Screening and Characterization of Hydrocarbonoclastic Bacteria Isolated from Oil-contaminated Soils from Auto Garages

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To cite this article:

Received: January 10, 2018; Accepted: February 1, 2018; Published: March 23, 2018

Abstract: Release of petroleum oil and its products into the environment is a worldwide concern. The present study focused on isolation, molecular identification, morphological and biochemical characterization of bacteria possessing hydrocarbon-degrading properties. The study also aimed at optimizing appropriate culture conditions for the isolates as well as screening for alkane hydroxylase enzyme. Out of twenty one microbes isolated, nine were selected based on their ability to utilize different hydrocarbons. The isolates were observed to mineralize heating oil, hexane, octane, toluene and diesel oil. PCR amplification of 16S rDNA gene revealed that the isolates belong to six different genera; Pseudomonas, Acinetobacter, Klebsiella, Enterobacter, Salmonella and Ochrobactrum. Based on their ability to degrade diesel oil, three isolates were selected and their growth conditions optimized. Optimum degradation was recorded at less than 1% substrate concentration, pH 7 and temperature range of between 30°C and 37°C and using yeast extract as nitrogen source. GC-MS analyses confirmed their diesel oil degrading properties. Alkane hydroxylase gene from one of the isolates (isolate 1C) was successfully amplified indicating its catabolic capabilities in degrading alkanes. Overall, the characterized isolates may constitute potential candidates for biotechnological application in environmental cleanup of petroleum contaminants.

Keywords: Oil-Contaminated Soils, Bacteria, Biodegradation, Petroleum Hydrocarbons

1. Introduction

Petroleum-based products are a principle source of energy for industries and daily life, making them a vital commodity central to the global economy [1]. They originate from crude oil whose main constituents are hydrocarbon compounds derived from ancient algae and plant remains found in reservoirs under the earth’s surface [2]. Petroleum products are divided into four classes: saturates, aromatics, resins and asphaltenes [3]. Accidental release of petroleum products occur regularly during exploration, production, refining, transportation, utilization and storage [4]. It is estimated that globally, approximately 1.7-8.8 million metric tons of petroleum hydrocarbons are released into marine ecosystems annually [5]. Water and soil pollution is a worldwide environmental problem that is of particular concern since it leads to uptake and accumulation of toxic substances including petroleum products in food chains consequently harming the flora and fauna [6]. Current conventional disposal methods of petroleum products include physicochemical techniques such as photo-oxidation, burying, dispersion, washing, incineration, thermal conversion and other pyrolysis techniques [5,7]. However, many of these methods are expensive and can result in incomplete decomposition of oil products. In addition, physicochemical methods such as volatization, photo-oxidation and chemical oxidation are rarely successful in rapid removal of hydrocarbon contaminants especially the aromatics [8].
Bioremediation is emerging as one of the most promising technologies for environmental removal of petroleum contaminants [1]. It is a process through which microorganisms metabolize contaminants through oxidative/reductive processes. Bioremediation can be carried out either through addition of oil degrading microbes into the soil in a process referred to as bio-augmentation or through provision of appropriate conditions and/or amendments (e.g. supplying oxygen, moisture and nutrients) for growth of the microorganisms in the soil, a process known as bio-stimulation [4]. Studies have shown that petroleum-based products can primarily be eliminated from the environment by microbes such as bacteria, yeast, fungi and microalgae [1]. Bacteria however play a major role in biodegradation of these hydrocarbon compounds. Some important bacterial species with this potential are of the genera *Bacillus*, *Arthrobacter*, *Halononas*, *Pseudomonas*, *Klebsiella*, *Proteus* among others [9]. These microbes completely degrade (mineralize) organic compounds into non-toxic end products that include carbon dioxide and water or organic acids and methane [6]. Bioremediation is therefore an effective technique that takes advantage of the versatility of microbes to completely degrade petroleum compounds into innocuous end products. Apart from being environmentally friendly, the method is also cost effective for treatment of oil pollution compared to physicochemical methods [10].

Although numerous studies have been conducted on microbial species capable of cleaning up petroleum contaminants around the world, Kenya has not adopted this emerging technology and no data exists on its potential application in environmental conservation. The significance of this work is that this is the first report on isolation and characterization of oil degrading bacteria in the country. With the ongoing oil explorations in East Africa region, accidental oil spills are likely to occur during drilling or transportation. The objective of the current study was therefore to screen, characterize (using morphological, biochemical and molecular methods) and optimize appropriate culture conditions for oil degrading microbes that would be best suited to degrade petroleum-based contaminants. This will provide insight on an effective strategy that can be used to clean up accidentally released petroleum oil and hence provide a possible effective oil spill response management strategy to oil prospecting, refining and transporting companies and the general public.

2. Materials and Methods

2.1. Sample Collection

Soil samples from the surfaces to a depth of about 15 cm were collected from six sites denoted as sites 1 to 6 in garages around Ngara area in Nairobi-Kenya. The collected samples were kept in sterile falcon tubes prior to transportation to the laboratory and stored at -20°C.

2.2. Bacteria Isolation

Isolation of hydrocarbon degrading bacteria was carried out using enrichment technique [11]. Soil sample (1 g) from each site was transferred into 250 ml Erlenmeyer flasks containing 100 ml sterile Bushnell Haas media (BHM) (g l\(^{-1}\): MgSO\(_4\) (0.2), CaCl\(_2\) (0.02), KH\(_2\)PO\(_4\) (1.0), K\(_2\)HPO\(_4\) (1.0), NH\(_4\)NO\(_3\) (1.0), FeCl\(_3\) (0.05) final pH 7) [12] supplemented with 1% used engine oil as sole carbon source and incubated at 37°C for 7 days in a rotary shaker (Gallenkamp, England) operating at 120 revolutions per minute (rpm). An inoculum was then picked from each BHM flask and streaked on to Luria Bertani (LB) agar plates (g l\(^{-1}\): Tryptone (10.0), Yeast extract (5.0), NaCl (10.0), Agar (15.0) final pH 7.2) and incubated overnight at 37°C. Colonies were picked from each plate, transferred into test tubes containing LB broth and incubated in a rotary shaker at 200 rpm for 24 hours at 30°C. Serial dilutions of up to 10\(^{-9}\) from each test tube was done and an aliquot of 100 µl plated on LB agar plates. Discrete colonies (from 10\(^{-6}\) to 10\(^{-4}\) dilutions plates) were then picked and purified by plating on fresh LB agar plates and later LB broth before storage at 4°C.

2.3. Screening for Degradation

Pure isolates were screened for their ability to metabolize heating oil using the procedure described by [11]. Minimal media supplemented with heating oil as the sole carbon source was used. Bacterial cells previously cultured overnight in LB media were washed twice with 0.85% NaCl solution before suspending in the same solution. An aliquot (100 µl) of bacterial culture was then transferred to test tubes containing 5 ml minimal media following autoclaving for 15 minutes at 121°C. Heating oil (0.5 ml) was then added and incubation carried out at 37°C in a shaker at 150 rpm. Minimal media composition g l\(^{-1}\): (NH\(_4\))\(_2\)SO\(_4\) (1.0), MgSO\(_4\).7H\(_2\)O (0.1), KH\(_2\)PO\(_4\) (0.5), K\(_2\)HPO\(_4\) (0.76). Composition of ml l\(^{-1}\) trace elements solution (mg l\(^{-1}\): ZnSO\(_4\) (100), H\(_3\)BO\(_3\) (300), CaCl\(_2\).2H\(_2\)O (134.2), FeSO\(_4\).7H\(_2\)O (2000), CuCl\(_2\).2H\(_2\)O (10), NaMoO\(_4\).2H\(_2\)O (30), NiCl\(_2\).6H\(_2\)O (20), MnCl\(_2\).4H\(_2\)O (30). The pH was adjusted to 7 using 1 M NaOH. Bacterial isolates were also screened for their ability to degrade toluene, octane and hexane. Pure bacterial cultures were streaked on Bushnell Haas agar plates that were kept in a desiccator containing 10% toluene and 90% hexadecane in a 25 ml beaker and incubated at 25°C for 14 days. Isolates were also separately exposed to 20% hexane in 80% hexadecane and 20% octane in 80% hexadecane and treated as described for toluene. Un-inoculated plates were kept as control.

2.4. Morphological Identification

Bacterial isolates were examined by Gram’s staining test to differentiate between Gram positive and Gram negative bacteria using the procedure described by [13]. Potassium hydroxide test was carried out as a confirmatory test for Gram’s iodine staining applying the procedure described by [14].

2.5. Molecular Identification

Isolates that showed growth with the different
hydrocarbons were selected for identification through PCR amplification of 16S rDNA. DNA extraction was carried out using phenol/chloroform/isoamylalcohol according to the procedure described by [15]. The quality of genomic DNA obtained was then analysed on 1% (w/v) agarose gel in 1X TAE buffer.

An aliquot of genomic DNA extracted from each isolate was used as a template to amplify 16S rDNA gene. The gene was amplified using two universal primers (Eurofins genomics, Germany): 16S F27, forward 5’…AGA GTT TGA TCC TGG CTC AG…3’ and R1492, reverse 5’…GTT TAC CTT GGT TAC CTT…3’ [16]. The PCR reaction was performed in PCR reaction tubes (25 µl) using 12.5 µl OneTaq® Quick-Load® 2X master mix with standard buffer (New England Biolabs, United States), nuclease free water (9.5 µl), 10 µM forward primer (0.5 µl), 10 µM reverse primer (0.5 µl) and genomic DNA template (2 µl). Thermocycler (MJ research - PTC 200, USA) was used and the conditions for the PCR were as follows: An initial denaturation step at 95°C for 3 min followed by 30 cycles of 95°C for 30 sec, 52°C for 45 sec and 68°C for 45 sec and a final extension step at 68°C for 5 min followed by holding at 4°C. A positive control (E. coli 16S rDNA) and negative control (water) were included in gel electrophoresis of PCR products. The PCR amplicons were then purified using Qiaquick PCR purification kit (Qiagen, Valencia, USA) and sent to Eurofins Genomics Ebersberg Germany, for sequencing. Obtained sequences were analyzed and deposited in Genbank.

Obtained 16S rDNA sequences were compared with already known 16S rDNA sequences at National Center for Biotechnology Information (NCBI) database using BLAST (Basic Local Alignment Search Tool) algorithm. All the sequences were then aligned using CLUSTALW algorithm in Geneious 9.1.4 and Phylogenetic trees constructed based on the nucleotide sequences with the Bayesian phylogenetic method in MrBayes software. The trees were then visualized using Fig tree version 1.3.1 software.

2.6. Biochemical Tests (Carbohydrate Fermentation, Starch Hydrolysis and Catalase Tests)

Glucose, fructose, maltose, and sucrose fermentation tests were carried out using phenol red carbohydrate broth according to the method described by [17]. The procedure described by [18] was applied with some modification for starch hydrolysis test. In this test, pure bacterial colony from each isolate was streaked in a straight line on minimal media agar plate containing 2% soluble starch and incubated at 37°C for 48 hours. The catalase test was performed according to the procedure described by [19].

2.7. Optimization of Growth Conditions for Diesel Oil Degrading Bacteria

Optimization of growth conditions for three selected bacterial isolates was conducted using diesel oil according to the procedure described by [20] with some modifications.

2.7.1. Effect of pH on Bacterial Growth During Biodegradation of Diesel Oil

The effect of pH on growth of three bacterial strains was determined using 100 ml BHM supplemented with 1% diesel oil as the sole carbon source. Bacterial inoculums (100 µl) previously cultured overnight in LB media were inoculated in autoclaved BHM with pH values equating to 3, 5, 7, 9 and 11 following washing with 0.85% NaCl. The pH values were adjusted appropriately using 1 M NaOH and 1 M HCl. The test was conducted at 37°C for 7 days in a shaker with a speed of 150 rpm. Control tests containing no bacterial inoculum were also included. Bacterial growth was then monitored using spectrophotometer readings at 600 nm and net dry biomass (g l⁻¹) determined simultaneously. An aliquot (5 ml) of culture media was pipetted into a pre-weighed centrifuge tube and spun at 16000 x g for 10 min. This was then washed twice with distilled water and dried overnight at 90°C before reweighing and the difference in weight and the volume used considered to obtain dry biomass.

2.7.2. Effect of Temperature on Bacterial Growth During Biodegradation of Diesel Oil

The test was conducted using 1% substrate concentration at varying temperatures (25°C, 30°C, 37°C, 45°C and 55°C) alongside control tests. Bacterial growth was determined after culturing for 7 days using spectrophotometer readings at 600 nm and the dry biomass (g l⁻¹) also determined simultaneously as described in section 2.7.1.

2.7.3. Effect of Various Concentrations of Diesel Oil on Bacterial Growth During Biodegradation

The influence of substrate concentration on growth of the three bacterial isolates was determined using sterile BHM at pH 7 supplemented with various concentration of diesel oil i.e., 0.5, 1, 3, and 5% at 37°C. Bacterial growth was determined after culturing for 7 days in a shaker at 150 rpm. Growth was determined using spectrophotometer readings at 600 nm and the dry biomass (g l⁻¹) determined simultaneously as described in section 2.7.1.

2.7.4. Effect of Nitrogen Source on Bacterial Growth During Diesel Oil Biodegradation

The influence of nitrogen source on growth of the three bacterial isolates was determined using autoclaved nitrogen-limited minimal media (100 ml) supplemented with 1% diesel oil at pH 7 and 37°C. Ammonium nitrate (NH₄NO₃), yeast extract and tryptone [0.1% (w/v)] were separately used as nitrogen sources. Bacterial growth was determined after culturing for 7 days in a shaker at 150 rpm using spectrophotometer readings at 600 nm and the dry biomass (g l⁻¹) determined simultaneously as described in section 2.7.1.

Statistical analysis

All analyses were carried out in triplicate and the experimental data analyzed using one way Analysis of variance (ANOVA) using Statistical Package for Social Sciences (SPSS), version 19.0 [21,22]. Differences among mean values for treatments at P<0.05 were evaluated using
Post hoc test (Tukey’s test).

2.8. Hydrocarbon analysis using GC-MS

Diesel oil in BHM inoculated with isolates 1C, 2C, 3A and 4A2 were analyzed after 21 days of incubation. An aliquot (100 µl) of bacterial cells was transferred to a 250 ml volumetric flask containing 100 ml sterile BHM supplemented with 1% diesel oil. Un-inoculated BHM flask was kept as a control. After 21 days of incubation, the hydrocarbons were analyzed according to the procedure described by [23] with some modifications. Hydrocarbon extraction from 30 ml BHM was done using an equal volume of dichloromethane on a separating funnel. This was repeated twice to ensure complete recovery of the hydrocarbons. The dichloromethane phases were then combined and treated with anhydrous Na$_2$SO$_4$ to remove emulsions and residual water. The resultant extracts were concentrated by evaporation under a stream of Nitrogen using a heidolph rotary evaporator (Goel Scientific, India). The residue obtained from each sample was then re-dissolved in dichloromethane. Hydrocarbon composition was analyzed by GC-MS using Shimadzu QP2010SE series GC-MS (Shimadzu, Japan) equipped with Zebron GC column (ZB-1MS) [30.0 m by 0.25 mm (inner diameter)] with a thickness of 0.50 µm. Helium was used as the carrier gas and a temperature program consisting of an initial oven temperature of 55°C for 3 minutes increased to 245°C for 5 min at a rate of 4°C/min. An aliquot of 10 µl was used as the sample. The injector and detector temperatures were maintained at 250°C and 260°C respectively.

2.9. Alkane hydroxylase Gene Amplification

An aliquot (2 µl) of genomic DNA extracted from three selected isolates was used to amplify alkB gene. The gene was amplified using two primers (Eurofins genomics, Germany): alk-3F (5’…TCG AGC ACA TCC GCG GCC ACC A…3’) and alk-3R (5’…CCG TAG TGC TCG ACG TAG TT…3’) [3]. The expected PCR product was 330 bp. The process was performed in PCR tubes (25 µl) by adding OneTaq® Quick-Load® 2X master mix with standard buffer (12.5 µl), nuclease free water (9.5 µl), 10 µM forward primer (0.5 µl), 10 µM reverse primer (0.5 µl) and genomic DNA template (2 µl). Thermocycler conditions for the PCR were as follows: An initial denaturation step at 94°C for 2 min followed by 30 cycles of 94°C for 1 min, 54°C for 30 sec and 72°C for 30 sec and a final primer extension step at 72°C for 5 min and a final hold at 4°C. PCR products obtained were electrophoresed alongside GeneRuler 1kb DNA ladder (ThermoFisher Scientific, USA) on a 1% agarose gel with 1X TAE buffer and visualized under a UV light transilluminator before purification using Qiaquick PCR purification kit and sent to Eurofins Genomics Ebersberg, Germany for sequencing. Phylogenetic analysis was then carried out as previously described for 16S rDNA in section 2.5.

3. Results

3.1. Bacteria Isolation

Twenty bacterial isolates and a fungal isolate were obtained following isolation and purification. These were denoted as 1C, 2C, 3C, 4C, 5C, 5CB, 6C, 1B, 2B, 3B, 4B, 5B, 6B, 1A, 2A, 3A, 4A, 4A2, 5A, 6A and 3AF (fungi) based on the sampling locations. Six sites (1 – 6) were sampled at three different soil depths {surface (1 cm), sub-surface 1(5 cm) and sub-surface 2 (15 cm) denoted as C, B & A respectively}.

3.2. Screening for Biodegradation Potential

Following isolation, selection of efficient hydrocarbon degraders based on the ability to grow in minimal media supplemented with heating oil was done. All the isolates grew. Isolates 4A2, 3A, 5A, 6A, 1B, 1C, 2C, 5C and 6C showed steady increase in cell density compared to the rest of the isolates and were therefore selected for further studies (Figure 1). Screening for degradation of single hydrocarbons was carried out by exposing isolates cultured in BHA plates to toluene, hexane and octane vapors and the results are shown in Table 1.

[Figure 1. Time course of growth for bacterial isolates cultured in minimal media supplemented with 1% heating oil for 7 days.]
Table 1. Growth of isolates cultured in BHA plates exposed to toluene, hexane and octane hydrocarbons after 14 days of incubation at 25°C.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>10% Toluene + 90% Hexadecane</th>
<th>20% Hexane + 80% Hexadecane</th>
<th>20% Octane + 80% Hexadecane</th>
</tr>
</thead>
<tbody>
<tr>
<td>1C</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2C</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>5C</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>6C</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>1B</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>3A</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>4A2</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>5A</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>6A</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

+ denotes growth, - denotes no growth

3.3. Morphological and Molecular Identification of Isolates

Table 2 shows Gram’s stain and KOH test results of some of the selected isolates. Variation in morphological characteristics of the colonies suggests that the selected isolates were different from each other. Figure 2 shows a gel photograph of 16S rDNA PCR amplicons (≈ 1500 bp DNA fragments) of the nine selected bacterial isolates. A phylogenetic tree based on the BLAST search was constructed and the topological robustness of the tree evaluated using percentages of posterior probabilities as shown in Figure 3.

Table 2. Morphological characteristics of selected oil degrading bacteria cultured on LB agar plates.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Colony forms</th>
<th>Color of colonies</th>
<th>Margin</th>
<th>Gram's iodine staining</th>
<th>KOH Test</th>
<th>Shape of cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1C</td>
<td>Irregular</td>
<td>Pale brown</td>
<td>Undulate</td>
<td>-</td>
<td>+</td>
<td>Rods</td>
</tr>
<tr>
<td>2C</td>
<td>Circular</td>
<td>Cream white</td>
<td>Entire</td>
<td>-</td>
<td>+</td>
<td>Rods</td>
</tr>
<tr>
<td>5C</td>
<td>Irregular</td>
<td>Cream yellow</td>
<td>Lobate</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>6C</td>
<td>Circular</td>
<td>White</td>
<td>Curled</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>1B</td>
<td>Irregular</td>
<td>Cream white</td>
<td>Undulate</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>3A</td>
<td>Circular</td>
<td>Cream white</td>
<td>Undulate</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>4A2</td>
<td>Irregular</td>
<td>Yellow</td>
<td>Lobate</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>5A</td>
<td>Irregular</td>
<td>Cream</td>
<td>Undulate</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
<tr>
<td>6A</td>
<td>Irregular</td>
<td>Cream</td>
<td>Curled</td>
<td>-</td>
<td>+</td>
<td>Cocci</td>
</tr>
</tbody>
</table>

+ denotes positive test, - denotes negative test

The nine bacterial isolates clustered with members of the following genera; *Pseudomonas, Acinetobacter, Ochrobactrum, Salmonella, Enterobacter and Klebsiella*. Closest relatives to these isolates together with their percentage identity as obtained from the BLAST search at the NCBI database is as shown in Table 3. The sequences were deposited in NCBI Genbank and awarded accession numbers as indicated in the table.
Table 3. Closest relatives of selected bacteria based on 16S rDNA gene sequences.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Closest hit</th>
<th>Phylum: Proteobacteria</th>
<th>Accession number</th>
<th>Identity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1C</td>
<td><em>Pseudomonas aeruginosa</em></td>
<td>Gammaproteobacteria</td>
<td>KX036860</td>
<td>99</td>
</tr>
<tr>
<td>2C</td>
<td><em>Klebsiella variicola</em></td>
<td>Gammaproteobacteria</td>
<td>KX036863</td>
<td>99</td>
</tr>
<tr>
<td>5C</td>
<td><em>Enterobacter cloacae</em></td>
<td>Gammaproteobacteria</td>
<td>KX036856</td>
<td>99</td>
</tr>
<tr>
<td>6C</td>
<td><em>Klebsiella pneumoniae</em></td>
<td>Gammaproteobacteria</td>
<td>KX036858</td>
<td>99</td>
</tr>
<tr>
<td>1B</td>
<td><em>Enterobacter cloacae</em></td>
<td>Gammaproteobacteria</td>
<td>KX036855</td>
<td>99</td>
</tr>
<tr>
<td>3A</td>
<td><em>Acinetobacter baumannii</em></td>
<td>Gammaproteobacteria</td>
<td>KX036861</td>
<td>99</td>
</tr>
<tr>
<td>4A2</td>
<td><em>Ochrobactrum anthropi</em></td>
<td>Alphaproteobacteria</td>
<td>KX036859</td>
<td>99</td>
</tr>
<tr>
<td>5A</td>
<td><em>Enterobacter cloacae</em></td>
<td>Gammaproteobacteria</td>
<td>KX036862</td>
<td>99</td>
</tr>
<tr>
<td>6A</td>
<td><em>Salmonella enterica</em></td>
<td>Gammaproteobacteria</td>
<td>KX036857</td>
<td>99</td>
</tr>
</tbody>
</table>

3.4. Biochemical Characterization

A summary of the carbohydrate fermentation, catalase, and starch hydrolysis tests is given in Table 4 below.

Table 4. Biochemical characteristics of the nine selected bacterial isolates.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Carbohydrate fermentation test</th>
<th>Catalase test</th>
<th>Starch test</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Glucose</td>
<td>Fructose</td>
<td>Maltose</td>
</tr>
<tr>
<td>1C</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2C</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>5C</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6C</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>1B</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3A</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>4A2</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>5A</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6A</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

+ denotes positive test, - denotes negative test
3.5. Optimization of Growth Conditions for Diesel Oil Degrading Bacteria

There was significant difference in diesel oil degradation depicted by microbial growth at the different pH values \( (p<0.05) \). Post hoc test was carried out using Tukey’s test at a significance level of 0.05. The optimum pH for microbial growth was 7 (Figure 4). At this pH, maximum biomass obtained was 0.594 g l\(^{-1}\), 0.742 g l\(^{-1}\) and 0.609 g l\(^{-1}\) for isolate 3A, 1C and 2C respectively. Below and above pH 7, growth of the three isolates was reduced. Isolate 3A however, showed slight tolerance to alkaline pH compared to isolate 2C during the growth period.

![Figure 4. Effect of pH on growth of the three selected isolates. Bacterial growth expressed in biomass (g l\(^{-1}\)) after 7 days of incubation using BHM supplemented with 1% diesel oil. Error bars have been displayed using standard error of the mean. Microbial growth was optimal at pH 7.](image)

To determine the influence of temperature on diesel oil degradation by the three selected isolates, growth was carried out at temperatures ranging from 25°C to 55°C and at the predetermined optimum pH (Figure 5). There was a significant variation in microbial growth at 25°C, 37°C and 55°C \( (p<0.05) \). However, there was no significant difference in growth at 25°C and 45°C \( (P = 0.153) \) and at 30°C and 37°C \( (P = 0.515) \). From these results, it is evident that a temperature range of between 30°C and 37°C is suitable for growth of the three selected bacterial isolates (1C, 2C and 3A). Maximum biomass was obtained at 37°C.

![Figure 5. Effect of temperature on diesel oil degradation expressed in biomass (g l\(^{-1}\)) after 7 days of incubation at pH 7. Error bars have been displayed using the standard error of the mean. Optimum temperature for diesel oil biodegradation indicated by microbial growth was recorded at 37°C for all the three isolates.](image)

Microbial growth decreased with increase in diesel oil concentration as shown in Figure 6. There was a significant difference in bacterial growth at diesel oil concentrations of 0.5%, 1%, 3% and 5% \( (P = 0.000) \). Notably, maximum biomass at 0.5% diesel oil concentration was obtained earlier (on the 4\(^{th}\) day) during the culturing period compared to the rest of the increased diesel oil concentrations for all the three isolates. At 0.5% diesel oil concentration, maximum biomass obtained for isolate 3A, 1C and 2C were 0.757 g/L, 0.885 g/L and 0.7843 g/L respectively.
The effect of nitrogen source on diesel oil degradation by isolates 1C, 2C and 3A is presented in Figure 7. Ammonium nitrate, yeast extract and tryptone were separately used as nitrogen sources at the predetermined optimal pH and temperature. Significant variation in microbial growth was observed for the three nitrogen sources (p<0.05). From these results, microbial growth was highest in presence of yeast extract. With yeast extract, maximum biomass obtained was 1.344 g/L, 1.163 g/L and 0.972g/L for isolate 3A, 1C and 2C respectively.

### 3.6. Analysis of Diesel oil Degradation by Gas Chromatography-Mass Spectrometry (GC-MS)

The data obtained shows that the isolates were capable of readily mineralizing most of the hydrocarbons present in diesel oil. This is indicated by decrease in size of hydrocarbon peaks (low relative abundance) at certain retention times (Figure 8). Comparison of chromatogram profiles of the control and the different isolates indicate that most of the branched chain and cyclic alkanes as well as aromatic hydrocarbons were completely degraded as is indicated by absent peaks. In contrast, peaks for linear alkanes were still present though at reduced amplitudes. New peaks indicating formation of metabolic intermediates were also observed. Mono-aromatics (such as Benzene, 1,2,3-trimethyl- and 5-Ethyl-m-xylene) and polycyclic aromatic hydrocarbons (such as Naphthalene, 2-ethyldecacydro- and Naphthalene, 1,2,3,4-tetrahydro-2,5,8-trimethyl-) were completely degraded by all the four isolates. Most branched chain alkanes (such as Decane, 2-methyl- and Eicosane, 2,4-dimethyl-) were also completely depleted. With the exception of Cyclohexane, (4-methylpentyl)-, the isolates were also able to completely degrade cycloalkanes present in diesel oil. Conversely, almost all linear alkanes were still present at the end of the incubation period though at reduced levels. Isoprenoid hydrocarbons, phytane (hexadecane, 2,6,10,14-tetramethyl) and pristine (pentadecane, 2,6,10,14-tetramethyl) were also not completely depleted.
3.7. AlkB Gene PCR Amplification

AlkB (Alkane hydroxylase) gene from genomic DNA of isolate 1C was amplified and sequenced. A PCR amplicon of 313 bp was obtained. PCR amplification of the gene was however unsuccessful for isolates 2C and 3A. Obtained sequences were compared with already known alkB gene sequences at the NCBI database using BLAST algorithm. The sequence was deposited at the NCBI database under the accession number, KX036864. A phylogenetic tree was constructed using sequences from BLAST search and the isolate’s alkB sequence. The topological robustness of the tree was evaluated using percentages of posterior probabilities. Figure 9 shows the phylogenetic tree for alkB gene sequences constructed using MrBayes software.

Figure 8. GC-MS profiles of diesel oil extracted from BHM after 21 days of incubation at pH 7 and 37°C with and without inoculation. (A) Control (uninoculated); (B) Isolate 1C; (C) Isolate 2C; (D) Isolate 3A and (E) Isolate 4A2. Hydrocarbon and metabolite peaks were identified through comparison of retention times with mass spectrophotometer database using Autochro-3000 software. Diesel oil extraction was performed twice using dichloromethane (DCM) as the solvent.

Figure 9. Phylogenetic relationship of isolate 1C alkB gene (shown in red) with alkB gene of 14 closely related bacterial strains from NCBI database. The tree is rooted at mid-point. The numbers at the node show bootstrap values as percentages obtained with 1000 resampling analyses. Branch length unit (0.04) represents the number of substitutions per nucleotide site.
4. Discussion

Increased oil explorations in the East African region and the anticipated oil spillages that do occur during routine operations and transportation have raised concerns of environment pollution [24]. In the present study, hydrocarbon degraders were isolated from auto garage contaminated soils from Ngara region in Nairobi-Kenya. A total of 20 bacterial and 1 fungal isolates were obtained from the oil-contaminated soils sampled. Nine isolates were selected for further studies and these exhibited variable morphological characteristics as shown in Table 2.

4.1. Screening for Hydrocarbon Degradation

During culturing in heating oil, isolates 3A, 5A, 6A, 1B, 1C, 2C, 5C, 6C and 4A2 demonstrated high growth as indicated by increase in optical density readings at 600 nm. Post hoc analysis (Tukey) showed that there was no significant difference in microbial growth of all isolates up to day 2 (p=0.05) as evidenced in the growth curves (Figure 1). This was probably due to unavailability of enzymes (before induction) involved in mineralization of the complex hydrocarbons and/or minimal production of secondary metabolites important for microbial growth [3]. Significant growth (p<0.05) was later noted on subsequent days after which growth remained fairly constant after the 7th day. This stationary phase may be attributed to the need by the cell to adapt continuously to more complex hydrocarbons, depletion of degradable substrates or accumulation of waste products. For isolate 4A2, the curve indicated a diauxic growth. For this isolate, the first stage of growth was observed between the first and second day of incubation in which simple hydrocarbons such as n-alkanes and some alkylic chains are possibly degraded. A stationary phase was then observed between the 3rd and 5th day and later a second exponential phase noted after the 5th day. More complex hydrocarbon molecules such as mono and poly aromatics are thought to be degraded during this phase. The ability of the bacterial isolates to utilize single hydrocarbons (hexane, octane and toluene) as carbon sources was evaluated. Results (Table 1) showed that isolate 4A2 was able to utilize all the three hydrocarbons. Bacterial strains 1C, 2C, 6C, 3A and 4A2 on the other hand were able to utilize hexane while only 1C, 3A and 4A2 could grow on octane. Only isolate 4A2 could grow on plates exposed to toluene indicating its ability to withstand toluene toxicity. Toluene is an aromatic hydrocarbon of the BTEX compounds (Benzene, Toluene, Ethylbenzene & Xylene), which are classified as priority pollutants.

4.3. Selected Hydrocarbon Degraders

Isolate 1C, identified as Pseudomonas aeruginosa was capable of utilizing heating oil, hexane, octane as well as diesel oil as the sole carbon source. Members of the genus Pseudomonas have been identified as the most predominant group in metabolism of hydrocarbons [27]. Ability of Pseudomonas to efficiently take up alkanes has been linked to production of rhamnolipid biosurfactants as was demonstrated by [27] using P. aeruginos DSVP20. This class of bacteria possesses a broad array of physiological and metabolic properties as well as a complex enzymatic system that enable them to utilize a wide range of aliphatic and aromatic compounds as their sole carbon source [4,27]. The metabolic versatility of Pseudomonas has been linked to presence of degradative plasmids such as OCT (octane), ALK (alkanes), TOL (toluene) XYL (xylene) and NAH (naphthalene) [28].

Two of the isolates (2C and 6C) that demonstrated excellent growth when cultured in heating oil belonged to the genus Klebsiella. The two were also capable of utilizing hexane vapour as well as diesel oil as the sole carbon source. Klebsiella species are well established in degradation of petroleum compounds. Among 45 hydrocarbon degrading isolates obtained from estuary sediments, [29] reported that bacteria of the genus Klebsiella were the most frequently encountered (46.7%) with some of them recording over 90% attributed to their lipopolysaccharide membrane which can play the role of a biosurfactant accelerating the biodegradation process [25]. PCR amplification of 16S rDNA gene (Figure 2) as well as sequencing revealed that the nine isolates belong to six different genera. From the tree topology (Figure 3), isolates 1B, 5A and 5C clustered with members of the genus Enterobacter while 2C and 6C clustered with members of the genus Klebsiella. Isolate 1C was found to be a member of the genus Pseudomonas while 3A, 4A2 and 6A clustered with members of the genus Acinetobacter, Ochrobactrum and Salmonella respectively. All these strains fall under the phylum proteobacteria with Ochrobactrum being an alpha proteobacteria while the rest are gamma proteobacteria (Table 3) as depicted also by clustering in the phylogenetic tree. Bacterial strains of subphyla α-, β- and γ proteobacteria are well established in their ability to degrade a wide variety of hydrocarbons [16,25].

As expected, all the isolates in this study were found to possess catalase enzyme activity since they were isolated from oxygenated environment (soil) where they are required to neutralize toxic oxygen compounds (Table 4). This implies that the isolates are either aerobic or facultatively anaerobic [26]. Some of the isolates were capable of fermenting the different carbohydrates tested while some were found to be non-fermenters. Carbohydrate fermenting microbes are essential in the oil industry as some have been reported to produce biosurfactants which find applications in biodegradation of petroleum wastes as well as in oil recovery.

4.2. Microbial Identification and Characterization

Gram staining and potassium hydroxide tests revealed that majority of the isolates in this study were gram negative cocci-shaped with a few gram-negative rods (Table 2), a finding that is in agreement with other studies. Dominance of Gram negative bacteria in oil contaminated sites has been

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degradation of toluene, xylene, nonane and naphthalene.

In the current study, isolate 3A, identified as Acinetobacter baumannii was found to utilize heating oil, diesel oil, as well as hexane and octane vapors as the sole carbon source. In a study conducted by [25], Acinetobacter species was found to be the most abundant group. Efficiency of Acinetobacter sp. in utilization of hydrocarbons could be attributed to their ability to produce biosurfactants as was observed in a study conducted by [30]. In this study, A. radioresistens KA53 was reported to produce alasan which is a high-molecular-weight bioemulsifier complex observed to accelerate mineralization of recalcitrant polycyclic aromatic hydrocarbons (PAHs) [30].

4.4. Determination of Optimum Conditions for Growth Using Diesel Oil

In the present study, an optimum pH of 7 was observed for all the three selected isolates. Isolate 1C however, displayed higher biomass production (0.742 g l\(^{-1}\)) compared to the other two isolates (Figure 4). A number of studies have also indicated optimal growth at or near pH 7 [20,31,32]. Maintenance of an optimal pH condition is very vital as variation in pH of the culture media caused by accumulation of metabolic waste products affects microbial growth [32]. Notably, Isolate 3A was slightly tolerant to alkaline pH compared to isolate 2C. This could be linked to soil pH where the isolate was sampled. Isolate 2C was isolated from the surface compared to isolate 3A from the sub surface 2, much deeper than the former. It has been reported that for oil contaminated soils, as soil depth increases, the pH value also increases [31].

The three bacterial strains were observed to grow well at 30°C and 37°C (p<0.05). At 37°C however, a higher microbial biomass was noted for all the three strains with isolate 1C again recording the highest biomass (0.718 g l\(^{-1}\)) indicating greater diesel oil utilization (Figure 5). Above 37°C, growth was reduced. Though the optimum temperature was found to be 37°C, P. aeruginosa AT18 strain was found to efficiently assimilate n-alkanes, naphthalene, toluene and crude oil at 41°C in a study conducted by [28].

In the current study, optimization of diesel oil concentration revealed that 0.5% substrate concentration provided excellent growth for the three isolates (Figure 6). A suitable range of diesel oil concentration was observed between 0.5 - 1% although isolate 1C was able to tolerate up to 3% diesel oil concentration. This is similar to a study conducted by [32] in which Pseudomonas and Bacillus species were observed to tolerate increased diesel oil concentration. Thus an optimum range of substrate concentration is very vital since biodegradation is not easily stimulated below the oil concentration range while above the range, growth inhibition may occur due to oxygen limitations as well as solvent toxic effect.

Significant difference in microbial growth was observed for the three nitrogen sources with the addition of yeast extract giving the highest growth among the three isolates (Figure 7). This is in agreement with a study performed by [31] in which Pseudomonas aeruginosa, Pseudomonas putida, Aeromonas hydrophila and Acinetobacter Iwoffi were observed to grow optimally using yeast extract and tryptone as the nitrogen source. Although yeast extract was observed to provide excellent microbial growth as a nitrogen source in the present study, growth was also observed for ammonium nitrate as nitrate possess high oxidation potential for elimination of hydrocarbon contaminants which normally exist in a reduced state [12].

4.5. GC-MS Analysis of Diesel Oil Degradation

Compared to straight chain alkanes, most branched-chain and cyclic alkanes were totally degraded as indicated by the GC-MS analyses in Figure 8. This is contrary to some studies which have reported slower degradation of branched chain alkanes compared to linear alkanes. Higher preference for straight chain alkanes compared to branched chain alkanes for instance was reported by [33]. In the present study however, most linear alkanes were still present at the end of the incubation period possibly due to their high concentration in diesel oil. Comparison of GC-MS profiles of control media with that inoculated with different isolates also revealed complete microbial degradation of aromatic hydrocarbons in spite of their persistent and recalcitrant nature. PAHs show greater resistance to degradation and are classified as persistent organic pollutants (POPs) [34]. Additionally, the ability of the isolates to reduce acyclic isoprenoid hydrocarbons, phytane and pristine normally used as internal biomarkers in environmental hydrocarbon analyses, suggest that the microbes possess multiple degradative genes which facilitate a diverse catabolic ability. Similar to the present study, numerous studies have also reported that phytane and pristine degradation remains slow until most alkanes are removed mainly due to their persistent nature [35,36].

4.6. AlkB Gene PCR Amplification

Alkane hydroxylase gene of isolate 1C was successfully amplified and sequenced indicating the isolate’s potential catabolic capability in degrading alkane fraction of petroleum oil. Isolate 2C did not give the expected PCR product size probably due to nonspecific priming while 3A showed no PCR product for the gene. Lack of expected PCR products for this catabolic gene could be due to existence of completely different gene sequences from those characterized from the bacteria. Alternatively, regions used to develop the primer pair may not have been well conserved due to existence of gene homologues in this bacteria [37]. Phylogenetic analysis (Figure 9) indicated that the gene sequence for this isolate (1C) clustered with alkB gene from other P. aeruginosa strains with a posterior probability of 63% further supporting that isolate 1C is a Pseudomonas. AlkB gene catalyzes the first step in aerobic degradation of alkanes (medium and long chain) in which oxygen atom originating from molecular oxygen is introduced into the alkane substrate to form an alcohol.
5. Conclusion

A total of twenty one microbes (20 bacteria and 1 fungus) were isolated from oil-contaminated soils sampled from garages around Ngara, Nairobi, Kenya. From these, nine efficient bacterial isolates were identified based on morphological and biochemical tests as well as 16S rDNA sequence analyses. The isolates were observed to utilize heating oil, hexane, octane and toluene as well as diesel oil as the sole carbon source. Optimization of culture conditions using three of the most efficient degraders revealed that optimal degradation of diesel oil was recorded with <1% substrate concentration at pH 7 and temperature range of between 30°C and 37°C. Additionally, yeast extract was found to be the best nitrogen source for diesel oil biodegradation. GC-MS analyses demonstrated that the isolated bacterial strains were capable of readily degrading different alkane and aromatic hydrocarbons present in diesel oil thus exhibiting a broad range of catabolic activities. These findings clearly indicate the prospect to develop an environmentally friendly mitigation strategy against petroleum hydrocarbon pollution using the obtained bacterial isolates.

Acknowledgements

This research was supported by National Commission for Science, Technology and Innovation (NACOSTI) grant (NACOSTI/RCD/ST & I/7thCALL/Msc/ 072).

References


