Research Article

Isolation and identification of plasmids of bacteria from petroleum products contaminated soil

Abstract

Isolation and identification of plasmids of bacteria from petroleum products contaminated soil was carried out. Hydrocarbon utilizing bacteria were isolated from petroleum-products contaminated soil from the tank farm of NNPC/PPMC depot in Benin City, Nigeria. Plasmid analysis of bacteria was carried out showing the presence of plasmids in seven isolates: Bacillus subtilis, Bacillus species, Arthrobacter species, Pseudomonas aeruginosa, Klebsiella aerogenes, Alcaligenes species, and Pseudomonas species. All the plasmids correspond to 23kb of the maker. The result of the DNA extracted and amplified in a PCR using the ndoB primers indicated no visible bands, showing that ndoB primers is not specific for bacteria isolates in the samples, hence there was no amplification. The gene specification of ndoB gene of isolates does not imply the absence of plasmids DNA that confers utilizing ability to it, but that the utilizing gene could possibly be borne on the chromosomal DNA which enabled its persistence in the contaminated soil or other genes in which were tested such as catechol 2,3 dioxygenase (C23DO) and monooxygenase could be present.

Keywords: bacteria, plasmid DNA, petroleum products, contaminated soil

Introduction

Many important processes influence the destination of hydrocarbons in the environment. Among these are sorption, volatilization, abiotic transformation (chemical or photochemical), and biotransformation. Biodegradation of oil contaminated soils, which exploits the ability of microorganisms to degrade and/or detoxify organic contamination, has been established as one of the efficient, economic, versatile and environmentally sound treatment. The study of identification of bacteria is important in microbial ecology, especially with molecular techniques. In particular, analysis of the microbial communities that partake in in-situ microorganisms biodegradation activities has been a challenge to microbiologist. Interest in this area has been catalyzed by the rapid advancement of molecular ecological methodologies. Hydrocarbon utilizing microorganisms are ubiquitously distributed in the environment following oil spills. These microorganisms are naturally degraded numerous contaminating petroleum hydrocarbons and cleansing of oil pollutants. The ability of microorganisms to degrade an organic compound is the result ultimately of the genetic makeup of the organisms. The chemical reactions involved in metabolism are mediated by enzymes. The range of enzymes which a bacterium has is a reflection of the specific genetic information in the cell. Genetic information in bacteria, as in all organisms, is stored in the form of DNA. The information is physically present in bacterial cells in two forms the chromosome and the plasmids. The bacterial chromosome is a single circular, highly folded double-strand of DNA. In addition to chromosomal DNA, a larger number of bacteria also have extra-chromosomal DNA in the form of plasmids. Many plasmids contain genes which code for the enzymes necessary for the derivable pathways important to bioremediation. Enzymes involved in the degradation of toluene, naphthalene, salicylate, octane have been shown to be plasmid encoded. This research focused on the Isolation and Identification of Plasmids of Bacteria from Petroleum Products Contaminated Soil.

Materials and methods

Plasmid extraction

Plasmid extraction was done using the alkaline lysis method. This was carried out by labelling Eppendorf tubes according to number of isolates and filled with 1ml of sterile water. Bacterial growth was mix by vortexing. One and half (1.5) ml of the broth was taken into the labelled tubes and centrifuged at 13,000rpm for 2mins. The supernatant was discarded by decanting gently leaving a little of the broth. This was then vortexed at high speed until pellet is completely suspended in broth. Three hundred (300µl) of TENS solution and mix by inverting tubes until the solution becomes slimy. A hundred (100µl) of ice cold absolute ethanol was added, vortex and centrifuge at 13,000rpm for 10minutes. White pellet is observed after centrifuge at 13,000rpm for 10minutes. Hundred (900µl) of ice cold absolute ethanol was added, vortex and centrifuge at 13,000rpm for 10minutes. White pellet is observed after centrifuge at 13,000rpm for 10minutes. Nine hundred (900µl) of ice cold absolute ethanol was added, vortex and centrifuge at 13,000rpm for 10minutes.

DNA extraction

DNA was extracted using the boiling method which is an alternative method for isolation of DNA from both Gram negative and Gram positive organisms. In this procedure, bacterial cells were harvested into 1ml of sterile water, vortex and centrifuge at 10,000rpm for 5minutes. Again, it was washed with 1ml of sterile water. 200µl of sterile water was added, vortex and boil at 95-100°C for 10minutes. It was vortex again centrifuge at 10,000rpm for 5minutes. The supernatant was transferred into another tube and stored at -20°C and use for PCR.
**PCR analysis**

The primer used for the DNA amplification was ndob and 5X FIREPol master mix (Solis BioDyne). In this reaction mix, samples were labelled accordingly on top of the Eppendorf tube. Fourteen (14µl) of sterile de-ionized water was taken and 4µl of PCR buffer (5X FIREPol Master mix), 0.5µl of the primer (ndob) and 1.5µl of extracted DNA were added and vortexed at low speed. The centrifuge was pulsed and placed in the thermal cycler. The constituted reaction was denatured at 95°C for 3 minutes. Cycling began with denaturing at 95°C for 30 seconds, annealing at 50°C for 30 seconds and 72°C for 1 min; and a final 10-min extension step at 72°C. All reaction mixtures were held at 4°C until analyzed. Following optimization, all reactions were performed in repeated cycles for a total of 30 cycles in an Eppendorf thermocycler. All experiments included controls without any added DNA. This involved four basic steps cell harvest, lyses, deproteination and decontamination.

**Results and discussion**

While a total of nine hydrocarbon-utilizing bacteria were obtained as shown in Table 1. Again, the hydrocarbon-utilizing isolates were highest in PMS while that of pure DPK was lowest. Figure 1 shows bands of plasmid DNA for Bacillus subtilis, Micrococcus varians, Corynebacteria spicie, Bacillus spicie, and Arthrobacter spicie. The bands indicate that Bacillus subtilis, Micrococcus varians and Bacillus spicie. Plasmid DNA with molecular weights corresponding to 23130bp. While Figure 2 shows DNA plasmids in Pseudomonas aeruginosa, Klebsiella aerogenes, Alcaligenes specie and Pseudomonas spicie. Corresponding to 23kpb markers. Figure 3 shows bands of amplified DNA of isolates using the naphthalene dioxygenase (ndoB) primers. There was no visible band observed for the nine isolates, showing that the organism does not have an ndoB gene that coners utilizing ability. These findings are not unexpected, because there are numerous reports to show genetic diversity among degradative genes.

**Table 1 Hydrocarbon utilizing bacteria (cfu/ml)**

| Samples   | Viable cell count (cfu/ml) |
|-----------|----------------------------|
| Pure DPK  | 6.7±1.45x 103              |
| Mixed DPK | 1.16±0.12×104              |
| AGO       | 1.93±0.15×104              |
| PMS       | 4.27±0.27×104              |

Key: DPK, dual purpose kerosene; AGO, automobile oil and gas; PMS, premium motor spirit.

**Figure 1** Plasmid result of Gram positive bacteria: lane 1: Bacillus subtilis, lane 2: Micrococcus varians, lane 3: Corynebacteria spicie, lane 4: Bacillus specie and lane 5: Arthrobacter specie.

**Figure 2** Plasmid results for Gram negative bacteria: lane 1: Pseudomonas aeruginosa, lane 2: Klebsiella aerogenes, lane 3: Alcaligenes specie, lane 4: Pseudomonas spicie.

**Figure 3** Result of PCR with ndoB-specific primers. Lane 1: Bacillus subtilis; lane 2: Micrococcus varians, lane 3: Pseudomonas aeruginosa, lane 4: Klebsiella aerogenes, lane 5: Alcaligenes specie, lane 6: Corynebacterium specie, lane 7: Bacillus specie, lane 8: Arthrobacter specie and lane 9: Pseudomonas specie.
The plasmid profile of the isolates which we classified into Gram positive and Gram negative shows that there are plasmids in lane 1, 2 and 4 (Bacillus subtilis, Bacillus specie, and Arthrobacter specie.) which corresponds to 23Kbp of the marker. While the Gram negative indicate that lane 1, 2 and 3 and 4 (Pseudomonas species, Klebsiella aerogenes, Alcaligenes specie. and Pseudomonas specie) also shows there were plasmids that correspond to 23kbp of the standard. PCR of the genomic DNA of all the nine isolates were negative to ndoB primers as can be seen on Figure 3. These findings are not unexpected, because there are numerous reports to show genetic diversity among degradative genes. Among 20 PAH-degrading isolates, strong hybridization signals with ndoB were only observed in seven of eight naphthalene-degrading Pseudomonas strains. In 1989, Furukawa found that only five Pseudomonas species of 15 biphenyl-degrading Pseudomonas, Achromobacter, Alcaligenes, Moraxella and Arthrobacter strains showed significant hybridization to phABC of Pseudomonas pseudoalcaligenes KF 707. Also observed no visible band for cuts made with EcoRV restriction enzyme for Klebsiella. From the foregoing, we cannot conclude that these bacteria do not have utilizing genes or that the degradation is plasmid mediated. But that other primers which were not available in the laboratory as of this work could detect their degradative genes.

Conclusion

Plasmid analysis of bacteria isolated show molecular sizes of the plasmid DNA molecules. No visible band was observed for the isolated bacteria indicating the organisms lack plasmid DNA that could be borne on the chromosomal DNA which enabled its persistence in the polluted soil. The presence of Pseudomonas species in all the hydrocarbon contaminated soil revealed its crude oil utilizing potential and hence should be considered the first choice in bioremediation of hydrocarbon polluted sites.

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None.

Conflict of interest

The author declares no conflict of interest.

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