Molecular Characterization of Low-Density Polyethene (LDPE) Degrading Bacteria and Fungi from Dandora Dumpsite, Nairobi, Kenya

Christabel Ndahebwa Muhonja,1 Gabriel Magoma,1 Mabel Imbuga,2 and Huxley Mae Makonde3

1Pan African University of Science and Technology, P.O. Box 62000-00200, Nairobi, Kenya
2Jomo Kenyatta University of Agriculture and Technology, P. O. Box 62000-00200, Nairobi, Kenya
3Department of Pure & Applied Sciences, Technical University of Mombasa, P. O. Box 90420-80100, Mombasa, Kenya

Correspondence should be addressed to Christabel Ndahebwa Muhonja; muhosh@gmail.com

Received 26 July 2018; Revised 22 September 2018; Accepted 2 October 2018; Published 3 December 2018

1. Introduction

Low-density polyethylene is a major cause of environmental pollution due to its high tensile strength, lightness, resistance to water, and microbial attack. The consumption of plastics in the country has increased to 4,000 tons per annum of polyethylene bags which together with hard plastics end up scattered in the environment creating “the plastics menace” [1]. Through the National Environmental Management Authority (NEMA), Kenya has embraced the 3Rs, reduce, reuse, and recycle, concept of solid waste management [2] and most recently the ban on the use of polyethylene carrier bags but this has not addressed the problem of polyethylene which remains scattered in the environment [3].

Biodegradation is the decomposition of substances through microbial activity and is a complex process which involves the following steps [4]: biodeterioration, depolymerization, assimilation, and mineralization. Bacteria and fungi of various genera have been implicated previously in the biodegradation of polyethylene albeit the low rates. Acinetobacter sp. was found capable of utilizing n-alkanes of chain length C10–C40 as a sole source of carbon as reported by [5]. Bacterial genera, namely, Pseudomonas, Acinetobacter, Brevibacillus, Rhodococcus, and Micrococcus [6, 7, 1], respectively, isolated from different sources proved to be the
potential organisms for polyethylene degradation. Fungal genera, Gliocladium, Cunninghamella, Penicillium, Aspergillus, Fusarium, Mucor, and Mortierella, from soil [1] were proven to have the potential to degrade polyethylene after analysis of degradation through various methods.

Plastic biodegradation as a result of the activity of certain enzymes causes cleavage of the polymer chains into monomers and oligomers. Enzymatically broken down plastic is further absorbed by the microbial cell to be metabolized. Aerobic breakdown produces carbon dioxide and water. The involvement of enzymes in microbial biodegradation of polyethylene has been investigated, and enzymes such as laccases and esterases have been confirmed to play a role in this process either directly or indirectly [8]. The production of enzyme laccase in the presence of polyethylene as the sole carbon source is a clear indication that laccase has a role in breaking down some of the intermediary products produced during this process. In this study, molecular characterization of bacteria and fungi that had been confirmed to degrade polyethylene was done as well as assessment of optimum pH, temperature, and sodium chloride concentration at which they can thrive. Presence of AlkB genes as the sole carbon source is a clear indication that laccase has a role in this process either directly or indirectly [8]. The PCR was performed on a GeneAmp PCR system 9600 (Applied Biosystems), using 1 µl Taq Polymerase (Applied Biosystems), 1 µl each of 10 PM concentrations of forward and reverse primers, 27 µl sterile deionized water, 8 µl PCR buffer containing dNTPs and MgCl₂, and 2 µl DNA template, for a total reaction volume of 40 µl. The cycling program used was as follows: 1 cycle of 94°C for 5 min; 30 cycles of 94°C for 30 sec, 55°C for 30 sec, and 72°C for 1.5 min; and a final extension of 72°C for 10 min. The PCR products were visualized through electrophoresis on a 1% agarose gel with ethidium bromide added directly. The 1.5 kbp products were subjected to Sangerideoxy sequencing using the forward primer and reverse primers at Macrogen DNA, Inc. (Netherlands). Sequence files were edited using Chromas version 2.6.2 and compared to the GenBank nucleotide database using the basic local alignment search tool (BLAST). Phylogenetic relationships were inferred using Mega 7 [11], and maximum-likelihood algorithms were available in Phylib. Maximum likelihood and parsimony-derived trees were bootstrapped using PHYML [12, 13].

2. Materials and Methods

2.1. Bacterial DNA Extraction. Total genomic DNA was isolated from the bacterial pure cultures grown to the late exponential phase by means of a standard protocol [9] as follows: 1.5 ml of the overnight bacterial culture (grown in the LB medium) was transferred to a 1.5 ml Eppendorf tube and centrifuged at 13000 rpm for 1 min to pellet the cells. The supernatant was discarded. The cell pellet was suspended in 600 µl TE buffer and centrifuged at 13000 rpm and the supernatant discarded. The cell pellet was resuspended in 200 µl TE buffer, and the following were added: 5 µl lysozyme (20 mg/ml), 5 µl RNase A (20 mg/ml), and 10 µl proteinase K (20 mg/ml) followed by overnight incubation at 37°C. In the next morning, the temperature was adjusted to 56°C for one hr and an equal volume of phenol/chloroform (1:1) was added and mixed well by inverting the tube until the phases were completely mixed. Spinning was done at 13000 rpm for 15 min at room temperature. The upper aqueous phase was carefully transferred to a new tube by using 1 ml pipette. This step was repeated twice to ensure all protein had been removed. An equal volume of chloroform and isoamyl (24:1) was added to the aqueous layer and centrifuged at 13000 rpm for 15 min. The aqueous layer was removed into a new tube. This step was also repeated to ensure all phenol is removed. An equal volume of isopropanol was added and stored overnight at −20°C. The samples were then defrosted and centrifuged at 4°C for 30 min to pellet the DNA. The pellet was washed in 70% ethanol and centrifuged at 13000 rpm for 5 min, and then, the ethanol was carefully pipetted out. The pellet was air dried on the bench for 20 min, and the isolated genomic DNA was viewed on a 1% agarose gel.

2.2. Bacterial DNA Amplification and Sequencing. Amplification of the 5′ end of the 16S rDNA gene was performed with universal primers (forward primer (8-F) 5′-AGAGTTTGATYMTGGCTCAG-3′) and reverse primer ((1942R) 5′-GTTACCTTGATACGGACTT-3′) [10]. The PCR was performed on a GeneAmp PCR system 9600 (Applied Biosystems), using 1 µl Taq Polymerase (Applied Biosystems), 1 µl each of 10 PM concentrations of forward and reverse primers, 27 µl sterile deionized water, 8 µl PCR buffer containing dNTPs and MgCl₂, and 2 µl DNA template, for a total reaction volume of 40 µl. The cycling program used was as follows: 1 cycle of 94°C for 5 min; 35 cycles of 94°C for 30 sec, 55°C for 30 sec, and 72°C for 1.5 min; and a final extension of 72°C for 10 min. The PCR products were visualized through electrophoresis on a 1% agarose gel with ethidium bromide added directly. The 1.5 kbp products were subjected to Sangerideoxy sequencing using the forward primer and reverse primers at Macrogen DNA, Inc. (Netherlands). Sequence files were edited using Chromas version 2.6.2 and compared to the GenBank nucleotide database using the basic local alignment search tool (BLAST). Phylogenetic relationships were inferred using Mega 7 [11], and maximum-likelihood algorithms were available in Phylib. Maximum likelihood and parsimony-derived trees were bootstrapped using PHYML [12, 13].

2.3. Fungal DNA Extraction. Fungal DNA extraction protocol reported by Gontia-mishra et al. [14] was used. Fungal mycelia were grown for 7 days at 55°C on potato dextrose agar. Mycelia were frozen in liquid nitrogen and ground to a fine powder using a mortar and pestle. The powder was transferred into 2 ml tubes, and 600 µl of preheated extraction buffer was added. The contents were incubated in a water bath at 65°C for 30 minutes with mixing after every 10 minutes. 270 µl volume of 5 M potassium acetate was added and centrifuged at 13000 rpm for 10 minutes. 700 µl of the supernatant was transferred into clean tubes, and 5 µl RNASE (10 mg) was added and then incubated for 30 minutes at 37°C. Chloroform and iso-amyl alcohol was prepared in the ratio of 24:1, and an equal volume was added to the mixture. 600 µl of supernatant was pipetted into clean tubes. DNA was precipitated by adding a tenth of the volume of 3 M potassium acetate and two thirds of the volume of isopropanol. This was incubated at −20°C for 30 minutes then centrifuged at 13000 rpm for 10 minutes. The pellet was washed using 70% ethanol followed by 10 minutes of centrifuging, and then, the DNA was eluted in 50 µl of RNASE-free water and stored at −20°C.

2.4. Fungal DNA Amplification and Sequencing. PCR was performed on a GeneAmp PCR system 9600 (Applied Biosystems), using 1 µl Taq Polymerase (Applied Biosystems), 1 µl each of 10 PM of forward and reverse primers, 27 µl deionized water, 8 µl PCR buffer containing dNTPs and MgCl₂, and 2 µl DNA template, for a total reaction volume of 40 µl. The cycling program used was as follows: 1 cycle of 95°C for 5 min; 35 cycles of 95°C for 30 sec, 60°C for 45 sec, and 72°C for 40 sec; and a final extension of 72°C for 5 min.
Primer pair F-566-5′-CAGCAGC CGCGTAAATTCC-3′ and for R- 1200-3′-CCGTGTTGAGTCA AATTAGC-3′ which amplify on average a 650 bp long fragment from the V4 and V5 regions were used [15]. The PCR products were visualized through electrophoresis on a 1% agarose gel with ethidium bromide added directly. The products were subjected to Sanger dyeoxy sequencing by Macrogen, Inc. (Netherlands). SeqMan Pro was used to assemble both the forward and reverse sequence files [16]. The sequences obtained were compared against the sequences available in the NCBI database using the basic local alignment tool (BLAST). The 18S rDNA gene sequences obtained in current study, together with those of the closest neighbor strains, were aligned using ClustaX version 2.1. Phylogenetic relationships were inferred using Mega 7 [11], and maximum-likelihood algorithms were available in Phylip. Maximum likelihood and parsimony-derived trees were bootstrapped using PHYML [12, 13].

2.5. Screening for Production of Enzymes. Bacterial isolates were screened for their ability to produce extracellular enzymes, i.e., laccases and esterases. The ability of the isolates to utilize substrates such as lignin and tween 20 exhibited their ability to produce the respective enzymes [17].

2.5.1. Determination of Presence of Enzyme Laccase. The media for selection of lignin-modifying fungi were prepared by the use of plain agar and minimal salt media with the incorporation of lignin (to encourage selection of ligninolytic fungi) and Guaiacol, which acts as a colorimetric indicator of the lignin-modifying enzymes laccase or peroxidases. All chemicals were obtained from Sigma Chemical Co., St. Louis. The presence of a reddish coloration after 3–5 days of incubation was an indication of laccase activity. The laccase assay per 1 liter: 400 µl Guaiacol, agar 15 g, 2 g malt extract, 0.5 g KH₂PO₄, 0.001 g ZnSO₄, 0.4 g K₂HPO₄, 0.02 g FeSO₄, and 0.2 g MgSO₄, 0.5 g KH₂PO₄, 0.1 g NH₄NO₃, 0.1 g KCl, 5 ml KOH, 0.25 g chloramphenicol, forming a reddish colored zone as a positive result.

2.5.2. Determination of Presence of Enzyme Esterase. The isolates were cultured on basal media (1% KH₂PO₄, 0.01% MgSO₄·7H₂O, 0.005% CaCl₂·2H₂O, 4% NaCl, and 1% Na₂CO₃) supplemented with 1% tween 20 (domestic grade) as the sole carbon source. The medium was then thereafter inoculated by the spotting of isolates per plate and incubated for at least 48 hours at 37°C for bacteria and at 28°C for 3–5 days for fungal isolates. The media were observed for zones of precipitation of calcium crystals around each isolate. Positive isolates for esterase production were indicated by the precipitation of calcium crystals around the colonies.

2.5.3. Screening for Genes Producing Alkane-Degrading Enzymes. Amplification was done using the sets of AlkB primers [18] shown in Table 1. The PCR was performed on a GeneAmp PCR system 9600 (Applied Biosystems) using Taq DNA polymerase. A total of 30 cycles of amplification was performed with template DNA denaturation at 94°C for 1 min, primer annealing at 40°C for 1 min, and primer extension at 72°C for 2 min [19]. The PCR products were visualized through electrophoresis on a 1% agarose gel with 2 µl ethidium bromide added directly.

2.6. Effect of Temperature on Growth of Fungal Isolates. Potato dextrose agar augmented with 250 mg/ml ampicillin to inhibit bacterial growth at pH 7.0 was prepared, sterilized, and dispensed in sterile Petri dishes. Each plate was inoculated with one fungal isolate and incubated at temperatures 20, 30, and 40°C. Growth of isolates was checked after 4 days of incubation. The level of growth was scored using the colony diameter, whereby 0 mm indicated no growth, 1-2 mm indicated minimal growth, 3-4 mm indicated average growth, and 5-7 mm indicated satisfactory growth while 8-10 mm indicated excellent growth.

3. Results and Discussion

3.1. Phylogenetic Relatedness of Low-Density Polyethylene Degrading Fungal Isolates. Amplification of fungal 18S rDNA using 1200R and 566F universal primers yielded the expected band size of approximately 640 bps from the PCR products of the amplified samples (Figure 1). These products were purified, sequenced, and analyzed. The results were used to obtain accession numbers from the NCBI GenBank. The analyzed sequences were aligned with those of the closest neighbors using ClustaX version 2.1. Phylogenetic relationships were inferred from phylogenetic comparison of the 18S rDNA sequences using Mega 7 and maximum-likelihood algorithms to generate the phylogenetic tree (Figure 2) which shows the phylogenetic relationships among the various Aspergillus species. The tree displays four clades in which the isolates have been clustered. From our previous study [20], Aspergillus oryzae (MG779508) resulted in a weight loss of 36.4 ± 5.53% which was the highest. Aspergillus oryzae is a promising biodegrader of polyethylene as it was able to degrade 30% of polyethylene in 200 days [21] in addition to formation of microcracks and increased embrittlement of the LDPE surface upon SEM analysis. In a study done using untreated LDPE incubated with A. oryzae, 5% weight loss was recorded compared with control (untreated and unexposed), Aspergillus fumigatus strain B2,2 (MG779513) recorded a weight reduction of 24 ± 3.26% which was the second highest in our previous study [20]. Aspergillus fumigatus is also among the species that have been investigated for their ability to degrade polyethylene and other polymers. In a study, three fungal species were investigated for their ability to degrade polyethylene, and A. fumigatus was the best degrader compared to A. terreus and F. solani following an analysis of the LDPE surface by SEM and FTIR [22]. Other fungi implicated in this study included Aspergillus nidulans, A. flavus, A. terreus, and A. neoflavipes which resulted in weight loss of the LDPE sheets. Use of weight reduction as a measure of the extent of polyethylene biodegradation has been widely accepted and used by many authors [23].
**Table 1:** Primers for AlkB genes encoding depolymerases responsible for alkane degradation [18].

| Primers and position | PCR product | Reference |
|----------------------|-------------|-----------|
| **alkB 1 set 1**     |             |           |
| 82 5'-TGGCCGGCTACTCGATGATCGGAATCTGG-3' 111  
951 5'-CGCGTGTTGATCGAGTGCGGTGGAGAGTTG3'-922 | 870 bp | Kok et al. |
| **alkB 1 set 2**     |             |           |
| 134 5'-CATTTCCCTGGATGTGATTG-3' 151  
851 5'-CCTCCTCGCCCTTTCGC-3' 834 | 718 bp | Stover et al. |
| **alkB 2**           |             |           |
| 134 5'-CTGGCTGGATCGAGTGCGGTGGAGAGTTG3'-151  
882 5'-CGAGTGTCTGCCGCGGTGG-3' 864 | 749 bp | Stover et al. |

**Figure 1:** PCR products for the amplification of 18S rDNA for the fungal isolates 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, and 16 using 1200R and 566F universal primers. L represents a 1kb ladder. The expected band size amplified is 640bps.

**Figure 2:** Phylogenetic tree of fungal isolates based on 18S rDNA sequences. All screened fungal isolates have NCBI accession codes in brackets. The scale bar refers to 0.007 substitutions per nucleotide position. Bootstrap values obtained with 1000 resampling are referred to as percentages at all branches.
3.2. Phylogenetic Relatedness of Low-Density Polyethylene Degrading Bacterial Isolates. Amplification of bacterial 16S rDNA using 1492R and 8F universal primers yielded 1420 bps fragments (Figure 3) which were purified, sequenced, and analyzed. The results were used to obtain accession numbers from the NCBI GenBank. The analyzed sequences were aligned with those of the closest neighbors using ClustaX version 2.1. Phylogenetic relationships were inferred from phylogenetic comparison of the 16S rDNA sequences using Mega 7 and maximum-likelihood algorithms to generate the phylogenetic tree (Figure 4) which shows the phylogenetic relationships among the genera and species. Brevibacillus, Bacillus, and Lysinibacillus are in one major clade while Pseudomonas, Ochrobactrum, and Celulosimicrobium are grouped in another major clade. Bacteria of the genera Bacillus, Brevibacillus, Ochrobactrum, Lysinibacillus, Celulosimicrobium, and Pseudomonas were identified as effective polyethylene degraders. Bacterial isolates A5,1a-Bacillus cereus (MG645256) produced the highest degradation effectiveness in terms of weight loss, i.e., 35.2%, followed by the isolate B2,2-Brevibacillus borstelensis (MG645267) showing 20.28% from an earlier study [20] while isolates B1,1a-Pseudomonas putida (MG645283) and D4,yn-Brevibacillus borstelensis strain (MG645261) produced a weight loss of 2.88% and −6.8%, respectively. The genus Bacillus was most frequently identified among the LDPE biodegrading genera in this study. Species identified under this genus include Bacillus cereus, Bacillus toyonensis, Bacillus thuringiensis, Bacillus subtilis, Bacillus pseudomycoides, Bacillus safensis, and Bacillus naciens. Various studies have been done to investigate the efficacy of genus Bacillus in polyethylene degradation, and different species under this genus have been found to have potential to degrade polyethylene [24, 25]. Bacillus cereus has been found to be a good bioremediation candidate in the biodegradation of polyethylene due to its ability to produce enzymes laccase and manganese peroxidase. In a comparative study, B. cereus was found to be more effective than B. sphericus in degrading photo-oxidized and thermos-oxidized LDPE [26]. According to [6], Brevibacillus borstelensis Accession number AH764129 was able to degrade 11% of nonirradiated polyethylene by weight in 30 days. Two bacterial isolates Bacillus amyloliquefaciens (BSM-1) and Bacillus amyloliquefaciens (BSM-2) were isolated from municipal soil and used for polymer degradation studies and were found to produce significant changes on LDPE in terms of weight loss, reduction of tensile strength, and appearance of new functional groups [27]. A novel strain of Pseudomonas, Pseudomonas citronellolis EMBS027, GenBank Accession number KF361478 was isolated by [28] from a municipal landfill in Indore, India, and it degraded 17.8% of polyethylene in 4 days. Different species of Pseudomonas were analyzed for their ability to degrade polyethylene and upon incubation for 120 days. Pseudomonas putida resulted in a weight loss of 9% [29].

3.3. Screening for Enzyme Production. Bacterial isolates were screened for production of enzymes laccase and esterase (Figure 5) which are among the enzymes implicated in LDPE degradation. Bacterial isolates Brevibacillus borstelensis strain B2,2, Brevibacillus parabrevis strain C2,2a, and Pseudomonas putida strain B1,1 exhibited the highest presence of laccase. Only two isolates Bacillus toyonensis and Bacillus macrolides were negative for laccase activity. Esterase activity was highest in isolates Brevibacillus borstelensis strain D4 yn, Bacillus naciens strain C4,1a, and Pseudomonas putida strain B1,1a. Fungal isolates were screened for production of enzymes laccase and esterase (Figure 6). Isolates B2, 2-Aspergillus fumigatus, A5,1-Aspergillus oryzae, and A4,2a-Aspergillus flavus exhibited the highest levels of laccase enzyme while the highest level of esterase enzyme was attributed to fungi Aspergillus Oryzae.

Production of extracellular enzymes plays an important role in polymer degradation through depolymerization, where the polymer is broken down into smaller subunits [30] which are then enzymatically degraded into intermediary products that can be assimilated into microbial cells [31] and utilized as carbon sources leading to production of energy, water, carbon dioxide, and methane in the case of anaerobic respiration [32]. In this study, production of extracellular enzymes, laccase and esterase, were investigated. Fungal and bacterial isolates in this study were scrutinized for their ability to produce laccase enzyme and isolate B2, 2-Aspergillus fumigatus (MG779513) which had a LDPE degradation effectiveness of 24% and had the highest diameter of coloration due to laccase production. This could be attributed to its ability to produce higher amounts of laccase and other extracellular enzymes which are believed to play a role in polyethylene degradation. According to [33], the production of this enzyme increases when the microbes are in close proximity with the polyethylene. Reference [34] was able to extract the crude laccase enzyme which was incubated with polyethylene and led to degradation as was evidenced through weight loss, FTIR, and SEM. Esterases catalyze the cleavage of ester bonds [35] of short-chain triglycerides or esters. Esters have been identified as part of the intermediary products produced during polyethylene degradation when the postincubation culture media are subjected to GC-MS analysis that can be assimilated into the microbial cells, undergoes hydrolysis to give rise to the subsequent carboxylic acid and alcohol that ultimately undergo respiration to produce energy [36]. Isolate A5,1-Aspergillus oryzae (MG779508) with a weight loss of 36.4% had a high activity of enzyme esterase of 10%. This could have contributed to its high degradation potential compared to other fungal isolates which had lower degradation.
3.4. Screening for AlkB Genes Producing Alkane-Degrading Enzymes. PCR to amplify AlkB genes was done using three sets of AlkB primers. Only one set of the AlkB primer was able to amplify the AlkB gene producing a fragment of size 870 bps. AlkB genes are responsible for production hydrolyase enzymes which are responsible for alkane degradation. The gene was amplified in 4 bacterial samples (Figure 7). A common feature of many alkane degraders is that they contain multiple alkane hydroxylases with overlapping substrate ranges [37]. AlkB- and alkB-related genes code for an alkane-degrading enzyme, alkane hydroxylase [38]. The analysis of the bacterial samples revealed presence of AlkB.
gene in 4 of the bacterial samples. Bacterial isolates that were positive for alkB 1 gene were D4 yn-
Brevibacillus borstelensis, B1,1-
Pseudomonas putida, and A5,a1-
Bacillus cereus. Alkane biodegradation is initiated through terminal oxidation to the corresponding primary alcohol, which is further oxidized by dehydrogenases to fatty acids which can enter the TCA cycle [39]. This genetic information is an indication of the genetic ability of the microorganisms to degrade long-chain alkanes through production of this enzyme.

3.5. Effect of Temperature on Growth of Fungal Isolates. The growth of fungi at different temperatures (20°C, 30°C, and 40°C) as shown (Figure 8) revealed that growth at 30°C
was significantly higher than growth at 20°C and 40°C with A. oryzae strain A5,1 having the highest growth (10 ± 0). However, isolate E4,1-A. nidulans grew optimally at 40°C. This could be attributed to the fact that the sampling site for these bacteria was from a dumpsite where the temperatures were generally ambient and hence favoring the growth of mesophilic microbes. Laccase production by fungi is influenced by type and concentration of carbon sources, pH, and temperature.

### 4. Conclusion

The present work indicates that soil bacteria and fungi isolated from the dumpsite have potential of degrading polyethylene. This is the first study on the isolation of local bacteria and fungi that can degrade LDPE which is the most common plastic in Kenya. Particularly, the application of Aspergillus oryzae strain A5,1 and Bacillus cereus strain A5,1 will be beneficial in the bioremediation of polyethylene as they exhibited significant degradation effectiveness. It was ascertained that the microorganisms are capable of producing enzymes laccase and esterase which have been confirmed to play a role in degradation of polyethylene. The isolates possess the alkane hydroxylase-producing gene (AlkB) which is the molecular explanation for the degradation of LDPE under investigation. Fungi in this study were found to grow optimally at the temperature of 30°C.

### Data Availability

The data (Figures 1–8 and Table 1) used to support the findings of this study are included within the article.

### Conflicts of Interest

The authors declare that they have no conflicts of interest.

### Acknowledgments

This work has been fully funded by the African Union Commission through the Pan African University Institute of Science, Technology, and Innovation. We highly appreciate the support we got from honest colleagues, Dr. Johnstone Neondo, Dr. Eliud Wafula, Steve Ogada, and Baba Ngom, who gave us sincere and constructive criticism that helped to shape up this final product. We thank the staff and management of the Institute of Biotechnology Research (IBR), Prof. A. B. Nyende, Richard Rotich, Catherine Eyiinda, and Grace Mungai, for providing an enabling environment for laboratory and research work for this study.
References

[1] B. Nowak, J. Pajak, M. Drozd-Bratkowicz, and G. Rymarz, "Microorganisms participating in the biodegradation of modified poly-ethene films in different soils under laboratory conditions," International Biodeterioration and Biodegradation, vol. 65, no. 6, pp. 757–767, 2011.

[2] M. C. Aurah, "Assessment of extent to which plastic bag waste management methods used in Nairobi city promote sustainability," American Journal of Environmental Protection, vol. 1, no. 4, pp. 96–101, 2013.

[3] J. P. Eubeler, M. Bernhard, S. Zok, T. P. Knepper, and T. P. Knepper, "Environmental biodegradation of synthetic polymers I. Test methodologies and procedures," Trends in Analytical Chemistry, vol. 28, no. 9, pp. 1057–1072, 2009.

[4] J. D. Gu, "Microbiological deterioration and degradation of synthetic polymeric materials: recent research advances," International Biodeterioration and Biodegradation, vol. 52, no. 2, pp. 69–91, 2003.

[5] R. Pramilla and V. Ramesh, "Potential biodegradation of low density poly-ethylene (LDPE) by Acinetobacter baumannii," African Journal of Bacteriology Research, vol. 3, no. 1, pp. 92–95, 2015.

[6] D. Hadad, S. Geresh, and A. Sivan, "Biodegradation of poly-ethylene by the thermophilic bacterium Brevibacillus borstelensis," Journal of Applied Microbiology, vol. 98, no. 5, pp. 1093–1100, 2005.

[7] S. Nanda and S. S. Sahu, "Biodegradability of poly-ethene by Brevibacillus, Pseudomonas, and rhodococcus spp.," New York Science Journal, vol. 3, no. 7, pp. 95–98, 2010.

[8] N. Lucas, C. Bienaimé, C. Belloy, M. Queneucde, F. Silvestre, and J. Nava-saucedo, "Chemosphere Polymer biodegradation: mechanisms and estimation techniques," Chemosphere, vol. 73, no. 4, pp. 429–442, 2008.

[9] F. M. Ausubel, R. Brent, R. E. Kingston et al., "Current protocols in molecular biology," Molecular Biology, vol. 1, p. 146, 2003.

[10] W. G. Weisburg, S. M. Barns, D. A. Pelletier, and D. J. Lane, "16S ribosomal DNA amplification for phylogenetic study," Journal of Bacteriology, vol. 173, no. 2, pp. 697–703, 1991.

[11] S. Kumar, G. Stecher, and K. Tamura, "MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets," Molecular Biology and Evolution, vol. 33, no. 7, pp. 1870–1874, 2016.

[12] B. G. Hall, "Building phylogenetic trees from molecular data with MEGA," Molecular Biology and Evolution, vol. 30, no. 5, pp. 1229–1235, 2013.

[13] N. Abdennadher and R. Boesch, "Deploying PHYLP software to a large scale distributed system," in Proceedings of Seventh IEEE International Symposium on Cluster Computing and the Grid, CCGrid 2007, pp. 673–678, Rio De Janeiro, Brazil, May 2007.

[14] I. Gontia-mishra, N. Tripathi, and S. Tiwari, "A simple and rapid DNA extraction protocol for filamentous fungal efficient for molecular studies," Indian Journal of Biotechnology, vol. 13, pp. 536–539, 2014.

[15] K. Hadzic-Matic, K. Lekang, A. Lanzen, I. Jonassen, E. M. Thompson, and C. Troedsson, "Characterization of the 18s rRNA gene for designing universal eukaryote specific primers," PLoS One, vol. 9, no. 2, Article ID e87624, 2014.

[16] A. Nayarisseri, M. Yadav, M. Bhatia et al., "Impact of next-generation whole-exome sequencing in molecular diagnostics," Drug Invention Today, vol. 5, no. 4, pp. 327–334, 2013.

[17] C. W. James Cappuccino, Microbiology Laboratory Manual (Eleventh), Pearson Education Limited, San Francisco, CA, USA, 1988.

[18] A. Belhaj, N. Desnoues, and C. Elmerich, "Alkane biodegradation in Pseudomonas aeruginosaisolated strains isolated from a polluted zone: identification of alkB and alkB-related genes," Research in Microbiology, vol. 153, no. 6, pp. 339–344, 2002.

[19] A. Vomberg and U. Klimmer, "Distribution of alkB genes within n-alkane-degrading bacteria," Journal of Applied Microbiology, vol. 89, no. 2, pp. 339–348, 2000.

[20] C. N. Muholja, H. Makonde, G. Magoma, and M. Imbuga, "Biodegradability of polyethylene by bacteria and fungi from Dandora dumpsite Nairobi-Kenya," PLoS One, vol. 13, no. 7, Article ID e0198446, 2018.

[21] A. Indumathi, T. Gayathri, E. Biotechnology, T. Nadu, and T. Nadu, "Plastic Degrading ability of Aspergillus oryzae isolated from the garbage dumping sites of Thanjavur, India," International Journal of Current Microbiology and Applied Sciences, vol. 3, no. 3, pp. 8–13, 2016.

[22] S. Zahra, S. S. Abbas, M. T. Mahsa, and N. Mohnsen, "Biodegradation of low-density poly-ethylene (LDPE) by isolated fungi in solid waste medium," Waste Management, vol. 30, no. 3, pp. 396–401, 2010.

[23] N. Ojha, N. Pradhan, S. Singh et al., "Evaluation of HDPE and LDPE degradation by fungus, implemented by statistical optimization," Scientific Reports, vol. 7, no. 1, 2017.

[24] P. P. Vimala and L. Mathew, "Biodegradation of polyethylene using Bacillus subtilis," Procedia Technology, vol. 24, pp. 232–239, 2016.

[25] K. Harshvardhan and B. Jha, "Biodegradation of low-density poly-ethylene by marine bacteria from pelagic waters, Arabian Sea, India," Marine Pollution Bulletin, vol. 77, no. 1-2, pp. 100–106, 2013.

[26] B. Suresh, S. Maruthamuthu, N. Palanisamy, R. Ragunathan, and K. Navaneetha Pandiyaraj, "Investigation on biodegradability of poly-ethylene by Bacillus cereus strain Ma-Su isolated from compost," International Research Journal of Microbiology (IRJM), vol. 2, pp. 292–302, 2011.

[27] M. P. Das and S. Kumar, "An approach to low-density poly-ethylene biodegradation by Bacillus amyloliquefaciens," Biotech, vol. 5, no. 1, pp. 81–86, 2015.

[28] M. Bhatia, A. Girdhar, A. Tiwari, and A. Nayarisseri, "Implications of a novel Pseudomonas species on low density poly-ethylene biodegradation: an in vitro to in silico approach," SpringerPlus, vol. 3, no. 1, p. 497, 2014.

[29] B. Miyint, K. Ravi, M. K. Sakharkar, C. S. Lim, and K. R. Sakharkar, "Biodegradation of low density poly-ethylene (LDPE) by Pseudomonas species," Indian Journal of Microbiology, vol. 52, no. 3, pp. 411–419, 2012.

[30] R. J. Müller, "Biodegradability of polymers: regulations and methods for testing," Biopolymers Online, vol. 4, pp. 365–374, 2005.

[31] L. D. M. Sahadevan, C. S. Misra, and V. Thankamani, "A bird’s eye view on bioremediation approaches of heavy metals contaminated soil regimes," Open Access Review Article, vol. 3, no. 1, pp. 1–18, 2013, http://environmentaljournal.org/3-1/uejrt-3-1-2.pdf.

[32] J. D. Hamilton, K. H. Reinert, J. V. Hagan, and W. V. Lord, "Polymers as solid waste in municipal landfills," Journal of the Air and Waste Management Association, vol. 45, no. 4, pp. 247–251, 2014.

[33] E. M. El-morsy and E. Ahmed, "Biodegradative activities of fungal isolates from plastic contaminated soils," Mycosphere, vol. 8, no. 8, pp. 1071–1086, 2017.
H. V. Sowmya, M. Ramalingappa, and B. T. Krishnappa, "Degradation of poly-ethene by Trichoderma harzianum-SEM, FTIR, and NMR analyses. Degradation of poly-ethene by Trichoderma," Environmental Monitoring and Assessment, vol. 186, no. 10, 2014.

H. Zhang, F. Li, H. Chen et al., "Cloning, expression and characterization of a novel esterase from a South China Sea sediment metagenome," Chinese Journal of Oceanology and Limnology, vol. 33, no. 4, pp. 819–827, 2015.

J. Zhang, X. Wang, J. Gong, and Z. Gu, "A study on the biodegradability of poly-ethene terephthalate fiber and diethylene glycol terephthalate," Journal of Applied Polymer Science, vol. 93, no. 3, pp. 1089–1096, 2004.

W. M. Abd El-Rahim, O. A. M. El-Ardy, and F. H. A. Mohammad, "The effect of pH on bioremediation potential for the removal of direct violet textile dye by Aspergillus Niger," Desalination, vol. 249, no. 3, pp. 1206–1211, 2009.

M. Gyung Yoon, H. Jeong Jeon, and M. Nam Kim, "Biodegradation of poly-ethene by a soil bacterium and AlkB cloned recombinant cell," Journal of Bioremediation and Biodegradation, vol. 3, no. 4, 2012.

Van Beilen, Z. Li, W. A. Duetz, T. H. M. Smits, and B. Witholt, "Diversity of alkane hydroxylase systems in the environment," Oil and Gas Science and Technology, vol. 58, no. 4, pp. 427–440, 2003.