Molecular Characterisation and Identification of Three Mushrooms Found in the Niger Delta Region

S. B. Chuku¹*, E. O. Nwachukwu¹, I. O. Agbagwa¹ and H. O. Stanley²

¹Department of Plant Science and Biotechnology, University of Port Harcourt, Rivers State, Nigeria.
²Department of Microbiology, University of Port Harcourt, Rivers State, Nigeria.

Authors’ contributions

This work was carried out in collaboration among all authors. Author SBC designed the study, performed the statistical analysis, wrote the protocol and wrote the first draft of the manuscript. Authors EON and IOA managed the analyses of the study. Author HOS managed the literature searches. All authors read and approved the final manuscript.

Article Information

DOI: 10.9734/ARRB/2020/v35i730253

Received 05 May 2020
Accepted 11 July 2020
Published 04 August 2020

ABSTRACT

Mushrooms are a group of fungi that are diverse and in the Niger Delta region; there are various species, some of which share similar morphological features. Over the years, mushrooms have been put to several important uses ranging from food, nutraceuticals, feed for livestock, and more recently bioremediation, hence, the need for an accurate approach of identification is eminent. This study aims to identify three mushroom samples with the potential for crude oil degradation using molecular tools. The molecular identification of the mushrooms was carried out with the ITS (Internal Transcribed Spacer) region to analyze their genetic diversity. The three mushroom samples were identified as Pleurotus ostreatus, Pleurotus floridanus and Lentinus squarrosulus. The blast results showed 86.0%, 97.9%, and 88.4% for Pleurotus ostreatus, Pleurotus floridanus and Lentinus squarrosulus respectively for sequence similarity. The samples were assigned unique accession numbers on GenBank. The use of molecular characterization gives reliable results to the species level. The results from this study have increased the database of Pleurotus and Lentinus DNA and serve as a basis for the identification of unidentified species by comparing their PCR amplified sequences with ITS primers.

*Corresponding author: E-mail: sobornatechuku@gmail.com;
Keywords: Molecular; characterization; Internal Transcribed Spacer (ITS); mushroom.

1. INTRODUCTION

There has been much global suffering due to increased environmental problems resulting from intensified industrialization and the use of chemicals for agricultural practices. This indiscriminate release of chemicals into the environment has caused a high level of pollution. Remediation of oil spill impacted sites in the Niger Delta after initial clean-up has so followed two (2) conventional methods viz: chemical and mechanical. These methods in themselves have had residual effects on the environment and are very expensive to carry-out. Hence, the need for eco-friendly and cost-effective methods of remediating spill sites [1]. Bioremediation uses the biological organisms’ metabolic potential to degrade or transform hazardous compounds in the environment into less toxic or harmless forms [2,3]. The use of fungi referred to as mycoremediation, has attained widespread acceptance due to its very low substrate specificity, so they can mineralize a wide range of highly recalcitrant and persistent organopollutants [4,5,6]. Prospection for fungi is owing to its ability to secrete high levels of lignin modifying enzymes and other enzymes with desired properties for biotechnological applications [7]. Mushrooms belong to these physiological group known as white rot fungi known to produce lignin modifying enzymes. The need to accurately identify these species is very important as it is the first step to utilize them for various biotechnological applications. This study aims to identify the three mushroom samples using molecular tools.

2. MATERIALS AND METHODS

2.1 Source of Fungi

The fungi species used for this work were obtained from (BIODEC) Odi Bayelsa State. The samples were labelled as samples 1, 2 and 3 for the collection of indigenous mushrooms. These were transported to the laboratory immediately in zip lock bags.

2.2 Media Preparation

Potato dextrose agar (PDA) was used for tissue culture. This was prepared following the manufacturer’s instruction using the potato dextrose agar powder (Hi-Media). To prepare 1 Litre of the media 39 grams of PDA was dissolved in 1 litre of distilled water and autoclaved at 121°C for 15 minutes. Chloramphenicol 0.1% (w/v) was added as an antibacterial agent [8].

2.3 Tissue Culture

This was carried out at the Regional Centre for Biotechnology and Bioresources Research laboratory, University of Port Harcourt, Choba, Rivers State, Nigeria. After surface sterilization, the inner tissue of the stipe was cut with a surgical blade already flamed and cooled around the flame area. The excised tissue was inoculated into the Petri dish with freshly poured media. The Petri dishes were immediately sealed with masking tape and cling film to avoid external contamination. The plates were then incubated for 14 days in the dark at room temperature 27±2°C. This was done for each species used [9,10]. Sub-culturing was done several times to obtain a pure culture.

2.4 Molecular Characterisation

2.4.1 DNA extraction

A slight modification of the cetyltrimethylammonium bromide (CTAB) Method [11] mycelium was scraped off and transferred to already sterilized mortar. DNA was extracted with Zymo Quick DNA Fungal/Bacterial miniprep Kit, with the following steps as stated by the manufacturer Zymo Research group CA, USA. For DNA extraction, a five (5) days old culture of the three samples (1, 2 and 3). The extracted DNA was stored in the freezer and maintained on ice to avoid denaturation.

2.4.2 DNA quantification

This step was done to determine the concentration and purity of the extracted DNA using the NanoDrop 2000c Spectrophotometer (Thermo Fisher Scientific Inc., Wilmington, Delaware, USA). The sample was vortexed briefly to homogenize. 1.7 µL of the extracted DNA was used. Then sample concentration µg/µl was measured absorbance at 260/280nm wavelength appropriate for nucleic acid sample. This reading was taken in triplicates [11,12].

2.4.3 DNA quality check

To determine the quality of extracted DNA, Agarose gel electrophoresis method was adopted to determine the quality of nucleic acid
extracted [13]. The gel was viewed in a gel documentation system by exposing it to UV light and pictures were taken to show the DNA Bands.

2.5 Polymerase Chain Reaction (PCR)

PCR Amplification of DNA samples was done using universal primers in a thermal cycler (GeneAmp PCR system 9700). ITS4: Reverse (5’- TCCTCGCTTATGGATATGC-3’) and ITS5: Forward (5’- GGAAGTAAAAGTCGTAACAGG - 3’) [14,15,16]. The total reaction volume of 25µl made up of 2.5µl of 10x PCR buffer, 1µl of 25mM MgCl$_2$, 1µl each of the forward primer and reverse primer, 1µl of DMSO, 2µl of 2.5 Mm DNTPs, 0.1µl of 5µ/µl Taq DNA polymerase, and 3µl of 10ng/µl DNA and 13.4µl Nuclease free water. The PCR cycling parameters: Initial denaturation at 94°C for 5mins, followed by 36 cycles of denaturation at 94°C for 30sec, annealing at 54°C for 30secs and elongation at 72°C for 45sec. Followed by a final elongation step at 72°C for 7 minutes. Amplified fragments were visualized on Safe view-stained 1.5% agarose electrophoresis gels and photographed under UV light in a gel documentation system. A 1kb ladder (Bioline) used as a marker for the gel run served as control.

2.6 Sequencing and Bioinformatic Analysis

The products of PCR were subjected to Sanger dideoxy sequencing at IITA, Ibadan, Nigeria. The sequences obtained were deposited to the GenBank database to compare with the National Center for Biotechnology Information (NCBI) using the basic local alignment search tool (BLAST) algorithm [17]. Phylogenetic trees were constructed for each organism using the best Basic Local Alignment Search Tool BLAST hits Multiple sequence alignment with the Clustal W algorithm and the phylogenetic trees showing maximum likelihood algorithm (Juke-Cantor) was obtained on (MEGA) version 8 [18].

3. RESULTS AND DISCUSSION

DNA concentration range from 26.6 ng/µL to 30.53 ng/µL for the samples while DNA purity range from 1.66 to 1.79 respectively. Quality check was then conducted on agarose gel which produced single bands of intact DNA when viewed on the gel with a gel documentation system, the result is as shown in plate 1. The results obtained indicated that the extracted DNA were qualified and appropriate for further molecular analysis. The results indicate contaminants like proteins or phenols are absent with the lower and higher range respectively [19,20].

3.1 Concentration and Purity of Extracted Nucleic Acid (DNA)

### 3.1.1 PCR amplification

Isolated DNA from the samples used was amplified with ITS primers (ITS-4 and ITS-5). The amplified products obtained were visualized on Safe view-stained 1.5% agarose electrophoresis gel. They appeared as single intact bands on the agarose gel with safe view-stain as shown in the plate below. The size of the PCR fragment for all the samples was deduced with the ladder used. Products of PCR migrated between 654 bp and 727 bp as shown in Plate 2. The result of all the mushrooms was within the range (350 and 880 bp) for fungi species using ITS 1 and ITS 4 [21] and the range of (651 and 800 bp) [22].

Table 1. Concentration and Purity of DNA extracted from the samples used

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>DNA concentration (ng/µL)</th>
<th>DNA Purity (260/280 nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>29.97 ±2.73</td>
<td>1.79 ±0.05</td>
</tr>
<tr>
<td>2</td>
<td>30.53 ±0.40</td>
<td>1.66 ±0.01</td>
</tr>
<tr>
<td>3</td>
<td>26.60 ±1.41</td>
<td>1.66 ±0.01</td>
</tr>
</tbody>
</table>

Each value is the mean of three replicates ± standard deviation.
products were used to generate phylogenetic trees generated by Maximum composite likelihood analysis showing sample 1 as *Pleurotus ostreatus*, sample 2 as *Pleurotus floridanus* and sample 3 as *Lentinus squarrosulus* respectively.

Plate 1. Agarose gel photograph of DNA extracted of the three samples used lane 1-3 representing samples 1-3 respectively

Plate 2. Amplicons obtained from a polymerase chain reaction of the internal transcribed spacer (ITS) region of the fungi understudy

$L$ – 1Kb DNA Ladder (Bioline). The numbers represent the samples used
Table 2. Taxonomic affinities of samples with blast searches of ITS sequences on GenBank

<table>
<thead>
<tr>
<th>Sample id</th>
<th>Taxonomic affinity (Gene Bank No.)</th>
<th>Percentage similarity (%)</th>
<th>Accession number</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><em>Pleurotus ostreatus</em> (MN049516.1)</td>
<td>86.00</td>
<td>MN968774</td>
</tr>
<tr>
<td>2</td>
<td><em>Pleurotus floridanus</em> (MN173381.1)</td>
<td>97.88</td>
<td>MN968775</td>
</tr>
<tr>
<td>3</td>
<td><em>Lentinus squarrosulus</em> (KT273380.1)</td>
<td>88.44</td>
<td>MN968776</td>
</tr>
</tbody>
</table>

Fig. 1. Phylogenetic tree generated by Maximum composite likelihood analysis based on the ITS 1-2 gene sequences

Fig. 2. Phylogenetic tree generated by Maximum composite likelihood analysis based on the ITS 1-2 gene sequences
Fig. 3. Phylogenetic tree generated by Maximum composite likelihood analysis based on the ITS 1-2 gene sequences

4. CONCLUSION

The mushrooms in the Niger Delta region are diverse. Most of them share identical features which could lead to conflicting results making it inappropriate to rely on morphological characterization for identification. The use of molecular characterization gives reliable results to the species level. The results from this study have increased the database of Pleurotus and Lentinus DNA and serve as a basis for the identification of unidentified species by comparing their PCR amplified sequences with ITS primers.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES


