes. Thereafter, 0.2 mL acetone solution containing 0.1 % w/v of the representative aromatic hydrocarbons (xylene, anthracene and pyrene) were aseptically pipetted and uniformly spread on the agar surface of the Petri dish plates. After the acetone evaporation, 0.1 mL aliquots of the 10^{-3} dilutions were spread plated on the surfaces of the solidified media with the aid of a glass spreader under aseptic conditions. The inoculated plates were sealed using adhesive tape and foil to prevent contamination and photolysis and later placed in black polythene bags. They were incubated in the dark at 28.00 ± 2.00 °C for 14 days [15, 18].

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137 **2.4 Preservation of Cultures**

Colonies that developed on MBA plates were sub-cultured onto new MBA plates and incubated for another 14 days. Forty - eight isolates that grew on these plates were selected as xylene, anthracene and pyrene degraders. They were later sub - cultured on Bijou bottles and preserved at 4.00 °C in refrigerator [18].

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143 2.5 Screening Test for the Most Potent Marine Degrading Bacterial Isolates

In order to screen and chose the most potent degrading bacterial strains, the growths of the forty-eight 144 145 isolates was tested by growing 5 mL of each isolates in large test tubes containing 25 mL of the modified mineral basal medium (MBM) augmented with 100 mg /L of xylene, anthracene and pyrene hydrocarbons 146 147 which were dissolved in acetone and added to each tube after autoclaving. Thereafter, the test tubes were incubated at 28.00 ± 2.00 °C for five days. After incubation, growths of bacterial cultures as 148 indicated by turbidity were measured at 600 nm using a UV - VIS spectrophotometer (Astell, UV - Vis 149 Grating, 752 W) and the cultures with the highest optical densities on each hydrocarbon in three studied 150 151 sites were chosen as the most potent xylene, anthracene and pyrene degrading bacteria [18, 19, 20]. 152

153 **2.6 Phenotypical identification of selected hydrocarbon utilizing bacterial isolates**

154 **2.6.1 Morphological characterization**

155 2.6.1.1 Colonial characteristics

Following sub - culturing and incubation, the colonial features such as shape, elevation, margin, optic,
texture, colour, size and surface descriptions of the nine chosen bacterial strains were observed and
noted [21].

160	2.6.1.2	Microscopic	characteristics
100	2.02		on a docorroction

- 161 The standard methods of Gram staining and endospore staining were carried out on the chosen bacterial
- 162 strains as described in Cheesbrough [22] and Health Protection Agency [23].
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164 **2.6.2 Biochemical characteristics**

The standard methods of testing for catalase, indole, motility, methyl red – Voges Proskauer, citrate, urease, starch hydrolysis, gelatin, nitrate reduction, coagulase, Hydrogen sulphide production, sugar fermentation, oxidase and casein hydrolysis tests were carried out as described in Willey *et al.* [21] and Cheesbrough [22].

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170 **2.6.3 Identification of the marine bacterial isolates**

Following morphological and biochemical characterization, the isolates were identified using Bergey's manual for determinative bacteriology, as described in Holt *et al.* [24].

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174 **2.7 Biodegradation Study and Analytical Technique**

175 Following the methods of Bennet et al. [25] and John and Okpokwasili [12] as modified in this study, the 176 degradation rates of bacterial isolates were determined using hydrocarbon supplemented MBM as 177 previously stated. Precisely, 1 mL of 48 hrs old cultures of each bacterial strains was inoculated into 28 178 sterile 200 mL capacity conical flasks (4 sets of 7 flasks) in triplicates comprising 100 mL of sterile MBM augmented with 100 mg /L of xylene, anthracene and pyrene aromatic hydrocarbons respectively. The 179 treatment set ups were incubated at 24.00 °C for 24 days. During incubation, 5 mL representative 180 samples from the different sets of flasks were pipetted at intervals of 0, 4, 8, 12, 16, 20 and 24 days and 181 182 the temperature, pH and bacterial growth (optical densities (OD_{600 nm})) of the media in different flasks 183 were measured. The residual aromatic hydrocarbons were determined spectrophotometrically using ethyl 184 acetate as the extraction solvent. For each sample, 5 mL of ethyl acetate was added and vigorously shaken manually. The organic and water layers from the media were separated by centrifugation at 5000 185 rpm for 20 mins. The water layers were disposed off while the organic layers were analyzed with UV - VIS 186

- 187 spectrophotometer at 240 nm wavelength (Astell UV Vis Grating, 752 W). The percentages of
 188 biodegradation of the various hydrocarbons were determined as follows:
- 189 % degradation $= \frac{a-b}{a} \times \frac{100}{1}$
- 190 Where a = the mean absorbance of the medium before incubation; b is the mean absorbance of the 191 medium after each 4th day of the incubation period.
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- 2.8 Thin layer chromatographic (TLC) analysis for the Degraded Aromatic Hydrocarbons 193 194 This was carried out according to the method of Bennet et al. [26] as modified in this study in order to 195 check whether 100 mg /L of the aromatic hydrocarbons from the MBM as stated above were degraded 196 and reduced into lesser and minor forms and parts by the existence of either small spots or by reduction in sizes in comparison to the controls. On a clean glass chamber, mixture of hexane/benzene: methanol 197 198 at 25 mL and 5 mL proportion was prepared as solvent system for the chromatogram. One microliter (1 199 μL) of the organic phases of the three strains with the highest percentage of degradation on each of the 200 three aromatic hydrocarbons was spotted and marked at the lower end of the TLC plate. Similarly, control 201 spots of xylene, anthracene and pyrene solutions with medium were placed. Plates were placed inside 202 the glass chamber and were again covered. Once the solvent front reached the top layer of the plate, the developed chromatogram was taken out and air - dried at 28.00 ± 2.00 °C. It was then visualized first with 203 204 UV light (235 nm) and lastly with iodine crystals. Test and control samples were compared from one another following the principles of light intensity and spot sizes. 205
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207 2.9 Analysis of Data

- 208 All values were expressed as mean ± standard deviation and analyzed using Graph-Pad Prism statistical
- 209 software version 7.00. Test of significance was performed on the data obtained using ordinary one-way
- 210 analysis of variance (ANOVA) followed by post Tukey's, multiple comparison test. The results were
- 211 considered statistically significant if the probability is less than .05 (P = .05) [19, 26].
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- 213 3. RESULTS
- 214 3.1 Screening Test

215 The result of the optical density (OD₆₀₀ nm) determination of the aromatic hydrocarbon bacterial 216 degraders isolated from Abonema sampled location is presented in Figure 4. From the result, strains 217 XYL2, ANT4 and PYR3 out of the 13 strains isolated were found to have the highest significant (P = .05) 218 optical densities of 0.952 \pm 0.004, 0.775 \pm 0.007 and 1.041 \pm 0.008 on xylene, anthracene and pyrene 219 <mark>aromatic</mark> hydrocarbons. The result of the <mark>optical density (OD₆₀₀ nm) determination</mark> of the aromatic 220 hydrocarbon bacterial degraders isolated from Nembe sampled location is presented in Figure 5. From 221 the result, strains XYL7, ANT1 and PYR5 out of the 17 strains isolated were found to have the highest significant (P = .05) optical densities of 1.055 ± 0.002, 0.816 ± 0.007 and 0.933 ± 0.007 on xylene, 222 223 anthracene and pyrene aromatic hydrocarbons. The result of the optical density (OD₆₀₀ nm) determination 224 of the aromatic hydrocarbon bacterial degraders isolated from Onne sampled location is presented in 225 Figure 6. From the result, strains XYL8, ANT6 and PYR9 out of the 18 strains isolated were found to have 226 the highest significant (P = .05) optical densities of 0.741 ± 0.007, 1.433 ± 0.013 and 0.871 ± 0.001 on 227 xylene, anthracene and pyrene hydrocarbons. On the basis of these results, strains ANT1, XYL2, PYR3, 228 ANT4, PYR5, ANT6, XYL7, XYL8 and PYR9 were chosen as the most potent degraders of xylene, anthracene and pyrene aromatic hydrocarbons. 229

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231 3.2 Taxon of the Degrading Bacterial Isolates

232 The result of the macroscopic description of degrading bacterial isolates is presented in Table 1. From the 233 result, most colonies were circular and irregular in shape, flat in elevation, undulate in margin, translucent 234 in optic, smooth in texture, creamy in colour, 4 mm in size and shiny in surface description. The result of 235 the microscopic and biochemical features of the degrading bacterial isolates is presented in Table 2. 236 From the result, most bacterial isolates were Gram negative in Gram reaction, rod shaped arranged in 237 single or pair, negative to spore, indole, methyl red, Voges Proskauer, urease, gelatin, nitrate reduction, 238 coagulase, hydrogen sulphide production, xylose, lactose, arabinose, maltose and casein hydrolysis tests 239 while positive to catalase, motility, citrate, starch hydrolysis, mannitol, glucose, sucrose, saccharose and 240 oxidase tests.

241 **3.3 Metabolism and Kinetics of Degradation**

242 The result of the changes in temperature (°C), pH and optical density (OD₆₀₀ nm) of medium during 243 aromatic hydrocarbon degradation by marine bacterial isolates are presented in Tables 3, 4 and 5 while 244 the results of the percentage weight reductions of xylene, anthracene and pyrene hydrocarbons by marine bacterial isolates are presented in Figures 7, 8 and 9. From the Tables 3 - 5 results, temperature 245 fluctuated between 25.5 and 26 °C for xylene and anthracene; and between 25 and 26 °C for pyrene 246 respectively while the control had 26 °C throughout the 24 days' period of the study. The pH fluctuated 247 between 6.85 and 7.15 for xylene; 6.97 and 7.11 for anthracene; 6.95 and 7.19 for pyrene respectively. 248 249 The pH remained almost neutral (7.05 - 7.00) in the control. The pH also changed a little within the 24 250 days' period of the study. The optical density (OD_{600} nm) increased ranging from 0.01 - 0.35 for xylene; 251 0.11 - 0.36 for anthracene and 0.17 - 0.54 for pyrene respectively as well as ranged from 0.10 - 0.15 in 252 the control. The optical density (OD₆₀₀ nm) increase from 0.01 - 0.54 within the 24 days' period of the 253 study. Similarly, from the Figures 7 - 9 results, Serratia marcescens XYL7 had the maximum percentage degradations and reductions in weight of 99.50 ± 0.05 % and 60.00 ± 0.02 % for residual xylene and 254 pyrene hydrocarbons respectively while Alcaligenes faecalis PYR5 had highest percentage degradation 255 and reduction in weight of 97.40 ± 0.01 % for residual anthracene hydrocarbons after 24 days 256 257 biodegradation study.

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259 **3.4 Thin Layer Chromatography Study of the Organic Layer**

260 The results of the thin layer chromatograms of xylene, anthracene and pyrene hydrocarbon degradations in the control set-up and their highest degraders are shown in Plates 1a - b, 2a - b and 3a - b. The 261 262 results revealed that all the control set ups possess high and large intensity spots while the highest 263 degraders Serratia marcescens XYL7 and Alcaligenes faecalis PYR5 had low and small intensity spots 264 with evidence of spot size reductions when visualized under UV illuminator at 235 nm. The result of the 265 retention factor of the aromatic hydrocarbons degraded by marine isolates as presented in Table 6 266 ranged from $0.27 \pm 0.08 - 0.84 \pm 0.01$, $0.74 \pm 0.08 - 1.00 \pm 0.12$ and $0.06 \pm 0.03 - 1.00 \pm 0.12$ compared 267 to controls which were 0.5 ± 0.01 , 0.63 ± 0.02 and 0.38 ± 0.01 for xylene, anthracene and pyrene 268 hydrocarbons.

270 4. DISCUSSION

271 The use of microbial bioremediation as an economical and environmental - friendly treatment tool for the 272 protection of certain petroleum - polluted oceans, seas, estuaries and shoreline has gained extensive 273 and growing interest since mechanical, physical and chemical alternatives are inadequate and ineffective 274 [8]. Petroleum polluted soils and waters are potent sources of efficient hydrocarbon degraders as they are 275 recommended for biodegradation and bioremediation treatment options [27]. In this study, a total of nine 276 (9) strains ANT1, XYL2, PYR3, ANT4, PYR5, ANT6, XYL7, XYL8 and PYR9 out of the 48 isolates (9/48) 277 suggesting 18.75 % of the isolates screened were chosen as the best adapted and potent degraders of xylene, anthracene and pyrene aromatic hydrocarbons which they use as sources of carbon and energy 278 279 and are indicated by the highest optical densities for each strains and hydrocarbons (Figures 4, 5 and 6). 280 The result in Table 1 and Table 2 showed that most of the marine bacterial isolates were Gram negative 281 in Gram reaction, rod shaped arranged in single or pair, with variable reactions to different biochemical 282 tests. The marine bacteria were identified as Providencia vermicola strain ANT1, Alcaligenes faecalis 283 strain XYL2, Brevundimonas diminuta strain PYR3, Alcaligenes faecalis strain ANT4, Alcaligenes faecalis 284 strain PYR5, Myroides odoratus strain ANT6, Serratia marcescens strain XYL7, Providencia sp. strain 285 XYL8 and Bacillus cereus strain PYR9 using Bergey's manual for determinative bacteriology by Holt et al. 286 [24]. These findings agree with the reports of Isiodu et al. [26] and Fagbemi and Sanusi [28] that two -287 third of most petroleum hydrocarbon degraders are Gram negatives with one - third being Gram positives. 288 Previous studies have reported that a lot of rod - shaped bacteria have been implicated in hydrocarbon 289 degradation studies and similar result was obtained in this study [26, 28, 29].

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The result in Table 3 revealed that the temperature remained in optimal level of 26 °C conducive for the bacteria activities which is also observable in the control. The strains isolated in this study were mesophilic in nature therefore proved that mesophilic bacteria can degrade hydrocarbons. Similar result were observed by Athar *et al.* [30]. The result contradicts the report of Irshaid and Jacob [16] that the decrease in temperature from 30 - 25 °C led to decrease in growth rates of xylene degrading bacteria from gasoline contaminated soil sites while whereas the growth rates almost terminated at 45 °C,. This contradiction might be due to the nature of the ecosystem as marine environment is generally 298 characterized with lower temperatures unlike the terrestrial ecosystem with higher temperatures. The 299 result in Table 4 showed that the drops in pH are probably because the degradation of the hydrocarbons 300 by these bacteria resulted in the discharge of acidic substances and intermediates (organic acids and 301 other metabolic products) which reduces the pH of the medium and is in agreement with the researches 302 carried out by previous studies [16, 25, 31]. The result in Table 5 revealed that the degradation and 303 utilization of these compounds resulted in increase in optical density (cell mass) of the organisms and 304 corroborates with findings of John et al. [18] and Akinbankole et al. [32]. The previous study reported that 305 cell growth study of anthracene and pyrene metabolizing bacteria revealed that only two strains KLA1022 306 (B. toyonesis) and JIP1005 (S. enterica) could utilize anthracene and pyrene hydrocarbons as source of 307 nutrient for their growth and metabolism because there was substantial rise in the quantity of cells for the 308 both strains. A decline in cell quantity was realized as the PAH concentration (sole carbon source) was 309 decreased [32]. These differences could be attributed to the acclimatization of these bacterial species to 310 exploit and thrive in the presence of petroleum or might be partly due to discrepancies among several 311 physicochemical environmental factors, including divergent ambient environmental conditions, sediment 312 organic carbons and structures [16].

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314 The results in Figures 7, 8 and 9 revealed that the degree of weight reductions was observed to rise with 315 increase in incubation period but varied with different microbial species tested. Serratia marcescens XYL7 316 significantly demonstrated the highest abilities to degrade the aromatic hydrocarbons with 99.50 \pm 0.05 % 317 and 60.00 ± 0.02 % weight reductions in xylene and pyrene respectively while Alcaligenes faecalis PYR5 318 significant degraded anthracene with 97.40 ± 0.01 % weight reduction after 24 days biodegradation study 319 with proof of increase low optical density (OD_{600 nm}) against their controls. However, the levels of xylene, 320 anthracene and pyrene degradations also included 5.20 \pm 0.03 %, 4.60 \pm 0.01 %, 11.10 \pm 0.06 % 321 degradation by abiotic factor as detected in controls with no bacterial inocula. After deduction of xylene, 322 anthracene and pyrene degradation by the abiotic factor, 94.30 ± 0.02 % and 49.90 ± 0.06 % of xylene 323 and pyrene degradation was in fact aided by Serratia marcescens XYL7 while 92.80 \pm 0.02 % of anthracene by Alcaligenes faecalis PYR5 during this period. Discrepancies in xylene, anthracene and 324 325 pyrene degradation abilities of the bacterial strains were evidently revealed even after 4 days of 326 incubation which was further intensified during incubation periods (4 – 24 days) with very strongly significant positive relationship (r = 0.897 - 0.996). Thus, xylene, anthracene and pyrene were degraded 327 328 by all the nine bacterial strains, but they varied extensively in their innate capabilities. These disparities 329 could be due to high molecular weight aromatic hydrocarbons that are resistant to microbial attacks and 330 the order of degradation is xylene > anthracene > pyrene. Biodegradation of aromatic hydrocarbons is 331 depended on their chemical configurations and equivalent physiochemical properties and low molecular 332 weight aromatic hydrocarbons are rapidly degrade rapidly than high molecular weight aromatic 333 hydrocarbons [33]. Similar results are obtained by Akpe et al. [5] who observed that the higher 334 percentage of hydrocarbons degraded, the lower the optical density value. Previous study reported that 335 Rhodococcus pyridinvorans NJ2 was the highest degrader (60 %) of pyrene, followed by Pseudomonas 336 sp. BP10 (44 %) and the least was *Ochrobactrum intermedium* P2 (42 %) in MSM with pyrene (50 µg 337 /mL) in 8 days [31]. In another previous study, it was reported that PAH (phenanthrene, flourene, 338 anthracene, pyrene) dissipation level ranged between 38.7 to 99.7 % with highest depletion recorded in 339 phenanthrene within seven days. The degradation rate of 3 - ring PAH was higher as compared to 4 - ring PAH (pyrene) by Serratia marcescens L - 11 [34]. The findings of these authors supported the results of 340 341 this study.

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343 In order to The results in Plates 1a – b, 2a – b and 3a – b evidently revealed that the 100 mg /L of 344 xylene, anthracene and pyrene hydrocarbons from the assay medium were degraded and reduced into 345 lesser, smaller and lighter fragments/sizes with clear confirmations of biodegradation losses of some 346 aromatic hydrocarbon components. These losses, disappearances or reduction in spot sizes could be 347 ascribed to the usage of aromatic hydrocarbons by marine isolates during the 24 days incubation period 348 under ideal cultural settings of pH and temperature as sole carbon sources. Moreover, the findings upheld 349 the results of the kinetics of degradation of the hydrocarbons by the marine bacterial isolates at day 24 350 with increasingly low OD monitored at 240 nm and is in agreement with published work of Bennet et al. 351 [25] who reported that the thin layer chromatography of the test anthracene showed small fragments with 352 low intensities when compared with control having larger fragments with high intensities as visualized at 353 235 nm OD - UV illuminator. Previous study reported that the thin layer chromatographic study of the

354	purified metabolites after the pyrene degradation course showed the presence of a single spot with Rf
355	value of 0.65, which is similar to the reference Rf value of protocatechuic acid [35] and conformed with
356	the pyrene Rf value in this study (Table 6). Also, in agreement with another previous study, the losses of
357	pyrene and anthracene in the medium showed that they have been metabolized by the degrading
358	bacteria [Akinbankole <i>et al.</i> [32].
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309	Table 1 Magrospopie description of the degrading besterial isolator
370	Table 1. Macroscopic description of the dediading pacterial isolates

Isolate		Colonial description									
	Shape	Elevation	Margin	Optics	Texture	Colour	Size	Surface			
ANT1	Circular	Flat	Undulate	Translucent	Smooth	Creamy	4.0 mm	Dull			
XYL2	Irregular	Raised	Erose	Translucent	Smooth	Creamy	3.0 mm	Dull			
PYR3	Circular	Flat	Undulate	Translucent	Smooth	Creamy	2.0 mm	Shiny			
ANT4	Circular	Flat	Erose	Translucent	Smooth	Creamy	5.2 mm	Shiny			
PYR5	Circular	Flat	Undulate	Translucent	Smooth	Creamy	4.0 mm	Dull			
ANT6	Rhizoid	Flat	Lobate	Translucent	Rough	Yellow	4.2 mm	<mark>Shiny</mark>			
XYL7	Irregular	Flat	Undulate	Translucent	Rough	Red	4.0 mm	Shiny			
XYL8	Irregular	Flat	Undulate	Translucent	Smooth	Creamy	4.2 mm	Shiny			
PYR9	Irregular	Raised	Undulate	Translucent	Smooth	Creamv	4.0 mm	Shiny			

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Property Isolate												
	ANT1	XYL2	PYR3	ANT4	PYR5	ANT6	XYL7	XYL8	PYR9			
Gram reaction	_	_	_	_	_	_	_	_	+			
Cellular morphology	Paired short rods	Singled long rods	Singled long rods	Singled short rods	Singled long rods	Single longer rods	Paired short rods	Single paired long Rods	Singled short rods			
Spore test	_	_	_	_	_	_	_	-	+			
Catalase	+	+	+	+	+	+	+	+	+			
Indole	+	_	_	_	_	_	_	+	+			
Motility Methyl red	+ _	+ _	+ _	+ _	+ _	_	+	+	+ +			
Voges - Proskauer	_	_	_	_	_		+	<u> </u>	_			
Citrate	_	+	_	+	+	+	+	_	+			
Urease	+	_	_	_	-	X	_	+	+			
Starch hydrolysis	+	+	_	+	-	+	_	+	+			
Gelatin hydrolysis	_	_	_	- /	5	-	+	_	_			
NO ₃ reduction	+	-	_		-	-	+	+	+			
Coagulase test	+	-	-	X	-	_	+	+	-			
H ₂ S production	+	_		-	_	+	-	+	+			
Mannitol	+	_		_	_	+	+	+	+			
Glucose	+	+	_	+	+	+	+	+	+			
Xylose	_		_	_	_	_	+	_	+			
Lactose	_	- X	_	_	_	_	_	_	+			
Sucrose	-	+	_	+	+	_	+	_	+			
Arabinose	+		_	_	_	+	+	_	_			
Maltose	-)	_	+/_	_	_	_	_	_	+			
Saccharose	+	+	_	+	+	_	+	+	+			
Oxidase	-	+	+	+	+	+	+	_	+			
Casein hvdrolvsis	_	_	-	_	_	-	+	_	_			

388 Table 2. Microscopic and biochemical features of the degrading bacterial isolates

NO = Nitrate; H S = Hydrogen sulphide; - = Negative result; + = Positive result 3

Isolate								Ahs										
			Xyl						Ant						Pyr			
	4	8	12	16	20	24	4	8	12	16	20	24	4	8	12	16	20	24
ANT1	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
XYL2	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
PYR3	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
ANT4	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
PYR5	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
ANT6	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
XYL7	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
XYL8	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
PYR9	26.00	26.00	25.50	26.00	26.00	26.00	26.00	26.00	26.00	25.50	26.00	26.00	26.00	25.00	25.50	26.00	26.00	26.00
Control	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	26.00	25.00	26.00	26.00	26.00	26.00

Table 3. Changes in temperature (°C) of medium during aromatic hydrocarbon degradation by marine bacterial isolates

Xyl = Xylene; Ant = Anthracene; Pyr = Pyrene; AHs = Aromatic hydrocarbons; 4 - 24 represents days of incubation.

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Isolate									AHs									
	Xyl									Ant								
	4	8	12	16	20	24	4	8	12	16	20	24	4	8	12	16	20	24
ANT1	7.05	7.11	7.05	7.04	7.15	7.02	7.06	6.98	7.00	7.07	7.02	7.01	7.06	6.97	7.00	7.00	7.03	6.99
XYL2	7.06	7.00	7.07	7.06	7.05	7.02	7.00	6.97	7.02	6.99	7.02	7.00	7.08	6.97	7.08	7.08	7.02	6.98
PYR3	6.88	6.86	6.85	6.87	6.95	6.96	7.03	7.01	7.06	7.00	7.03	7.02	7.09	6.97	7.05	7.05	7.01	7.01
ANT4	7.03	7.00	7.03	7.05	7.01	7.02	7.03	7.01	7.09	7.05	7.02	7.01	7.06	6.96	7.03	7.03	7.02	6.98
PYR5	7.02	6.97	7.01	7.08	7.02	7.04	7.03	6.98	7.10	7.09	7.04	7.01	7.07	6.99	7.19	7.07	7.00	6.97
ANT6	7.05	6.97	7.03	7.06	7.04	7.05	7.04	7.00	7.06	7.05	7.01	7.02	7.09	6.98	7.01	7.06	7.00	6.98
XYL7	7.06	7.02	7.03	7.03	7.02	7.03	7.04	7.00	7.06	7.05	7.01	7.02	7.06	6.95	7.11	7.01	7.00	7.03
XYL8	7.06	7.03	6.99	7.03	7.01	7.04	7.02	6.99	7.11	7.00	6.99	6.97	7.06	6.95	7.13	7.04	6.98	7.04
PYR9	6.99	6.96	6.97	7.01	6.99	7.02	7.05	7.04	7.07	7.02	7.01	6.98	7.04	6.95	7.16	7.02	6.99	6.99
Control	7.05	7.05	7.05	7.05	7.05	7.05	7.03	7.03	7.03	7.03	7.03	7.03	7.00	7.00	7.00	7.00	7.00	7.00

Table 4. Changes in pH of the medium during aromatic hydrocarbon degradation by marine bacterial isolates

Xyl = Xylene; Ant = Anthracene; Pyr = Pyrene; AHs = Aromatic hydrocarbons; 4 - 24 represents days of incubation

Isolate									AHs									
			Xyl						Ant	Ant					Pyr			
	4	8	12	16	20	24	4	8	12	16	20	24	4	8	12	16	20	24
ANT1	0.17	0.18	0.19	0.20	0.21	0.22	0.20	0.21	0.23	0.24	0.25	0.26	0.17	0.19	0.20	0.23	0.25	0.27
XYL2	0.03	0.05	0.06	0.08	0.09	0.10	0.14	0.15	0.19	0.20	0.22	0.23	0.25	0.26	0.27	0.30	0.44	0.54
PYR3	0.01	0.02	0.05	0.07	0.08	0.09	0.25	0.26	0.27	0.28	0.29	0.30	0.19	0.20	0.23	0.24	0.26	0.51
ANT4	0.11	0.12	0.13	0.14	0.15	0.35	0.15	0.16	0.19	0.20	0.25	0.28	0.21	0.22	0.24	0.26	0.28	0.34
PYR5	0.20	0.25	0.28	0.30	0.32	0.34	0.11	0.13	0.15	0.17	0.19	0.21	0.18	0.20	0.22	0.24	0.26	0.28
ANT6	0.11	0.14	0.17	0.20	0.23	0.35	0.24	0.25	0.27	0.30	0.35	0.36	0.18	0.19	0.20	0.21	0.23	0.25
XYL7	0.11	0.12	0.13	0.14	0.15	0.16	0.11	0.14	0.17	0.19	0.20	0.21	0.30	0.31	0.35	0.37	0.40	0.51
XYL8	0.13	0.14	0.15	0.17	0.18	0.20	0.16	0.17	0.19	0.21	0.22	0.23	0.30	0.30	0.35	0.37	0.40	0.45
PYR9	0.18	0.20	0.22	0.23	0.24	0.25	0.14	0.16	0.18	0.20	0.21	0.23	0.25	0.27	0.30	0.35	0.40	0.45
Control	0.10	0.10	0.10	0.10	0.10	0.10	0.15	0.15	0.15	0.15	0.15	0.15	0.15	0.15	0.15	0.15	0.15	0.15

Table 5. Changes in the optical density (ODeco nm) of the medium during aromatic hydrocarbon degradation by marine bacterial isolates

Xyl = Xylene; Ant = Anthracene; Pyr = Pyrene; AHs = Aromatic hydrocarbons; 4 - 24 represents days of incubation.

Isolate		AHs	
	Xylene	Anthracene	Pyrene
ANT1	0.27 ± 0.08	0.74 ± 0.08	0.34 ± 0.08
XYL2	0.69 ± 0.07	0.86 ± 0.01	0.24 ± 0.02
PYR3	0.76 ± 0.08	0.86 ± 0.01	1.00 ± 0.12
ANT4	0.82 ± 0.08	0.98 ± 0.08	0.83 ± 0.08
PYR5	0.82 ± 0.08	1.00 ± 0.12	0.92 ± 0.01
ANT6	0.71±0.01	0.41 ± 0.01	0.59 ± 0.01
XYL7	0.84 ± 0.01	0.82 ± 0.08	0.38 ± 0.01
XYL8	0.32 ± 0.01	1.00 ± 0.12	0.79 ± 0.01
PYR9	0.30 ± 0.01	0.88 ± 0.01	0.06 ± 0.03
Control	0.50 ± 0.01	0.63 ± 0.02	0.38 ± 0.01

Table 6. Retention factor of the aromatic hydrocarbons degraded by marine bacterial isolates

 $Retention \; factor \; (R_F) = \frac{\text{distance travelled by solute}}{\text{distance travelled by solvent}}$

AHs= Aromatic hydrocarbons



Α



A. Thin layer chromatogram of xylene hydrocarbon degradation in control sample with arrow showing large high intensity spot visualized under UV illuminator at 235 nm B. Thin layer chromatogram of Serratia marcescens XYL7 on xylene hydrocarbon degradation with arrow showing small low intensity spot visualized under UV illuminator at 235 nm.

В



Plate 2a – b. Thin layer chromatograms of anthracene hydrocarbon degradation

A. Thin layer chromatogram of anthracene hydrocarbon degradation in control sample with arrow showing large high intensity spot visualized under UV illuminator at 235 nm B. Thin layer chromatogram of Alcaligenes faecalis PYR5 on anthracene hydrocarbon degradation with arrow showing almost cleared low intensity spot visualized under UV illuminator at 235 nm.





A. Thin layer chromatogram of pyrene hydrocarbon degradation in control sample with arrow showing large high intensity spot visualized under UV illuminator at 235 nm B. Thin layer chromatogram of Serratia marcescens XYL7 on anthracene hydrocarbon degradation with arrow showing small low intensity spot visualized under UV illuminator at 235 nm.



Fig 1. Geoeye satellite image (2016) showing the Abonema sample points



Fig 2. Geoeye satellite image (2016) showing the Nembe sample points







Marine bacterial strains



449 from Abonema sampled location

450 *= Isolates with highest degradability; values are mean ± standard deviation of triplicate determination.





Marine bacterial strains

461 Fig 6. Optical density (OD₆₀₀ nm) determination of the aromatic hydrocarbon bacterial degraders isolated from Onne sampled location * = Isolates with highest degradability; values are mean ± standard deviation of triplicate determination.

454





Fig 7. Percentage weight reductions of xylene hydrocarbons by marine bacterial isolates * = Isolates with highest percentage degradability; values are mean ± standard deviation of triplicate determination.



468 469 470

Fig 8. Percentage weight reductions of anthracene hydrocarbons by marine bacterial isolates * = Isolates with highest percentage degradability; values are mean ± standard deviation of triplicate determination.





Fig 9. Percentage weight reductions of pyrene hydrocarbons by marine bacterial isolates *= Isolates with highest percentage degradability; values are mean ± standard deviation of triplicate determination.

476 **5. CONCLUSION**

- 477 This study has shown that all the studied sites are sources of potent aromatic hydrocarbon degrading
- 478 bacterial strains belonging to the genera: Providencia, Alcaligenes, Brevundimonas, Myroides, Serratia,
- 479 and Bacillus. It also showed that the isolated bacteria especially Serratia marcescens XYL7 were able to
- 480 significantly (*P* = .05) degrade mono and polyaromatic hydrocarbons. The growth rates for these isolates
- 481 were moderately affected by temperature, pH and concentration of aromatic hydrocarbons. There were
- 482 clear confirmations of biodegradation losses, disappearances or reduction in spot sizes of the aromatic
- 483 hydrocarbon components indicating that the hydrocarbons in the medium have been reduced into
- 484 smaller, lesser and lighter fragments by these bacteria. Thus, the promising potentials of these bacterial
- 485 strains especially Serratia marcescens XYL7 with regards to in vitro degradations and reductions of these
- 486 pollutants could be exploited in bioremediation campaigns in Nigeria.
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