Endophytic Fungal Diversity in *Hardwickia binata*: Bridging the Gap between Traditional and Modern Techniques

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Abstract: Endophytic fungus is crucial for maintaining plant health and defense mechanisms, acting as protective barriers against pathogens, and producing medicinally beneficial bioactive compounds. Genome sequencing and metagenomics have significantly enhanced the understanding of fungal diversity and metabolic capabilities, enabling the identification of new genes and substances. Traditional culture-dependent methods have been complemented by culture-independent techniques, offering a more comprehensive view of fungal diversity. Using both culture-dependent and culture-independent techniques, the present research investigation explored the diversity of endophytic fungi encountered in the foliage of *H. binata*. The study examined the topographical characteristics and nutritional content of soil samples collected from the locality of the selected plant sample, *H. binata*, to better comprehend the effects on the plant’s growth. The balanced nutrient constituted approximately a pH of 7.2, which suggested an alkaline nature and promoted plant development. The ratio of nitrogen, phosphorous, and potassium remained 3:1:1. A total of 25 fungal isolates, categorized into 17 morphotypes, were obtained using the culture-dependent approach; *Curvularia* and *Nigrospora* emerged as the most common genera. Furthermore, the prediction of the ITS2 secondary structure supports the identification of species, highlighting a wide variety of fungal species present in *H. binata*. The culture-independent approach generated 69,570 high-quality sequences, identifying 269 Operational Taxonomic Units (OTUs). The dominant Ascomycota phylum, along with various genera, indicated a rich fungal community associated with *H. binata*. This study advances the understanding of the endophytic fungus communities that are associated with *H. binata* and the nature of soil ecology. The findings emphasize the significance of holistic techniques in the study of microbial dynamics within plant systems as well as their implications for ecosystem management and plant health.  

Keywords: endophytic fungi; culture-dependent; ITS2; OTUs; microbial dynamics

1. Introduction

Endomycetous fungi thrive in healthy plant tissues and are symbiotically associated with most plants in their natural ecosystems. This allows for the identification of endophytic microbes within every examined plant genus [1,2]. Endophytic microorganisms provide bioactive chemicals that enhance plant development, adaptation, uptake of nutrients, fixation of nitrogen, stress tolerance, and disease control, thereby establishing long-lasting mutually beneficial relationships [3]. Fungal abundance, on the other hand, varies substantially among habitats and ecosystems [4]. Fungi are extremely varied and difficult to predict accurately, which makes academic research particularly challenging.
Fungal endophytes work with host plants to regulate development, allowing researchers to assess the diversity and species of fungal communities [5]. Despite the diversity of plant species found in tropical climates, research into strategies for isolating endophytes from these habitats has recently increased [6,7]. Surrounded by a wide variety of native flora and animals, the Nagamalai hills, 12 km south of Madurai city, possess untapped assets. Popular by several names, “Anjan,” or *Hardwickia binata* Roxb, is a deciduous tree that has been used in traditional medicine to cure a wide range of conditions, including leprosy, dyspepsia, leucorrhea, cancer, diarrhea, and Gram-positive and Gram-negative bacteria [8,9]. As far as we can determine, *H. binata*, one of the most common woodland plants, has no record of endophytic fungus.

Microbial research, especially on fungi, is challenging because of the wide range of species and the challenge of developing accurate forecasts. However, different researchers have made different estimates about the number of fungal species and the expected diversity of fungal ecosystems on Earth [10]. Analyzing the structure and composition of the microbial communities associated with hosts has been made easier with the gradual development of many approaches [11]. There are two different approaches that have been used to identify endophytic fungi: cultivation-dependent screening, which uses traditional microbiological methods to isolate microbes on solid media, and cultivation-independent screening, which uses metagenomic analysis [12]. Plant endophyte communities can be studied using both direct sequencing and culture-dependent methods. Culture-dependent methods include putting sterile tissue on an agar medium, promoting endophyte development, and keeping cells in pure culture for genetic and morphological identification. Direct sequencing is an additional possible method. A combination of primers is used in PCR to get the necessary sequences from the DNA extracted from the surface-sterilized plant tissue. Sequences having a “known identity”, which is usually (but not always) discovered in a public database, are compared to the obtained sequences in the next phase [9].

From culture-based techniques to molecular biology, endophytic diversity research has progressed, providing new insights. The majority of fungal identification methods require viewing fungi in their natural habitat or grown on growth media, whereas mycological research uses morphological, phylogenetic, and ecological aspects for identification. Although studying fungal variation can greatly benefit from conventional methods, molecular approaches offer more sophisticated methodologies. For taxonomic identification, cultivation is insignificant; however, molecular methods can be beneficial for reliably distinguishing fungi by examining their morphology [13,14]. Mycologists were able to investigate newly obtained as well as previously maintained microbial samples with greater accuracy via a combination of more conventional methods (morphological assessments) with amplification technology. Concerning this, several new taxa have been proposed or established by fungus taxonomists.

A non-cultivational approach to studying genomes from ambient microbes is known as metagenomics [15]. Almost 99% of the microorganisms in an environmental sample may be identified using this method [16]. Interpreting the environment and its inhabitants has been completely transformed by the concept of exploring the entire microflora. The genetic study of environmental samples has emerged as a key technique for obtaining a deeper understanding of the ecological diversity, operational and structural variation, and evolutionary history of the species [17]. A well-known option for high-throughput genome sequencing includes the Illumina MiSeq and HiSeq technologies. The platforms can generate millions of measures within just one cycle and process hundreds of samples simultaneously. These methods have examined fungal diversity in a variety of samples based on recorded data [18]. The main objective of the current study is to comprehend the diversity and cultivation potential of the fungal community in *H. binata* leaves. To this end, the two approaches: one that is culture-dependent and involves cultivating and identifying endophytic fungi, and another that is culture-independent and involves direct molecular identification of endophytic fungi. The results shed information on the factors influencing endophyte diversity and how to optimize endophyte discovery efficiency.
2. Materials and Methods

2.1. Materials Used

The chemicals and media components used in the current study were obtained from HiMedia, Mumbai, India. The internal transcribed spacer (ITS) primers for molecular identification were obtained from Sigma-Aldrich, Bangalore, India.

2.2. Description of the Site, Selection of Plants, and Soil Analysis

An increasing interest among research organizations in the isolation of endophytes prompted an investigation into the isolation of endophytic fungi in different regions of the Nagamalai hills in Madurai. The research site, located at an altitude of 150 m, and a latitude of 9°54′ N, and a longitude of 78°00′ E, contains a wide variety of plant species distributed across its nearly four kilometers of 1500-foot-tall terrain [19]. The hill’s subhumid eco-climate is slightly dry due to limited drainage capacity, despite its rich flora and fauna. The endophytic fungal diversity analysis focused on *Hardwickia binata* Roxb, a woodland plant in the Fabaceae family and Caesalpinioideae (Figure 1). The confirmation of the botanical samples’ authenticity was conducted by the Botanical Survey of India, Coimbatore. Following the procedure guidelines indicated by Tamilnadu Agricultural University (TNAU) (https://agritech.tnau.ac.in/agriculture/agri_soil_sampling.html, accessed on 31 March 2022), soil samples were obtained from the *H. binata* research location. Topographic features were documented. Soil physicochemical parameters were analyzed by sending soil samples to the Agriculture Development, ICAR Krishi Vigyan Kendra, CENDECT, Soil and Water Analysis Center, Theni (Dt), Tamilnadu, India.

![Hardwickia binata Roxb.](image)

**SYSTEMATIC POSITION**

*Kingdom: Plantae*

*Phylum: Spermatophyta*

*Class: Dicotyledonae*

*Order: Fabales*

*Family: Fabaceae*

*Genus: Hardwickia.*

*Species: Hardwickia binata Roxb.*

Figure 1. Taxonomic classification of the selected plant.

2.3. Collection of Plant Samples

About sixty completely developed, dark green leaves were hand-picked from the lowest branches of *H. binata* to directly collect the endophytic fungus populations. The samples were collected according to strict aseptic standards and promptly transported to the laboratory in sterile bags to reduce the chance of contamination. The endophytic fungus identified in the foliage of *H. binata* was isolated and studied after being properly sterilized and diced into three hundred segments via a clean, sterile blade. The resulting segments were subsequently rinsed under running water.

2.4. Culture-Dependent Approach

2.4.1. Endophytic Fungi Isolation and Morphological Characterization

The leaf segments were rigorously surface sterilized under the [20] methodology to remove any exterior contaminants and make it less difficult to isolate endophytic fungi. The process included providing the foliage sections with a complete rinse with normal water before submerging the leaf bits in a 70% ethanol and 4% NaOCl solution for disinfection. After being air-dried using sterile filter papers, the sterilized foliage segments have been
rinsed in sterilized distilled water to remove any remaining moisture residue. Potato dextrose agar plates were used to examine the effectiveness of the surface sterilizing process. Effective sterilization can be determined by the absence of microbial growth. To avoid bacterial and fungal contamination, the sterilized leaf segments were laid out on a medium encompassing potato extract, dextrose, agar-agar, and streptomycin. Subsequent to an incubation period of 7 days, the fungal isolates were subcultured onto new PDA plates at 24 °C following a 10-day period. Pure cultures were kept at 4 °C for further examination and preservation once morphological characterization was completed.

2.4.2. Assembling Sequences and Phylogenetic Tree Construction

The protocol described by [21] was followed for the genomic DNA extraction from fungal mycelia. The isolated DNA was run through agarose gel electrophoresis (0.8%) to determine its purity. Using fungal specific primers (ITS1-F (5′-CTTGGTCATTTAGAGGAAGTA-3′) and ITS4-R (5′-CTTGGTCATTTAGAGGAAGTA-3′), the Internal Transcribed Spacer (ITS) region was amplified. A polymerase chain reaction (PCR) was executed in a 25 µL reaction consisting of 1 µL of DNA template, 1.25 µL of forward and reverse primers, and 12.5 µL of 2 × DNA master mix (Amplicon). The PCR amplification was performed using the Prima-96™ Thermocycler instrument from Himedia, Mumbai, India. The reaction conditions included denaturation at 94 °C for 4 min, followed by 30 cycles of denaturation at 94 °C for 30 s, annealing at 58.2 °C for 1 min, and extension at 72 °C for 2 min. Finally, there was a 7 min extension at 72 °C. Confirmation of the amplicons was executed on 1% agarose gels, followed by purification and sanger sequencing performed by Bio kart India Pvt. Ltd., located in Bangalore, India. Analyzed using NCBI and BLAST, the amplified sequence yielded valuable insights. Additionally, a phylogenetic tree was meticulously crafted using MEGA X [22].

2.4.3. ITS2 Secondary Structure Prediction

By using GenBank annotation or Hidden Markov models (HMMs) executed via the web server (http://its2.bioapps.biozentrum.uni-wuerzburg.de, accessed on 15 April 2022), the ITS2 area was determined and delineated [23]. Subsequently, the BLAST algorithm was used to do sequence similarity searches. A comparative analysis was conducted between each consensus sequence and published sequences obtained from reference species in various culture collections. The phylogenetic analysis only used sequences that were associated with verified species. The phylogenetic analysis used the ITS2 sequences acquired from reference species with the best coverage and identity using the BLAST search. The ribotypes of the endophytic isolates and the reference taxa were annotated using the ITS2 Database [24]. To generate synchronous alignment of sequence and secondary structures, 4SALE V1.7.1 was used with the ITS2 sequence and secondary structure for all of the sequences. The present study has modeled the genera-specific consensus secondary structure and analyzed compensatory base change (CBC). As mentioned earlier, the phylogenetic tree was rebuilt using the Phangorn program, which is part of the R framework. Using the Phangorn tool in the R environment, the phylogenetic tree was recreated. The General Time Reversible (GTR+I+G) substitution model was used to build the tree using the maximum likelihood (ML) approach. Phylogenetic tree branch support values were computed using 1000 repetitions.

2.5. Culture-Independent Approach

DNA Extraction and Metagenomic Analysis

The genomic DNA was extracted from the homogenized leaf samples using the QIAGEN soil kit. Afterwards, the concentration and purity of DNA were evaluated using a 1% agarose gel. Then, PCR amplification was carried out with specific primers (ITS1: 5′TTGTCATTAGGAAAGTA-3′ and ITS2: 5′GCTGCGTTCTCATCGATGC3′) with unique barcodes to amplify the nuclear ribosomal Internal Transcribed Spacer (ITS) region. PCR reactions were performed using TAQ Master Mix, with cycling conditions consisting...
of 30 cycles at 95 °C for 45 s, 56 °C for 45 s, and 72 °C for 60 s, followed by a final extension at 72 °C for 10 min. Products of around 400–450 bp were examined for further analysis. Subsequently, the PCR products were quantified by measuring the 260/280 ratio using Nano Drop readings. To remove any remaining primers, the purified products were treated with Ampure beads. After conducting an additional 8 cycles of PCR, Illumina barcoded adapters were used to prepare the sequencing libraries. The libraries were then purified and quantitated using the Qubit dsDNA High Sensitivity assay kit. Sequencing was performed using the Illumina Miseq platform and a 2X00PE sequencing kit, resulting in the production of 250 bp paired-end reads for subsequent analysis. The culture-independent documentation was performed according to the standard procedure followed by Biokart Pvt. Ltd., Bangalore, India.

2.6. Data Analysis in the Pipeline

Paired-end reads were matched to samples using their distinct barcodes, and then the barcode regions and primer sequence were removed. Paired-end reads were combined, and splicing sequences were acquired. Comprehensive filtering was conducted on the splicing sequences to ensure the acquisition of pristine, high-quality tags [25]. The tags were compared with a reference database to detect and remove any chimeric sequences. Sequence analyses of the clean tags were conducted using the QIIME2 pipeline, a powerful tool for analyzing genetic data. Sequences that shared a similarity of at least 97% were grouped together into Operational Taxonomic Units (OTUs). Sequences from each OTU were carefully examined for additional annotation. For taxonomic assignment of the representative sequences, the Unite Database (https://unite.ut.ee/, accessed on 30 April 2022) [26] was used with the blast algorithm. Individual data points were eliminated from the dataset. All further analyses were conducted using this normalized dataset.

3. Results

3.1. Soil Analysis

The soil sample’s nutritional content and topographical characteristics were examined in the investigation (Table 1). The findings showed that the soil at the H. binata location had a pH of 7.2, which is somewhat alkaline. The soil seems to be rich in organic matter, as shown by the measured organic carbon content of 7.4%. A modest amount of salt was indicated by an electrical conductivity of around 0.65 dS/m. The soil had a balanced nutritional profile that was good for plant development, with a nitrogen-phosphorus-potassium (NPK) ratio of 3:1:1. In addition, the amounts of iron (Fe), manganese (Mn), zinc (Zn), copper (Cu), and boron (B) were determined to be 2.34, 1.74, 1.35, 1.5, and 0.35 units, respectively, when testing the soil for micro-nutrient availability. These results shed light on the soil’s fertility and its ability to sustain plant life, in particular the development of H. binata.

Table 1. Summary of soil nutrients at sampling locations of the study.

<table>
<thead>
<tr>
<th>Soil Nutrients</th>
<th>Hardwickia binata</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPK</td>
<td>2:1:1</td>
</tr>
<tr>
<td>Electrical conductivity</td>
<td>0.65 dS/m</td>
</tr>
<tr>
<td>pH</td>
<td>7.4</td>
</tr>
<tr>
<td>Fe</td>
<td>2.34</td>
</tr>
<tr>
<td>Mn</td>
<td>1.74</td>
</tr>
<tr>
<td>Zn</td>
<td>1.35</td>
</tr>
<tr>
<td>Cu</td>
<td>1.59</td>
</tr>
<tr>
<td>B</td>
<td>0.35</td>
</tr>
</tbody>
</table>
3.2. Culture-Dependent
Cultured Endophytic Fungus: Diversity and Community Structure

The investigation into endophytic fungal diversity within the foliage of *H. binata* employed a culture-dependent approach, yielding 60 fungal isolates from symptomless leaves. Subsequent morphological analysis revealed 25 distinct isolates, which were further categorized into 17 morphotypes based on cultural traits (Figure 2). This classification served to streamline the dataset by eliminating redundant cultures, enhancing the precision of subsequent analyses. Morphological and phylogenetic analyses, utilizing subsets of ITS sequence data, provided insights into the taxonomic composition of endophytic isolates sourced from *H. binata*. Genus-level classification revealed enrichment in taxa such as *Fusarium* 8%, *Phoma* 4%, *Didymella* 4%, *Talaromyces* 4%, *Sphaeropsis* 4%, *Curvularia* 16%, *Rhytidhysterom* 8%, *Diaporthe* 4%, *Phyllosticta* 4%, *Aspergillus* 8%, *Sordaria* 4%, *Nigrospora* 12%, *Xylaria* 4%, *Pestalotiopsis* 4%, *Daldinia* 4%, and *Pseudopithomyces* 4%. These taxa span diverse fungal classes, including Sordariomycetes, Dothidiomycetes, Ascomycetes, Eurotiomycetes of the phylum Ascomycota, and Agaricomycetes of the phylum Basidiomycota. The taxonomic distribution of morphotypes isolated from the plant samples is detailed in Figure 3. Among these, the genus *Curvularia* and *Nigrospora* emerge as the predominant genera, constituting 16% and 12%, respectively, of the isolated endophytic fungi. These morphotypes were further identified by the use of ITS2 secondary structure prediction.

Daldinia (167 bp), and Sordaria (170 bp), Phyllosticta, Xylaria psidii, 20-
curvata, and others (Supplementary Figure S1 and Table S1). Phylogenetic analysis coupled with ITS2 secondary structure prediction led to the identification of Curvularia lunata, Nigrospora lacticolonia, and Daldinia eschscholtzii, alongside the recognition of 25 isolates encompassing diverse fungal species within H. binata Table 2. By analyzing the RNA molecules, the distinct genetic structures in the conserved nuclear region known as the ITS2 secondary structures may be able to help distinguish between the endophytic fungal individuals at the genotypic level.

Figure 3. Phylogenetic relationships between reference strains and representative isolates as determined by the ITS region (outgroup as Plenodomus). Species distribution along genus lines as determined by the culture-dependent approach.

3.3. Prediction of ITS 2 Secondary Structure

Comparative analysis revealed a similarity in taxonomy preferences between the Internal Transcribed Spacer (ITS) and ITS2 regions in fungi. Notably, ITS2 exhibited promising potential as a standalone marker for species delimitation within fungal communities. Utilizing sequences retrieved from the ITS2 database, phylogenetic reconstruction was conducted to ascertain fungal species diversity. Within H. binata, 17 morphotypes underwent ITS2 secondary structure prediction (Figure 4), exhibiting lengths ranging from 155 to 207 bp. Notably, Fusarium (156 bp), Rhytidhysteron (158 bp), and Curvularia (171 bp) presented with three helices, while others, including Phoma (160 bp), Didymella (160 bp), Talaromyces (160), Sphaeropsis (165 bp), Diaportha (162 bp), Phyllosticta (167 bp), Caprinellus (207 bp), Aspergillus (170 bp), Sordaria (155 bp), Nigrospora (159 bp), Xylaria (164 bp), Pestalotiopsis (170 bp), Daldinia (167 bp), and Pseudopithomyces (161 bp), formed four helices with elongated third helices. Compensatory Base Changes (CBC) analysis indicated the absence of CBCs among isolates like Fusarium, Rhytidhysteron, Curvularia, and others (Supplementery Figure S1 and Table S1). Phylogenetic analysis coupled with ITS2 secondary structure prediction led to the identification of Curvularia lunata, Nigrospora lacticolonia, and Daldinia eschscholtzii, alongside the recognition of 25 isolates encompassing diverse fungal species within H. binata Table 2. By analyzing the RNA molecules, the distinct genetic structures in the conserved nuclear region known as the ITS2 secondary structures may be able to help distinguish between the endophytic fungal individuals at the genotypic level.
**Figure 4.** Isolated fungi from the culture-dependent method: a consensus model for their ITS2 secondary structure.

<table>
<thead>
<tr>
<th>Phylum</th>
<th>Class</th>
<th>Order</th>
<th>Family</th>
<th>Genus</th>
<th>Species</th>
</tr>
</thead>
<tbody>
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<td>Sphaeriales</td>
<td>Sordariaceae</td>
<td>Sordaria</td>
<td>Sordaria</td>
<td>Sordaria fimicola</td>
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<td>Dothideomycetes</td>
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<td>Patellariaceae</td>
<td>Rhytidhysteron</td>
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<td>Psathyrellaceae</td>
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<td>Coprinellius</td>
<td>Coprinellius radians</td>
</tr>
</tbody>
</table>
3.4. Culture-Independent Examination of Endophytic Fungal Community Structures Using a Culture-Independent Approach

A total of 69,570 high-quality sequences were obtained from Illumina Miseq sequencing analysis after quality control and filtering. A total of 269 OTUs were generated from the obtained sequences, with a GC content of 56.5%. After chimeric removal and filtering, the ITS libraries seem to have covered a significant portion of the fungal variety in the samples utilized in this research. This indicated that the sequence data correctly mirrored the structural makeup of the endophytic fungus samples. Utilizing the UNITE fungal ITS database, which is validated by QIIME, we classified every OTU to the species level, exposing variations in the mycobiome.

3.5. Examining Operational Taxonomic Units

Figure 5 shows the fraction of fungal genera’ abundances at the phylum and genus levels. The fungal community presented in _H. binata_ was identified through a culture-independent method and attributed to 10 phyla, 37 classes, 133 families, and 189 genera. The phyla include Ascomycota (71.3%), Basidiomycota (17.8%), Mortierellomycota (2.2%), Mucromycota (1.4%), Olpidiomycota (1.4%), Neocallimastigomycota (1.1%), Rozellomycota (1.1%), Glomeromycota (1.1%), and Zoopagomycota (0.3%). At the genus level, _Tilletia_ (26%) was the most dominant, followed by _Ramichloridium_ (23%), _Dissoconium_ (13%), _Aspergillus_ (10%), _Penicillium_ (6%), _Cladosporium_ (5%), _Pichia_ (4%), _Talaromyces_ (4%), and _Fusarium_ (4%). These results suggest that the abundance of the culturable endophytic communities was similar. Furthermore, a wide range of microfungi were included in all OTUs with a sequence similarity of ≥97% that had sequences that were made publicly available with species identification (Supplementary Table S2). The Ascomycota phylum was found to be predominant, and one percent of the fungi belong to unidentified and unclassified phyla. As such, the sequences are not easily allocated to any recognized category using the present taxonomy references database.

![Figure 5. Phylum- and genus-level endophytic fungal community composition and relative abundance (column length reflects genus percentage and column color reflects fungal genus color) in the sample. For sequences that could not be categorized well into any existing category, the label “unassigned” was applied.](image-url)
3.6. Rarefaction Analysis and Richness Estimates

As the curves began to approach saturation, suggesting that the chosen sequence data accurately matched the fungal abundance of these samples, the rarefaction curves demonstrated that the sequencing diligence was more comprehensive in covering the fungal variety (Figure 6). To put it another way, the rarefaction curve indicated that almost every species in the community of the samples was covered by the sequencing data. Conversely, the endophytic microbial component distributions in *H. binata* were found at several taxonomic levels, and the endophytic community complexity was estimated using various fungal diversity indices, such as Shannon and Simpson (Table 3).

![Rarefaction Curve](image)

**Figure 6.** Endophytic fungal rarefaction curves of the plant sample.

<table>
<thead>
<tr>
<th>Culture-Dependent Approach</th>
<th>Culture-Independent Approach</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungal Isolates</td>
<td>Shannon Diversity Index</td>
</tr>
<tr>
<td><em>H. binata</em></td>
<td>25</td>
</tr>
</tbody>
</table>

Table 3. The community composition and richness of endophytic fungal isolates from the leaves of *H. binata*.

3.7. Culture-Dependent versus Culture-Independent

Culture-dependent and culture-independent methods are employed in studying endophytic fungal diversity, each offering unique insights into community structure and composition. Comparing the number of independent isolated fungus species (25 total) obtained in this work to the total number of OTUs (269 total) found via sequencing would seem to validate this approach. However, several of the proportionately dominant taxa revealed by the sequencing method were really represented by isolates. In general, the ITS sequences of mycobial endophytes found using both techniques yielded a direct comparison that showed 97% similarity or higher between all fungal taxa found using the conventional methods and even fungal OTUs found using the culture-independent methods.

The fungal community analysis using a culture-independent and culture-dependent approach revealed a dominant Ascomycota phylum, accounting for 71.3% of the community. A small fraction, approximately 1%, could not be classified under known phyla and were categorized under the “uncultured” group. This highlights the complexity and di-


versity of the fungal community and the limitations of current taxonomic classifications in fully characterizing microbial populations. Both methods identified a number of frequently found fungus taxa, including Aspergillus, Nigrospora, and Fusarium. In the mycobiome, there are several highly or often discovered genera that were missed using the conventional approach. These include Dissoconium, Pichia, Lasiodiplodia, Cladosporium, Tilletia, Ramichloridium, and Trichothecium. The common and distinct fungi among the samples were determined using a Venn diagram. A total of 189 fungal taxa have been identified in the comparative study using culture-independent approaches. Furthermore, 17 fungal genera were discovered using culture-dependent approaches. Remarkably, the overlap between the two methods revealed that 17 fungal genera were shared by both datasets. This convergence highlights the value of using alternative methodologies to thoroughly investigate fungal communities by indicating a degree of consistency between culture-independent and culture-dependent methods in capturing a fraction of the fungal diversity (Figure 7). Comparing the two approaches, the culture-dependent method provided detailed insights into cultivable fungal species, allowing for precise morphological characterization and taxonomic classification. However, it may underestimate fungal diversity due to limitations in cultivability. On the other hand, the culture-independent approach offered a broader view of fungal diversity by detecting non-cultivable taxa, albeit with challenges in taxonomic assignment. Integrating both methods can enhance the comprehensive analysis of endophytic fungal communities, providing a more holistic understanding of their structure and composition.

![Figure 7. A column graph and a Venn diagram depicting the total count of endophytic fungi obtained through culture-dependent and culture-independent methods. (A) Show the total number of fungal isolates from H. binata that were acquired by culture-dependent and culture-independent methods. (B) Genomic level comparison utilizing culture-dependent and culture-independent methods.](image)

4. Discussion

Endophytic fungi are commonplace within numerous plant tissues and are essential to the health and defense systems of plants. Microbial colonization of plant tissues acts as a barrier to prevent pathogen invasion, increasing the resistance of the host plant [27–29]. Moreover, endophytic fungi are renowned for their production of bioactive metabolites with diverse pharmaceutical properties, including antiviral and anticancer activities [30,31]. Recent advancements in genomics and metagenomics have revolutionized our understanding of fungal diversity and metabolic potential, facilitating the discovery of novel genes and metabolic compounds [32,33].
Traditionally, culture-dependent methods have been employed to isolate and identify endophytic fungi based on morphological and phenotypic characteristics [34,35]. However, limitations inherent to these methods have spurred the development of culture-independent techniques, such as metagenomics, enabling the direct examination of microbial communities from environmental samples [36,37]. By integrating both culture-dependent and culture-independent approaches, researchers can achieve a more comprehensive understanding of fungal diversity, revealing species that may elude traditional cultivation methods [38]. This integrated methodology not only enhances fungal species identification but also facilitates in-depth investigations into their ecological roles and potential applications across various industries, including biotechnology, agriculture, and medicine.

The study’s findings shed light on important soil properties and nutrient content for H. binata’s development. A soil with a high organic carbon content and a slightly alkaline pH is ideal for plant development. Soil micronutrient availability and a balanced nitrogen-phosphorus-potassium (NPK) ratio suggest the soil may maintain healthy growth despite moderate saline levels. Results from a study by [39] were similar. The two locations that were examined are located in an area that has a history of Pb-Zn mining dating back over 300 years. Pb, Zn, and Cd were significantly present in the soils and plants of both locations throughout the sample period. Soils from the slag heap had a greater level of HMs than those from the wasteland, on average. Soil organic matter, total nitrogen, phosphorus, potassium, accessible potassium, and hydrolyzable nitrogen were all substantially reduced in the slag heap soils compared to the wasteland soils, with the exception of available phosphorus. To maximize H. binata’s development in the research region, however, more research into its unique needs with regard to soil salinity, pH, and nutrient levels is required. Insightful management techniques may be built upon these results, which add to our knowledge of soil fertility and help ensure the long-term viability of H. binata populations.

The study represents a pioneering investigation into the endophytic fungal population inhabiting H. binata leaves, utilizing both traditional culture-dependent methods and modern culture-independent techniques. Precise taxonomic identifications were derived from DNA analysis, enabling a comprehensive comparison of taxa obtained from both methods using the same samples. This study marks the first report on the identification and isolation of endophytic fungi within H. binata. The culture-dependent method yielded 60 fungal isolates from symptomless leaves, subsequently identified into 17 morphotypes encompassing diverse taxa, including Fusarium, Phoma, Didymella, Talaromyces, Sphaeropsis, Curvularia, Rhytidhysteron, Diaporthe, Phyllosticta, Aspergillus, Sordaria, Nigrospora, Xylaria, Pestalotiopsis, Daldinia, and Pseudopithomyces genera. Comparable findings were noted in previous research by [40], who isolated 15 endophytic fungi from mature E. pachyclada leaves, identifying species such as Penicillium crustosum, Penicillium chrysogenum, Penicillium commune, Penicillium corylophilum, Alternaria infectoria, Alternaria alternata, Alternaria tenuissima, Aspergillus flavus, and Aspergillus niger. Additionally, previous studies by [41,42] have identified endophytic fungi from medicinal plants and explored the importance of molecular phylogeny in fungal taxonomy, highlighting the challenges associated with DNA barcoding methods such as ITS sequencing due to sequence diversity within and between species. For phylogenetic and taxonomic placement analyses of fungi, the protocol of choice has been the sequencing analysis of the ITS rRNA region. However, building a reliable and repeatable phylogenetic connection is hampered by the sequence diversity of the ITS1 region across species belonging to the same genus [43]. For fungus, the internal transcribed spacer region is the most chosen DNA barcode. But in certain fungal groupings, the inter-and intra-specific distances in ITS sequences differ greatly; as a result, it is not a completely trustworthy method for distinguishing between species [44].

A common core secondary structure of the nuclear ribosomal internal transcribed spacer 1 (ITS1) region in fungi was proposed by Koetschan et al. [45], using a helix-wise folding approach in conjunction with ITS1 sequence annotation based on the Hidden Markov Model. This innovative technique facilitated automated sequence-structure-based taxonomy of ITS1 for environmental fungi and incorporated structural data into phylo-
genetic analysis. Their findings suggested that ITS2 secondary structure prediction could serve as a valuable tool for species delimitation within fungal communities. Comparative analysis indicated similar taxonomy preferences between ITS and ITS2 regions in fungi, with ITS2 demonstrating potential as a standalone marker. Phylogenetic reconstruction utilizing ITS2 sequences enabled the identification of fungal species diversity within *H. binata*, revealing insights into genetic diversity and evolutionary relationships among fungal isolates. Specific fungal species, including *Curvularia lunata*, *Nigrospora lacticolonia*, and *Daldinia eschscholtzii*, were identified, along with 25 isolates encompassing diverse fungal species within *H. binata*. Additionally, [24] discovered distinct helix patterns in isolates from various genera, with some exhibiting a 3-helix pattern and others a 4-helix pattern, highlighting structural variations across species. The ITS2 ribotype findings we obtained are consistent with previous research [22,46] on the genera *Colletotrichum*, *Fusarium*, *Xylaria*, *Aerobasidium*, *Diaporthe*, and *Aspergillus*. Ref. [47] explored the utility of ITS2 molecular morphometrics for delineating Ascomycota species, emphasizing the effectiveness of this approach in elucidating fungal species boundaries and potentially easing challenges associated with molecular characterization. Integration of secondary structure information with primary sequencing data significantly enhances species identification within plant genera and sheds light on their evolutionary history. Furthermore, phylogenetic analysis of ITS2 at higher taxonomic levels leverages the conservative secondary structure qualities of ITS2 and aids in nucleotide sequence evolutionary studies, contributing to improved structure prediction and taxonomic understanding.

The diversity of fungal communities in various habitats, such as soils, plants, and environmental samples, has been thoroughly investigated using cultivation-independent methods [48–50]. When studying fungal species diversity, high-throughput sequencing technologies were able to identify more fungi than the culture-based technique would have in the past. As much as one percent of all microorganisms may be cultivated, according to research by [51]. This study employed a metagenomics approach to comprehensively characterize the endophytic fungal community inhabiting *H. binata*, leveraging the power of culture-independent methods to elucidate its taxonomic composition and diversity. Through high-throughput Illumina MiSeq sequencing analysis, a substantial dataset comprising 69,570 high-quality sequences was generated, leading to the identification of 269 Operational Taxonomic Units (OTUs). The observed GC content of 56.5% further confirmed the typical fungal DNA composition, enhancing the reliability of the obtained results. The utilization of paired-end Illumina sequencing in similar studies, such as that by [52], has provided comparable insights into fungal diversity within endophytic communities, reinforcing the robustness of the methods employed. Classification of OTUs to the species level using the UNITE fungal ITS database revealed a rich and diverse fungal community associated with *H. binata*, spanning 10 phyla, 37 classes, 79 orders, 133 families, and 189 genera. The dominance of the Ascomycota phylum, followed by Basidiomycota, underscores the complexity of the fungal community inhabiting *H. binata*. These findings align with similar research on endophytic fungal communities in related plant species, as demonstrated by [53], analyzed the endophytic fungal assemblages in *H. contortus* shoots and roots using a multiscale approach. *H. contortus*’s endophytic fungal flora includes 4 phyla, 11 classes, 17 orders, 21 families, and 18 genera in the shoot sample and 10 classes, 18 orders, 22 families, and 20 genera in the root sample. This suggests that both sections are host to a diverse range of fungal endophytic species. At the genus level, *Tilletia* emerged as the most dominant genus, followed by *Ramichloridium*, *Dissoconium*, *Aspergillus*, *Penicillium*, *Cladosporium*, *Pichia*, *Talaromyces*, and *Fusarium*. The presence of these genera indicates a diverse assemblage of endophytic fungi colonizing *H. binata* leaves. Moreover, the detection of OTUs with high sequence similarity to publicly available sequences further validates the taxonomic assignments and highlights the presence of various microfungi within the endophytic community. However, because of advancements in high-throughput sequencing techniques, an increasing number of endophytic fungi have been identified, bringing the total number of fungal species to 5.1 million. The identification and traits
of the majority of fungi are still unknown, nevertheless, since only around 10,000 species have been studied and categorized [54]. In a comparable way, our research [55] found that ten of the twenty-two most common fungal OTUs could be assigned to specific genera. *Phyllosticta*, *Talaromyces*, *Aspergillus*, *Aporospora*, *Cladosporium*, *Alternaria*, *Fusarium*, *Penicillium*, and *Allophaeosphaeria* were among the fungi that were identified. The majority of these endophytic fungi, like *Fusarium*, are often found in nature and may be either harmful pathogens or endophytes of many different types of crops [56].

Our use of Illumina sequencing has allowed us to get some insight into the fungal endophytes in this medicinal plant, and we have also detected a number of fungi that have not yet been classified or recognized, suggesting the presence of novel microbial resources. Rarefaction analysis and richness estimates provided insights into the comprehensiveness of the sequencing effort, demonstrating saturation of rarefaction curves and indicating adequate coverage of fungal diversity within the sampled *H. binata* population. The results were in line with those of the study by [57], which established that all samples had reached an equilibrium level in terms of sequencing depth and that taxonomic designations were accurate. Based on the numbers of OTUs, diversity, and coverage indices, [2] demonstrated that fungal sequence diversity was compared in each sample. Fungal communities in the subterranean regions (rhizosphere and root) are exceedingly diverse, according to the diversity index that was calculated for each possible combination of microhabitats. Furthermore, research showed that the fungal endophyte biodiversity in the above-ground components (leaves and stems) is not very great. Endophytic fungal species were most reliably identified in the root sample, rhizosphere, leaves, and stems, according to the Chao 1 and Shannon indices. The presence of a varied array of fungal species in the rhizosphere was shown using Simpson’s diversity index. It is clear that the stems have the lowest diversity indices throughout all parts of the sample, which indicates that the community’s phylotypes are most unevenly distributed. Based on this, our study also estimated the fungal diversity indices, such as the Shannon and Simpson indices, which underscore the complexity of the endophytic microbial community inhabiting *H. binata*, indicating a diverse assemblage of fungal taxa.

The investigation of endophytic fungal diversity often employs both culture-dependent and culture-independent methods, each offering distinct advantages in revealing community structure and composition. In this study, comparing the results obtained from these two approaches provides valuable insights into the fungal community associated with the host plant, *H. binata*. The culture-dependent method yielded a total of 25 independent, isolated fungal species, whereas the culture-independent method identified 269 Operational Taxonomic Units (OTUs) through sequencing. This discrepancy underscores the complementary nature of these techniques, with culture-dependent methods providing detailed insights into cultivable fungal species, including precise morphological characterization and taxonomic classification. According to research by [58], most genera found using the culture-independent approach are saprotrophs, while most genera found using the conventional method are phytopathogens. Although in six instances the identification is only up to the genus level owing to a data shortage, they were able to identify several taxa to the species level using the conventional strategy that combined morphological and genetic methods. However, it is important to note that several proportionately dominant taxa revealed by sequencing were also represented by isolates, indicating a degree of consistency between the two methods in capturing fungal diversity.

The fungal community analysis revealed a dominant Ascomycota phylum, comprising 71.3% of the community, with approximately 1% remaining unclassified under known phyla. This highlights both the complexity and diversity of the fungal community associated with *H. binata*, as well as the limitations of current taxonomic classifications in fully characterizing microbial populations. Both methods identified commonly found fungal taxa such as *Aspergillus*, *Nigrospora*, and *Fusarium*. However, the culture-independent approach unveiled additional frequently encountered genera that were missed using the conventional approach, including *Dissoconium*, *Pichia*, *Lasiodiplodia*, *Cladosporium*, *Tilletia*,...
**Ramichloridium**, and **Trichothecium**. The comparison of common and distinct fungi among the samples using a Venn diagram revealed a total of 189 fungal taxa identified through culture-independent approaches, with 17 fungal genera discovered using culture-dependent methods. Remarkably, 17 fungal genera were shared by both datasets, indicating a degree of consistency between culture-independent and culture-dependent methods in capturing a fraction of fungal diversity. Our findings were in line with those of [52], who also found that most of the OTUs were consistent between culture-dependent and culture-independent methods of detection, with the exception of **Aspergillus** (which was more common in the former but less so in the latter) and **Cadophora** sp. (a fungal taxon that was commonly found in the latter but not in the former). Because **Aspergillus** sp. may grow rapidly, even at low concentrations in the tissue samples, this finding might be due to their presence. So, they may proliferate rapidly and yield many isolates in the culture-dependent method, but they would go undetected in the next generation of sequencing.

Combining effective high-resolution phylogenetic fingerprinting with high-throughput culture methods allows us to recover previously unknown species of microorganisms, allowing us to discover and get certain unique or difficult-to-cultivate microbes [59]. The functional information of the fungal OTUs derived from culture-independent approaches may be better understood by comparing conventional and culture-independent data [58]. Integrating both culture-dependent and culture-independent methods enhances the comprehensive analysis of endophytic fungal communities. While culture-dependent methods provide detailed insights into cultivable species, they may underestimate fungal diversity due to limitations in culturability. Conversely, culture-independent approaches offer a broader view of fungal diversity by detecting non-cultivable taxa, albeit with challenges in taxonomic assignment. By combining these methodologies, researchers can gain a more holistic understanding of endophytic fungal community structure and composition, contributing to our knowledge of fungal ecology and its implications for plant health and ecosystem dynamics. Overall, this study has demonstrated the promise of combining both methods in a single study to connect NGS datasets with morphologically and phylogenetically identified fungal isolates from culture. This complementary approach allows for accurate taxonomic labeling of fungal endophytes found using NGS, which is important because NGS studies without reference cultures can be misleading and a lot of the data from the past needs to be viewed critically.

### 5. Conclusions

This study explores the diversity of endophytic fungal communities within *H. binata* leaves using both culture-dependent and culture-independent methods. Culture-dependent methods allowed for the isolation and identification of cultivable fungal species, while culture-independent techniques allowed for the detection of non-cultivable taxa. The study found that endophytic fungi play a crucial role in plant health and defense mechanisms in the presence of the Ascomycota phylum. The identification of bioactive metabolites produced by these fungi also suggests their potential pharmaceutical applications in medicine and biotechnology. The study used high-throughput sequencing technologies to bridge the gap between NGS datasets and taxonomically identified fungal isolates, ensuring accurate taxonomic labeling and enhancing the reliability of fungal community studies. The study also highlighted the complementary nature of these techniques in elucidating fungal ecology. The combined use of culture-dependent and culture-independent methods holds promise for advancing our understanding of fungal ecology and its implications for plant health and ecosystem dynamics. Future research should continue to use both methodologies to unravel the complex relationships between endophytic fungi, their host plants, and the environment.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/microbiolres15020053/s1, Figure S1: ITS2 Secondary structure predicted phylogenetic tree of H. binata; Table S1: Compensatory base changes (CBC) analysis in the sequences of H. binata. Table S2: Culture-independent molecular approaches reveal the diversity of fungal endophytic communities in leaves of H. binata.

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