INTRODUCTION

Kingdom Fungi is the second largest kingdom after insects [1]. They are found all over the planet, even in the most hostile and uninhabitable places such as in Antarctic ice, mountain surfaces, rocks, seawater and habitable places like the tropics and temperate regions [2]. Fungi possess arrays of many chemical molecules with biochemical activities. They produce several types of enzymes and secondary products of metabolism, which include antibiotics, vitamins, polysaccharides and organic acids [3]. Fungi play very important roles in human daily life, which ranges from its use in many industrial processes to agriculture, medicine, food processing, textile, biomedical production, natural cycling, and production of bio-fertilizers and so on. Therefore, fungal biotechnology has become a very important part of human welfare [4]. Fungi are fast-growing organisms that do well on the surface of most plant and other organic substances. They have adapted to survive on many substrates in other to survive, reproduce and compete with other life forms. Fungi have divers mood of nutrition such as symbiotic, parasitic and saprophytic. Apart from humans, fungi also play a beneficial and necessary part in the survival and growth of many plants by suppressing plant root diseases, protecting the plants from pathogens; they also improve their host plant ability to acquire nutrients by antagonism through the production of plant repellent. Fungi also provide for their host plant many vitamins needed for plant growth. Some of the metabolite produced by fungi within a plant host increases the plant ability to withstand the invasion of predators, parasites and diseases and may facilitate reproductive processes. One of the known characteristics of fungi is their nature as food spoilers that causes damage to raw and cooked food. Some species of fungi are plant pathogens such as the causative organism of Potato late blight caused by Phytophthora infestans, which was responsible for the Potato famine of Ireland in 1845-1849, and rice blast which caused the infamous Bengal famine of 1943. The fungi ergot formed by mycelium of Claviceps spp especially C. purpureus have been implicated for the poisoning of cattle. The mycotoxins aflatoxins and ochratoxin A produced by many species of Apergillus are a common cause of food poisoning in many countries. Nevertheless, fungi have been found to be beneficial to man as producers of many different metabolites such as antibiotics, enzymes such as cellulase, lipase and lginolysinolytic enzymes. They are also known for the production of alkaloids such as ergot alkaloids from Claviceps, pigments such as anthraquinone and betalains. Fungi have also been reported in biological control of nematodes, also are good sources of aroma and flavor. Some edible fungi are also a source of nutrition as they provide essential elements such as selenium, potassium, riboflavine, niacin and vitamin D. In addition, edible fungi can also prevent or treat some diseases such as Parkinson, Alzheimer, hypertension, cancer and stroke [2, 5]. Fungi could perform these functions due to array of metabolites which they possess such as: esters, aldehydes, amonic compounds, organic acids, etc [6].

MATERIALS AND METHODS

Soil samples collection

Soil samples were collected at various locations in the Horticultural garden of LAUTECH and then taken to Microbial Resources and Research Laboratory, Ogbomoso. This was serially diluted and inoculated onto a prepared Potato Dextrose Agar (PDA) plate containing antibacterial (Chloramphenicol) to inhibit bacteria growth on the plate with one plates serving as control. They were incubated at 28 °C for 24–72 h. pure fungi isolates were obtained by sub-culturing The Potato Dextrose Broth (PDB) was prepared by dissolving 24 g in 1 liter of distilled water, it was mixed and dissolves by heating with frequent agitation, and it was boiled for some minutes until complete dissolution is obtained according to the manufacturer’s guidelines. Then it was dispersed into appropriate bottle; 500 ml in eight places, sterilized in an autoclave at 121 °C for 15 min.
Submerged fermentation (smf) of soil fungi isolates

Submerged fermentation (smf) was done using Potato Dextrose Broth (PDB). The eight isolates were coded as OT, AS, OS, GT, AT, GS, BT, BS and were inoculated into the broth medium in a 500 ml Erlenmeyer flasks. All experiments were carried out aseptically inside laminar air-flow chamber. The flasks were placed in a rotary-shaker incubator at 28 °C for seven (7) days while biomass and metabolite production were monitored daily.

Extraction of the fungi metabolite

The organisms produced their metabolites intracellularly-that is within its cells. Extraction of the metabolites was performed after 21 d of culture. The biomass was filtered using sterile muslin cloth (sterilized in an autoclave at 121 °C for 15 min) while the metabolites were extracted using Ethanol at ratio 2:1 of the culture media. The metabolates were obtained after centrifugation at 5,000 rpm for 10 min. The metabolates were obtained for each of the soil fungus and then kept at 4 °C for further use.

Molecular characterization of the soil fungi

Fungal genomic DNA extraction

Fungal isolates were grown on cellophane membrane fixed on Potato Dextrose Agar and were incubated at 25 °C. The fungal isolates weighing about 50-100 mg (wet weight) were placed in a sterile ZR Bashing Lysis Tubes containing 750 µl of lysis buffer, the mixture was put in a bead fitted 2 ml tube holder assembly and crushed at the highest speed of 5 min, the preparation was swirled briefly in the ZR Bashing Bead™ Lysis Tube then centrifuged at 10,000 rpm for one minute a micro-centrifuge. Resultant supernatant of about 400 µl was transferred into a Zymo-Spin™ IV Spin Filer (orange top) in a collection tube and centrifuged for one minute at 7,000rpm, a 1200 µl measure of fungal DNA binding buffer was added to the filtrate in the collection tube. Then, 800 µl of the mixture was transferred to a Zymo-Spin™ IIC Column in a Collection Tube and centrifuged 10,000 rpm for 1 minute period. The flow-through was discarded then. 200 µl DNA pre-washed buffer was added to the Zymo-Spin™ IIC Column in new collection tube; this was centrifuged at 10,000 rpm for 1 minute. After centrifugation, 500ul fungal DNA wash buffer was added to a Zymo-Spin™ IIC Column and centrifuged at 10,000 rpm for one minute. Zymo-Spin™ IIC Column was transferred to a clean 1.5 ml micro-centrifuge tube and 100 µl (35 µl minimum) DNA elution buffer was added directly to the column matrix and centrifuged at 10,000rpm for 30 seconds to elute the fungal DNA. DNA was then suitable for PCR and other downstream applications.

Polymerase chain reaction (PCR) of the extracted fungal DNA

The extracted fungal DNA were amplified in a thermal cycler (Gene Amp PCR system 9700); the PCR reaction mixture consisted of 190ul extracted fungal DNA 2.5ul 10x PCR loading buffer, 2.0ul 2.5Mm DNTPs mix, 1.0ul 25Mm mgcl2, 1ul DMSO, 5ul of tag polymerase and 1ul each of the forward and reverse primer;

ITS1F: 5’-TCCGATAGTGAACCTGCGG-3’ and ITS2R: 5’-GCTGCGTATTCTCATGGATGC-3’

Antimicrobial susceptibility test

Four (4) clinical isolates namely: Staphylococcus aureus, Klebsiella spp, Candida albicans and Escherichia coli isolated from clinical samples at LAUTECH, Ogbomoso were used to test for the antimicrobial activities of the metabolites (the metabolites were tagged as: OT, AS, OS, GT, AT, GS, BT, BS). Muler Hilton agar was prepared and poured into Petri dish. After it has cooled down, the clinical isolates were evenly streaked onto the surface of the plate. 100 µl of each of the metabolates was impregnated onto a paper filter paper disc and then placed on the center of the plates. All the eight (8) metabolates were tested against the four (4) clinical isolates. They were incubated for 18-24 h at 37 °C. The inhibitory activities of the metabolates against the pathogenic organisms was determined and measured in mm.

Protein structure and function of the fungal isolates

After the DNA nucleotide sequences were obtained, the sequence was transferred to European Molecular Biology open software suit (EMBOSS) to translate the sequence into amino acid. The amino acid sequence was transferred to PHYRE 2 for the possible protein prediction and viewed using Pymol software.

RESULTS

Isolation and prevalence soil fungi

The soil samples were collected inside foil papers using spatula to dig through the soil at a depth of 1-5m. Serial dilution was performed for the fagi isolation. The diluted of 10^-8 and 1.0^8 test tubes was inoculated into the prepared Potato Dextrose Agar (PDA) plate containing antibacterial (Chloramphenicol) to inhibit bacteria growth on the plate. The plates were incubated at 28 °C for about 3 d. Growth pattern was observed from the third day. The cultures were transferred onto slant bottles for biochemical characterization and identification. (Plates 1-2). In this study, isolates of the Aspergillus genus had the highest occurrence (62.5%) followed by Aspergillus aculeatus (50%), Aspergillus Carbonarias (12.5%) followed by Penicillium genus (25%) Penicillium verrucosum (12.5%), Penicillium cryoenogen (12.5%) and Talaromyces albobilverticillus (12.5%).

Molecular identification and evolutionary relationship of soil fungi

The nucleotide sequences obtained were blasted at the National Center for Biotechnological Information (NCBI) Genbank FASTA formats were obtained of other related sequences and used to plot a phylogenetic tree that established the evolutionary relationship with the query sequence (fig. 1-7). The soil fungal isolates were identified to be: Penicillium verrucosum, Talaromyces albobilverticillus and Aspergillus spp which were A. aculeatus and carbonarius (table 1).

Plate 1: Culture of soil fungi on PDA plate

Plate 2: Cultures of soil fungi preserved on PDA slants
Table 1: Soil fungi identity

| Code    | Molecular identity          |
|---------|----------------------------|
| LAU-MAJ-ASF | Penicillium verrucosum   |
| LAU-MAJ-BSR | Aspergillus aculeatus    |
| LAU-MAJ-BTF | Aspergillus carbonarius  |
| LAU-MAJ-GXF | Talaromyces alboplicicillus |
| LAU-MAJ-GTF | Aspergillus aculeatus    |
| LAU-MAJ-OSF | Aspergillus carbonarius  |
| LAU-MAJ-OTF | Aspergillus aculeatus    |
| LAU-MAJ-ATF | Penicillium chrysogenum  |

Fig. 1: Phylogenetic relationship of LAU-MAJ-ASF and Penicillium species

Fig. 2: Phylogenetic relationship of LAU-MAJ-BSR and Aspergillus species

Fig. 3: Phylogenetic relationship of LAU-MAJ-BTF and Aspergillus species
The evolutionary history of the isolates was determined by UPGMA [7]. The sum of the branch length of the optimal tree was shown to be 0.61519578. Evolutionary distances were determined using the maximum composite method [8, 9]. The analyses include 16 nucleotide sequences. Every ambiguous positions were deleted for each sequence pair. A total of 628 positions were recorded for the final dataset. Evolutionary analyses were conducted in MEGA X [9].

![Fig. 4: Phylogenetic relationship of LAU-MAJ-GSF and Talaromyces species](image)

![Fig. 5: Phylogenetic relationship of LAU-MAJ-GTF and Aspergillus species](image)

![Fig. 6: Phylogenetic relationship of LAU-MAJ-OSF and Aspergillus species](image)

**Protein sequence of metabolite**

After the DNA nucleotide sequence was obtained, the sequence was transferred to European Molecular Biology Open Software Suite (EMBOSS) to translate the sequence into amino acid. The amino acid sequence was transferred to PHYRE 2 software that was used to model the 3D protein structure and viewed with Pymol Software (Plate 3a-h). In terms of functions, the proteins were found to be structural protein, proton transporter, oxidoreductase and hydrolase/hydrolase inhibitors.
Fig. 7: Phylogenetic relationship of LAU-MAJ-OTF and Aspergillus species

Plate 3 (a-g) Modelled 3D Protein structure (a-Penicillium verrucosum; b-Penicillium chrysogenum; c-Aspergillus aculeatus; d-Aspergillus carbonarius; e-Talaromyces albolibiverticillus; f-Aspergillus aculeatus; g-Aspergillus aculeatus) showing the amino acid residues
The need for new and more effective antibiotics is on the rise as infectious agents of health concern.

**DISCUSSION**

**Morphological identification of the soil fungi**

There have been several reviews on successful methods of isolating fungi from nature. There are several fungal species distributed in soil, plant debris, etc. The soil acts as a growth media for fungi in various parts of the globe [10]. Many researches have focused on the prevalence and isolation of keratinophilic fungi from soil samples [11-13]. Recently, saprophytic fungi have also been isolated from soil as it causes diseases with people with low immunity [14].

Soil is the most essential source for the isolation of fungi [15]. For example, species belonging to the genera Aspergillus, Penicillium and Mucor have been isolated from roots of soils in different agricultural sites. The results of another study conducted [16] showed that several factors such as pH, organic carbon, moisture content, sulphur, nitrogen and potassium have great effects on soil microbial diversity [17, 18]. However, these microbes have great potentials applications in drug discovery.

**Antimicrobial susceptibility test**

The need for new and more effective antibiotics is on the rise as cases of antibiotic resistance strains of pathogens are increasing daily. This needs have been met for decades largely by the production of semi-synthetic product from the natural product, but in recent years, advancement in technology have rebuffed researchers in the discovery of natural product and antibiotics from microbial sources, this include finding new antibiotics from older ones as well as from new sources. This breakthrough has led the discovery of new antibiotics very novel mechanisms of action capable of forming the bases of new antibiotic classes useful in treating pathogens. Fungi are known for their characteristics of synthesizing a wide range of bioactive metabolites [9]. Naturally occurring antibiotics are mostly isolated from soil microbes. These antibiotics substances contribute vital roles in their establishment on (rhizoplane) and around (rhizosphere) the roots of plants. Therefore, to achieve novel compounds, search are to be made from soil microbes; such isolates are potential sources of natural products that possess secondary metabolites and enzymes [20-22].

**CONCLUSION**

In conclusion, the continuous exploration of soil microbes, which are reservoirs of drug for the development of novel products is essential for new drug discovery and development. In addition, biomolecules from natural products which are eco-friendly have proven to be abundantly available for utilization as active bio-products against infectious agents of health concern.

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**AUTHORS CONTRIBUTIONS**

The authors have contributed equally in preparing the work colation of data and writing of the manuscript.

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### Table 1: Zone of inhibition by the soil fungi metabolites against clinical isolates

| Fungi metabolites | Zone of inhibition in diameter (mm) |
|-------------------|------------------------------------|
|                   | Staphylococcus aureus | Escherichia coli | Candida albicans | Klebsiella spp |
| Penicillium verrucosum(AS) | - | - | - | - |
| Penicillium chrysogenum (AT) | - | - | - | - |
| Aspergillus aculeatus (BS) | 23±0.55 | 9±0.47 | - | 24±2.45 |
| Aspergillus carbonarius (BT) | 21±2.67 | - | - | - |
| Talaromyces albobiverticillus (GS) | - | - | - | - |
| Aspergillus aculeatus (GT) | - | - | - | - |
| Aspergillus carbonarius (OS) | - | - | - | - |

Resistant (-)

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**Determination of zones of inhibition against clinical isolates**

The antimicrobial activities of the metabolite of the eight (8) isolates were carried out against four (4) pathogenic organisms of clinical origins namely: Staphylococcus aureus, Escherichia coli, Candida albicans and Klebsiella spp. Out of all the metabolite tested against the pathogens, the metabolite of Aspergillus aculeatus (BS) showed the highest and prominent inhibition against S. aureus (23±0.55 mm), E. coli (9±0.47 mm), Klebsiella spp (24.00±2.45 mm) but no zone of inhibition against C. albicans. Metabolite of A. carbonarius had a zone of inhibition of 21±2.67 mm against S. aureus but no zone of inhibition against C. albicans, Klebsiella spp and E. coli. The clinical isolate showed resistance to metabolites of Penicillium verrucosum, Penicillium chrysogenum, Talaromyces albobiverticillus and Aspergillus aculeatus (GT) as presented in table 1.

**Fig. 8: Comparing fungi metabolites with synthetic antibiotics using Klebsiella spp and Escherichia coli as test organisms**
CONFLICT OF INTERESTS

Authors do not have any conflict of interest in any way.

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