

Journal of
Fungi

Fungal Endophytes in Plants

Edited by
Gary A. Strobel

Printed Edition of the Special Issue Published in *JoF*

Fungal Endophytes in Plants

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Special Issue Editor

Gary A. Strobel

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Special Issue Editor

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This is a reprint of articles from the Special Issue published online in the open access journal *Journal of Fungi* (ISSN 2309-608X) in 2018 (available at: http://www.mdpi.com/journal/jof/special-issues/fungal_endophytes)

For citation purposes, cite each article independently as indicated on the article page online and as indicated below:

LastName, A.A.; LastName, B.B.; LastName, C.C. Article Title. *Journal Name Year, Article Number,*
Page Range.

ISBN 978-3-03897-246-4 (Pbk)

ISBN 978-3-03897-247-1 (PDF)

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Contents

About the Special Issue Editor	vii
Gary Strobel	
Special Issue: Fungal Endophytes in Plants	
Reprinted from: <i>J. Fungi</i> 2018, 4, 104, doi: 10.3390/jof4030104	1
Tyler Maxwell, Richard G. Blair, Yuemin Wang, Andrew H. Kettrig, Sean D. Moore, Matthew Rex and James K. Harper	
A Solvent-Free Approach for Converting Cellulose Waste into Volatile Organic Compounds with Endophytic Fungi	
Reprinted from: <i>J. Fungi</i> 2018, 4, 102, doi: 10.3390/jof4030102	3
Joël E. T. Ateba, Rufin M. K. Toghueo, Angelbert F. Awantu, Brice M. Mbanning, Sebastian Gohlke, Dinkar Sahal, Edson Rodrigues-Filho, Etienne Tsamo, Fabrice F. Boyom, Norbert Sewald and Bruno N. Lenta	
Antiplasmoidal Properties and Cytotoxicity of Endophytic Fungi from <i>Symphonia globulifera</i> (Clusiaceae)	
Reprinted from: <i>J. Fungi</i> 2018, 4, 70, doi: 10.3390/jof4020070	18
Yuemin Wang and James K. Harper	
Restoring Waning Production of Volatile Organic Compounds in the Endophytic Fungus <i>Hypoxyylon</i> sp. (BS15)	
Reprinted from: <i>J. Fungi</i> 2018, 4, 69, doi: 10.3390/jof4020069	27
Dong-Hui Yan, Xiaoyu Song, Hongchang Li, Tushou Luo, Guiming Dou and Gary Strobel	
Antifungal Activities of Volatile Secondary Metabolites of Four <i>Diaporthe</i> Strains Isolated from <i>Catharanthus roseus</i>	
Reprinted from: <i>J. Fungi</i> 2018, 4, 65, doi: 10.3390/jof4020065	39
Yu-Ling Huang, Naupaka B. Zimmerman and A. Elizabeth Arnold	
Observations on the Early Establishment of Foliar Endophytic Fungi in Leaf Discs and Living Leaves of a Model Woody Angiosperm, <i>Populus trichocarpa</i> (Salicaceae)	
Reprinted from: <i>J. Fungi</i> 2018, 4, 58, doi: 10.3390/jof4020058	55
Andreea Cosoveanu, Samuel Rodriguez Sabina and Raimundo Cabrera	
Fungi as Endophytes in <i>Artemisia thuscula</i> : Juxtaposed Elements of Diversity and Phylogeny	
Reprinted from: <i>J. Fungi</i> 2018, 4, 17, doi: 10.3390/jof4010017	70
Sunil K. Deshmukh, Manish K. Gupta, Ved Prakash and M. Sudhakara Reddy	
Mangrove-Associated Fungi: A Novel Source of Potential Anticancer Compounds	
Reprinted from: <i>J. Fungi</i> 2018, 4, 101, doi: 10.3390/jof4030101	91
Sunil K. Deshmukh, Manish K. Gupta, Ved Prakash and Sanjai Saxena	
Endophytic Fungi: A Source of Potential Antifungal Compounds	
Reprinted from: <i>J. Fungi</i> 2018, 4, 77, doi: 10.3390/jof4030077	130
Andreea Cosoveanu and Raimundo Cabrera	
Endophytic Fungi in Species of <i>Artemisia</i>	
Reprinted from: <i>J. Fungi</i> 2018, 4, 53, doi: 10.3390/jof4020053	172

- Eyob Chukalo Chutulo and Raju Krishna Chalannavar**
Endophytic Mycoflora and Their Bioactive Compounds from *Azadirachta Indica*:
A Comprehensive Review
Reprinted from: *J. Fungi* 2018, 4, 42, doi: 10.3390/jof4020042 190
- Brian R. Murphy, Fiona M. Doohan and Trevor R. Hodkinson**
From Concept to Commerce: Developing a Successful Fungal Endophyte Inoculant for
Agricultural Crops
Reprinted from: *J. Fungi* 2018, 4, 24, doi: 10.3390/jof4010024 202
- Gary Strobel**
The Emergence of Endophytic Microbes and Their Biological Promise
Reprinted from: *J. Fungi* 2018, 4, 57, doi: 10.3390/jof4020057 213

About the Special Issue Editor

Gary A. Strobel was born and raised in Massillon, Ohio. He completed a B.S. degree at Colorado State University in 1960, and a PhD at the University of California, Davis in 1963. Since that time, he has been on the faculty of Montana State University. His research and academic interests have centered on microbe–higher plant relationships. More recently, he has begun to examine endophytic fungi and bacteria for their novel bioactive compounds and their unique biology. He has lectured at over 400 institutes and universities worldwide. Moreover, he has published over 360 articles in scientific journals and holds over 50 USA and international patents. He has received numerous awards including an NIH Career Development Award, the Wiley Award, Special Recognition from the Royal Nepal Chemical Society, and the MSU-VP Award for Technology and Science. He is an elected fellow of the American Academy of Microbiology and was elected as a full fellow of the Explorer's Club of the World. Also, he has been awarded a senior BARD fellowship from the government of Israel and a senior CSIRO fellowship from the Australian Government. From 1979 to 2000 he was chief of the Montana NSF EPSCOR program, which encourages and promotes science at all levels of society. In 2012 he was presented the E.O Wilson award for his work on endophyte biodiversity. He was also elected a member of the National Academy of Inventors in 2014 and was presented with the Lowell Thomas Award in 2015 by the Explorer's Club of the World.

Editorial

Special Issue: Fungal Endophytes in Plants

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Received: 24 August 2018; Accepted: 28 August 2018; Published: 1 September 2018

Once in a while scientific developments occur that represents a complete shift in the paradigm of a scientific discipline. The emerging field of endophyte biology is an example of this phenomenon. One hundred years ago the search was on to describe the pathology and causal agent of every plant disease on the planet. Every microbe associated with a plant was suspect. Those not having a pathological etiology were discarded as uninteresting and not worthy of attention. By the mid-twentieth century some investigators began to systematically isolate, describe and name the non-pathogens of plants and they were designated as endophytes if they were found in the living tissues of the plants. Soon it was learned that some of these organisms associated with certain grasses were the cause of abortions and death in livestock. The name endophyte took on a whole new meaning as an unwanted biological case. With the advent of the discovery of certain endophytes making the drug-taxol there was another shift in the thinking and potential importance of these microorganisms. The field of endophyte biology took on a whole new persona. Soon it was realized that there are literally thousands of endophytes making hundreds of compounds that may have uses in industry, medicine and agriculture. Further studies have revealed that plants have a microbiome just as humans and other animal species and that this assemblage of microbes has an important role in the ultimate health and survival of plant species. We are just now putting resources and time into the study and ultimate utilization of these microbes to the benefit of mankind. Some endophytes seem to provide protection to the plant from other microbes, insects and herbivores, others promote growth while others are known that allow the plant to withstand environmental stresses such as heat and drought. The microbiome of plants mainly represents fungi but bacterial species also occur with some regularity.

My career began as a student of forestry at Colorado State University in the mid-1950s. I soon learned that this discipline was not organized to address the fundamental questions of science so I became a botany major with a minor in chemistry. Eventually, I went on the UC-Davis in California for a PhD in plant pathology. Again, I felt that chemistry, biological chemistry and microbiology held the answers to the future of plant biology. I became an assistant professor of plant pathology at Montana State University in 1963. I focused on questions relating to the fundamental aspects of plant disease physiology and biochemistry. Phytotoxins, plant receptors and other major topics concerning the biochemistry of plant disease were the targets of my time and energies. In the early 90s we successfully isolated and characterized taxol from a novel endophytic fungus associated with a pacific yew tree growing in Northern Montana. Soon we began to examine endophytes from forests all over the planet. Those trips took us to some of the most beautiful, enchanting, and challenging places and those experiences have left an indelible mark on my memory. By now some of these discoveries have found their way into the real world such as the use of *Muscador albus* as a soil treatment as a replacement for the use of methyl bromide. This endophytic organism has been approved by the US-EPA and is now on the market. Furthermore, the volatile antimicrobial compounds of other *Muscador* isolates will soon appear in the marketplace.

It has been an honor to me to have been asked to serve as the editor of this special volume on endophytes. It caps a long career dedicated to an understanding of how microbes associate with plants and how we can utilize this information to improve crop production as well as our own lives. Certainly, the efforts of the *Journal of Fungi* staff including managing editor Aimar Xiong, assistant

editors Ashlynn Wang and Monica Cui are acknowledged as being helpful while being at the same time being professional and courteous. As someone once said—"If I had to do this career all over again- I would." The strange and interesting world of fungi offers to us a never-ending challenge of excitement and discovery.

Author Contributions: G.S. prepared the manuscript.

Funding: None.

Conflicts of Interest: The author declares no conflict of interest.



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Article

A Solvent-Free Approach for Converting Cellulose Waste into Volatile Organic Compounds with Endophytic Fungi

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Received: 25 June 2018; Accepted: 23 August 2018; Published: 26 August 2018

Abstract: Simple sugars produced from a solvent-free mechanocatalytic degradation of cellulose were evaluated for suitability as a growth medium carbon source for fungi that produce volatile organic compounds. An endophytic *Hypoxyylon* sp. (CI-4) known to produce volatiles having potential value as fuels was initially evaluated. The growth was obtained on a medium containing the degraded cellulose as the sole carbon source, and the volatile compounds produced were largely the same as those produced from a conventional dextrose/starch diet. A second *Hypoxyylon* sp. (BS15) was also characterized and shown to be phylogenetically divergent from any other named species. The degraded cellulose medium supported the growth of BS15, and approximately the same quantity of the volatile compounds was produced as from conventional diets. Although the major products from BS15 grown on the degraded cellulose were identical to those from dextrose, the minor products differed. Neither CI-4 or BS15 exhibited growth on cellulose that had not been degraded. The extraction of volatiles from the growth media was achieved using solid-phase extraction in order to reduce the solvent waste and more efficiently retain compounds having low vapor pressures. A comparison to more conventional liquid–liquid extraction demonstrated that, for CI-4, both methods gave similar results. The solid-phase extraction of BS15 retained a significantly larger variety of the volatile compounds than did the liquid–liquid extraction. These advances position the coupling of solvent-free cellulose conversion and endophyte metabolism as a viable strategy for the production of important hydrocarbons.

Keywords: endophytic fungi; Mechanocatalysis; cellulose degradation; volatile organic compounds; myco-diesel; *Hypoxyylon*

1. Introduction

The endophytic fungi are organisms that colonize the tissue of living plants. In most cases, this relationship is asymptomatic and may even provide benefits to plants [1]. Endophytes have been studied extensively and found to produce a remarkable variety of natural chemical products [2]. While much of the interest has focused on bioactive compounds, the production of other important compounds has also been reported. A recent noteworthy discovery is that certain endophytes can produce hydrocarbons that have the potential to be used as fuels or fuel additives [3]. These products have been compared to diesel fuel and even described as “myco-diesel”, because they include

compounds normally associated with diesel fuel. Over the past decade, interest in fungi producing volatile organic products with the potential for use as fuels has increased, and several studies have identified potentially useful fungi [4–10]. Related work has also identified fungi producing volatile products but has not focused on their potential usefulness as fuels [11–25].

The availability of hydrocarbon fuels from fungi complements fuel products produced by other organisms. For example, certain algae produce aliphatic fatty acids and considerable effort has been expended into developing these into viable biofuels [26]. Likewise, yeast fermentation has been prominently utilized to convert carbohydrates from corn into ethanol for fuel [27]. In general, fungal products contain a more complex variety of volatile compounds than either algae or yeast, including ketones, esters, alcohols, and a remarkable variety of hydrocarbon products. All of these biofuels complement more conventional fuels and thus represent important pathways worthy of exploration given the current interest in developing alternative fuels. However, one of the concerns that exists when producing hydrocarbon fuel from fungi is that they require a refined carbohydrate source (e.g., sucrose) in their diet. There has been debate regarding the suitability of devoting carbohydrates to fuel production. A solution to this dilemma would be to find an alternative food source for the fungi.

Recently, a “green chemistry” mechanocatalytic method has been reported that allows cellulose waste products to be converted into simple carbohydrates [28]. This process involves ball milling performed in the solid state and is thus entirely solvent free and capable of rapidly producing large quantities of carbohydrates (see experimental). The major water-soluble products from this process have been shown to be glucose, fructose, and levoglucosan. No oligosaccharides larger than dimers survive the milling, even after short processing times (e.g., 30 min) [28]. This process has been successfully demonstrated using a remarkable variety of cellulose-based feedstock materials from plants (Table 1) and includes numerous materials normally regarded as unusable waste such as orange peels, cherry pits, coffee grounds, and discarded newspaper.

At the present time, however, it has not been demonstrated that fungi can actually grow on the carbohydrates created from the cellulose breakdown process. The aim of this manuscript is to demonstrate that carbohydrates produced from this solvent-free degradation process are a suitable carbon source for fungal growth and that the volatile products produced from the degraded cellulose closely match compounds produced from a more conventional diet. In the following, sugars from only one cellulose source (oak) are tested. The results from other materials in Table 1 are expected to give similar results, as it has been demonstrated that cellulose from various sources consistently breaks down into simple sugars [28].

Table 1. Waste materials containing cellulose that can be converted into simple sugars.

Feedstock	Percent Hydrolyzed ^a
Cherry pit	95.7
Flint corn kernal	93.4
St. Augustine grass	92.5
Oat	90.3
Orange peel	85.0
Corn cobs	81.5
Bamboo	75.1
Cedar	74.0
Red Oak	72.4
Maple	72.0
Douglas Fir	71.1
<i>Nannochloropsis</i>	69.2
Aspen	68.0
Poplar	66.9
Yellow pine	65.3

Table 1. Cont.

Feedstock	Percent Hydrolyzed ^a
Wheat	65.0
<i>Miscanthus</i> grass	64.7
White pine	64.4
Mixed yard waste	58.1
Switch grass	57.9
Hickory	55.9
Paper, newsprint	54.7
Flint corn stover	52.1
Banana leaf	52.0
Big blue stem grass	50.1
Little blue stem grass	48.9
Coffee grounds	45.2

^a This value represents hydrolysis of the holocellulose present in the material.

2. Materials and Methods

The cellulose employed in this study to create the simple sugars was obtained from water oak (*Quercus nigra*) sawdust sourced from a local sawmill. The oak was dried at room temperature to a moisture content of <10% and cut into 2 cm or smaller pieces. Delaminated kaolinite (Kaopaque 10, IMERYS) was used as received.

The mechanical processing of cellulose employed 8000M and 8000D mixer mills (SPEX Certiprep, Metuchen, NJ, USA). Two grams of a 1:1 mixture of the kaolinite clay catalyst and biomass source were processed for two hours in 65-mL vials (1.5" ID × 2.25" deep) made of 440C steel, utilizing three 0.5-inch diameter balls composed of the same material as the milling vial. Energy was applied in 30-min intervals with 30 min of cooling time to minimize the effects of frictional heating. Hydrolysis of hemicellulose and cellulose (holocellulose) was monitored gravimetrically. Conversion of holocellulose to water-soluble oligosaccharides was determined by stirring 0.1 g of the reaction mixture in 30 mL of water. The production of water-soluble products was measured by filtration through a 47-mm diameter Whatman Nuclepore® track etched polycarbonate membrane filter with a pore size of 0.220 µm. The residue was dried in a 60 °C oven for 12 h and then weighed.

The potato dextrose broth and agar were purchased from Becton Dickinson. Ammonium sulfate, acetonitrile, ethyl acetate, anhydrous magnesium sulfate, and methanol were purchased from Fisher Scientific. A sample of 1,8-cineole was obtained from TCI chemicals (Portland, OR, USA). The yeast nitrogen base was purchased from Sigma Aldrich. HyperSep C-18 solid-phase extraction columns (1 g bed weight) were purchased from Thermo Scientific. Potato dextrose agar was purchased from Microtech Scientific. All the reagents were used as received.

The isolation of the *Hypoxyylon* sp., BS15, was from branch clippings of a *Taxodium distichum* (Bald Cypress) gathered near Orange City, Florida, USA. The branches were treated with 70% ethanol, flame sterilized, and then dried in a sterile laminar-flow hood. A sterile knife blade was then used to cut away the outer tissue from the clipping, and a square wedge of the inner tissue was placed on water agar. This dish was incubated, and any fungal hyphae observed growing from the sample were transferred onto separate plates of potato dextrose agar.

The potato dextrose broth was prepared by adding 2.4 g of the potato dextrose broth to 100 mL of purified water in a 500-mL Erlenmeyer flask. The flask was then sealed with aluminum foil and autoclaved for 15 min for sterilization. The fungi of interest (CI-4 or BS15) were then added to the sterile broth, and it was resealed with foil. Cellulose broth was prepared using 250 mL of purified water, 5 g of degraded cellulose, 1.5 g of ammonia sulfate, and 1.7 g of the yeast nitrogen base without amino acids. In both growth media, the fungi were then left to grow for 25 days in the lab at 20–25 °C

without stirring. Each broth was then vacuum filtered twice through Whatman Grade 4 filter paper to remove all particulates.

A control sample containing cellulose not subjected to the mechanocatalytic degradation process was prepared by adding 2 g of finely ground cellulose powder and 0.5 g of ammonium sulfate to 250 mL of distilled water. This medium was autoclaved for 15 min, and, after cooling, two separate solutions were prepared by adding CI-4 or BS15 to the liquid. This culture was allowed to grow for 2 weeks at 20–25 °C without stirring.

For the solid-phase extraction of the fungal volatile compounds, a C-18 cartridge was first washed with 4 mL of methanol and then with 4 mL of water. Filtered fungal broth (50 mL) was then passed through the column slowly under vacuum. The column was washed again with 4 mL of water to remove any contaminants and then dried by drawing air through the column for 15 min. The retained compounds were then eluted by passing acetonitrile through the column. A clear brown solution was typically recovered from this process. The eluent was then filtered with a 0.22-μm syringe filter prior to analysis.

For the liquid–liquid extractions, a total of 300 mL of the filtered fungal broth was shaken in a separatory funnel with 50 mL of ethyl acetate. The ethyl acetate was then separated from the water and dried over anhydrous magnesium sulfate. The solution was then filtered with a 0.22-μm syringe filter prior to analysis.

The gas chromatography/mass spectrometry (GC/MS) analysis for the volatile compounds was performed using a method described previously with slight modification [29,30]. An Agilent 6850 was used with a 5975CVC MS detector and a Restek Rxi-5HT capillary column (30 m × 0.25 mm, film thickness 0.25 μm). The carrier gas was ultrahigh purity helium with a one cm³/min constant flow rate and an initial column head pressure of 77 kPa. The injector split was set to 250 °C at a 20:1 split ratio with 1-μL volume per injection. The column oven temperature was programmed to 45 °C with an initial temperature hold for 1 min with a 10 °C/min ramp to 100 °C and hold for 5 min, followed by a 5 °C/min ramp to 200 °C and a hold for 5 min. The detector was set at a constant 280 °C and set to scan 30–350 m/z. The data acquisition and processing were performed on Agilent MSD ChemStation software. The identification of the compounds was made via library comparison using the National Institute of Standards and Technology (NIST, Gaithersburg, MD, USA) database.

For DNA extraction from BS15, a small sample of the fungal tissue (50–100 mg) was collected into a microcentrifuge tube from the surface of a potato dextrose agar plate after 1 week of growth at room temperature. The tissue was lysed using a FastPrep Homogenizer (MP Biomedicals, Santa Ana, CA, USA) by zirconia–silica bead beating in 1 mM of sodium dodecyl sulfate, 5 mM of EDTA, and 10 mM of Tris-HCl, pH 8.0 with 10 μg/mL RNase A. The lysate was centrifuged, and then, the DNA was purified from the supernatant by silica column binding in guanidinium thiocyanate [31].

Diagnostic gene sequences used for identification by genetic barcoding were amplified by a routine polymerase chain reaction (PCR) with Taq [32,33]. Primers ITS1-F_KYO1 and ITS4_KYO1 were used to target the internally transcribed spacers (ITS1 and ITS2) and the flanking portions of the ribosomal RNA encoding genes (SSU, 5.8S, and LSU) [34]. The protein-coding genes α-actin and β-tubulin were amplified by primers ACT-512F/ACT-783R and T1/T22, respectively [35,36]. The PCR products were visualized by agarose gel electrophoresis, similarly purified by silica column binding, and then sequenced commercially (GENEWIZ, Plainfield, NJ, USA). The sequences were deposited in GenBank under accession number MH223406 (ITS), MH465497 (actin), and MH465498 (tubulin).

The ribosomal gene sequences were analyzed with a series of BioPython-based scripts [37]. First, full-length ITS sequences were extracted via ITSx and used to locally query the UNITE+INSDC fungal database by BLAST search [38–40]. Based on these search results, relevant taxa were selected, and a list was compiled of all the unambiguous binomial species within these taxa. The corresponding UNITE records were pooled and analyzed by ITSx. For each species, a single representative full-length ITS2 record was chosen for alignment. Relevant α-actin and β-tubulin records used in alignment were retrieved from GenBank [41].

The phylogenograms were generated using MEGA software [42]. The sequences were aligned by the MUSCLE algorithm and then clustered by the maximum likelihood method with 1000 bootstrap replicates [43,44]. Both the α -actin and β -tubulin sequences were treated as protein-coding during the phylogenetic analyses, while ITS sequences were not. All other settings in MEGA were unchanged and no manual modifications were made during the alignment or clustering. The resultant phylogenograms were exported and visualized via Interactive Tree of Life (iTOL) web software [45]. The nodes were pruned on the basis of relatedness to BS15 and intra-generic species richness. The alignments and phylogenograms were deposited in TreeBase under submission number 23089.

3. Results and Discussion

3.1. Growth of the *Hypoxyylon* CI-4 on Degraded Cellulose

As an initial test of the feasibility of using carbohydrates from mechanocatalytic cellulose degradation as a fungal diet, a *Hypoxyylon* sp. was added to a growth medium consisting of the degraded cellulose as the sole carbon source (see experimental). A control sample was also prepared, having the fungus on a conventional diet of potato dextrose broth. The particular *Hypoxyylon* fungus used for this study (designated CI-4) was selected because it has been previously shown to produce a diverse variety of volatile organic hydrocarbons [46,47]. Both cultures exhibited similar fungal growth and were incubated for three weeks. The hydrocarbon fraction was extracted from the growth media using a solid-phase extraction process (see experimental). A gas chromatography/mass spectrometry (GC/MS) analysis exhibited a diverse range of volatile products, as expected from the previous study [46]. A comparison of the volatile compounds produced from each growth condition is illustrated in Figure 1 and demonstrates that the degraded cellulose material produces the same major products as a conventional carbohydrate-rich diet. A notable difference, however, between the growth media is that the amounts of the volatile compounds produced from the cellulose degradation products were roughly two–five times less than the same products produced from the potato dextrose broth.

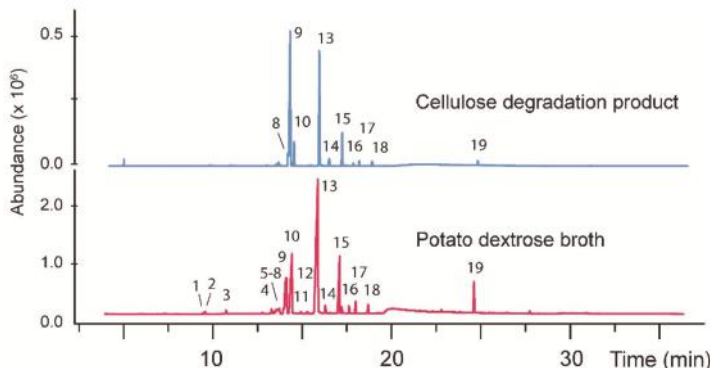


Figure 1. A gas chromatogram showing the volatile organic products produced by the fungus CI-4 growth on a conventional media (**bottom**) versus a diet containing carbohydrates produced from cellulose degradation (**top**). The nominal masses for each numbered peak are given in Table 2. In each case a control sample was also analyzed consisting of the growth medium without fungi added. This solution was processed identically to the fungi-containing samples. In each case, no peaks from the control samples corresponded to any of the peaks shown above.

The molecular masses and tentative identification of individual compounds from CI-4 were made by comparing the mass spectrum of each peak against the data in the NIST database. Although the nominal masses were obtained in all cases, most compounds were not identifiable. All the results are

summarized in Table 2. Also included in Table 2 are the results from the analysis of a second fungus (BS15, described below).

Table 2. A list of the volatile compounds produced by CI-4 or BS15 grown on either the potato dextrose broth (PD) or the degraded cellulose (DC), showing tentative compound identification where possible.

Fungus	Peak # ^a	R.T. (min)	Area (%) PD, DC ^b	Tentative Identity ^c	Mol. Mass (Da)	Qual. ^d
CI-4	1	9.58	0.1, -	Unknown	126	-
CI-4	2	9.72	0.3, -	Unknown	138	-
CI-4	3	10.88	0.5, -	Unknown	124	-
CI-4	4	12.93	0.2, -	Unknown	152	-
CI-4	5	13.43	0.8, -	Unknown	122	-
CI-4	6	13.57	0.3, -	Unknown	154	-
CI-4	7	13.68	1.4, -	3-Ethenyl-2-methylene cyclopentanecarboxylic acid	152	50
CI-4	8	14.11	0.6, 2.8	Unknown	152	-
CI-4	9	14.25	10.6, 45.7	Unknown	150	-
CI-4	10	14.57	15.5, 6.0	Unknown	154	-
CI-4	11	15.08	0.3, -	Unknown	154	-
CI-4	12	15.44	0.3, -	Unknown	152	-
CI-4	13	16.02	43.8, 2.0	Unknown	168	-
CI-4	14	16.45	1.1, 2.0	1-Acetyl-2-(1-hydroxyethyl)-cyclohexene	168	50
CI-4	15	17.24	10.4, 7.6	Unknown	150	-
CI-4	16	17.79	1.0, 0.8	Unknown	170	-
CI-4	17	18.14	1.5, 1.2	Unknown	170	-
CI-4	18	18.84	1.1, 1.1	3-Isopropoxy 5-methyl-phenol	166	61
CI-4	19	24.78	4.0, 1.0	2,3-Dimethoxy-naphthalene	188	85
BS15	1	4.82	1.1, 2.7	Furfuryl alcohol	98	72
BS15	2	5.07	9.9, 2.4	Methyl 4-oxo-2-butenoate	114	94
BS15	3	7.85	9.3, 1.7	Benzeneacetaldehyde	120	70
BS15	4	8.22	1.6, -	4-methoxy-2,5-dimethyl-3(2H)-furanone	142	77
BS15	5	8.45	36.7, 26.1	2,5-furandione dihydro-3-methylene	112	55
BS15	6	8.65	-, 21.5	Levoglucosanone	126	78
BS15	7	9.65	20.9, 12.8	2-Phenyethanol	122	86
BS15	8	12.24	3.0, -	Unknown	158	-
BS15	9	12.34	-, 20.3	5-(Hydroxymethyl)furfural	126	91
BS15	10	13.84	17.8	Unknown	86	-
BS15	11	14.56	2.5, -	Phenylacetic acid	136	75
BS15	12	16.23	3.0, -	Unknown	138	-
BS15	13	16.45	-, 3.2	Unknown	142	-
BS15	14	18.20	3.7, -	Unknown	154	-
BS15	15	20.47	-, 8.1	Unknown	162	-
BS15	16	22.73	-, 3.2	2,4-dihydroxy-3,6-dimethyl Benzoic acid, methyl ester	196	72
BS15	17	25.58	-, 2.1	Dihydro-5-(2-oxocyclohexylidene) 2(3H)-furanone	180	70
BS15	18	26.63	1.7, -	Furo [3, 4-f][1,3] benzodioxole-5,7-dione	192	65
BS15	19	26.78	2.9, -	Unknown	97	-
BS15	20	27.06	8.7, -	Unknown	127	-
BS15	21	28.04	1.8, -	Unknown	127	-

^a Peak numbers correspond to the numbering shown in Figure 1 (CI-4) or Figure 4 (BS15). ^b The labels PD and DC refer, respectively, to the potato dextrose broth and the degraded cellulose. The areas listed are the relative peak areas. ^c All the assignments of structure were made on the basis of the match to the National Institute of Standards and Technology (NIST) database. ^d Qual. refers to the highest listed quality value for the peaks that occur in both the growth media or, for the peaks that occur only in a single medium, to the value from that solution.

To verify that the volatile compounds produced from CI-4 and grown on the degraded cellulose are the result of the presence of simple sugars rather than residual cellulose, a control containing cellulose not degraded by the mechanocatalytic process was also prepared for comparison (see Materials and Methods). After two weeks of incubation on this medium, CI-4 showed no growth.

A notable difference between the compounds extracted here by the solid-phase extraction and the previous study of CI-4 [46,47] is that the solid-phase extraction failed to recover some of the early eluting peaks. As discussed below, a liquid–liquid extraction demonstrated that these compounds are,

in fact, present, and the cause of their omission from the solid-phase extraction sample is currently under investigation.

3.2. Phylogenetic Characterization of a New *Hypoxyylon* sp., BS15

Recently a second fungus producing volatile organic products was isolated from a Bald Cypress (*Taxodium distichum*) near Orange City, FL (USA). This fungus, designated BS15, was selected for study based on the serendipitous observation that compounds having a distinctive odor were produced.

The identification of BS15 involved extracting genomic DNA, amplifying and sequencing its ribosomal internally transcribed spacer regions (ITS), and then applying an improved bioinformatics analysis based on existing methods. The detection of flanking ribosomal genes in the BS15 sequence by ITSx allowed for the extraction of full-length ITS1 and ITS2 sub-sequences, a critical factor for producing alignments where gap site data is utilized in the phylogenetic analyses [48]. Independent BLAST searches using these sub-sequences to query the UNITE+INSDC database returned alignments with species exclusively of the taxonomic family *Xylariaceae*. Therefore, all public sequence records pertaining to the family *Xylariaceae* were comprehensively screened. The ITS sequences were detected by ITSx in 3443 of 3470 records from 394 unique binomial species.

A notable discrepancy regarding the naming and classification of organisms described in this work is the recent recognition of the family *Hyopxylaceae* by INSDC, whose members were previously included within *Xylariaceae* [49]. However, these records have not yet been updated in UNITE at this time. For the present work, non-*Hyopxylaceae* species were included in the alignment and clustering but pruned from the ITS phylogram with the exception of *Xylaria hypoxylon*, presented as a rooted out-group (Figure 2).

The relative richness of the full-length sequence records and the consistency in the sequence length made ITS2 a more favorable target for multiple alignment than ITS1 for the family *Xylariaceae*. An analysis of ITSx outputs revealed a bias for sequences containing the large ribosomal subunit sequence (LSU) compared with the small subunit (SSU) sequences among the UNITE records for the family *Xylariaceae*. Because the detection of these flanking ribosomal sequences is required for the full-length extraction of ITS sequences by ITSx, there were nearly twice the number of full-length ITS2 sequences ($n = 2165$) available for alignment compared with ITS1 ($n = 1212$). The sequence lengths were considerably less variable for ITS2 (SD = 5) than ITS1 (SD = 51).

Our taxonomic evaluations are consistent with other authors who found protein-coding genes more congruent with phenotypic observations than non-coding ITS sequences for *Hypoxyylon* and related genera [41]. The phylogenograms generated from the ITS2 sequences were remarkably unresolved regardless of the alignment and clustering methods, with several genera not clustered into the monophyletic groups (e.g., *Annulohypoxylon* spp., *Daldinia* spp.) (Figure 2). Both the α -actin and β -tubulin genetic analyses were able to fully resolve these taxa, albeit with fewer specimens ($n = 78$) than ITS (Figure 3). For all three genetic markers, the fungal strain BS15 was consistently clustered among *Hypoxyylon* spp. and most closely associated with *H. investiens*.

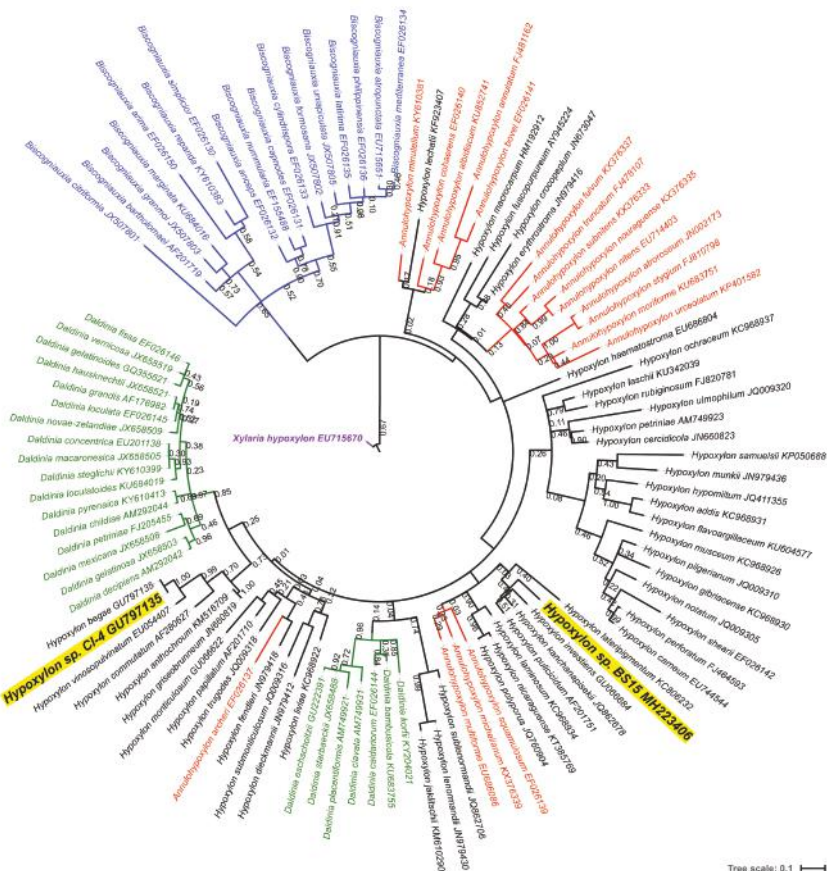


Figure 2. Phylogenetic reconstruction of *Hypoxylon* sp. BS15 and related organisms generated from maximum likelihood clustering of MUSCLE-aligned ITS2 sequences. Branch lengths are drawn to scale, representing the average number of nucleotide substitutions per site between the sequences. 121 nodes were selected for inclusion in the present figure from 331 nodes in the original phylogram. The bootstrap values at the nodes are from 1000 bootstrap iterations.

3.3. Growth of BS15 on Degraded Cellulose and Analysis of Volatile Hydrocarbons

In order to more generally evaluate the suitability of the degraded cellulose as a carbon source for fungi, BS15 was also evaluated for its ability to grow on the material. The procedure described above using two separate diets was employed with BS15 growth. The first included the degraded cellulose as the sole carbon source (see experimental), and the second contained potato dextrose broth. Both cultures exhibited strong fungal growth with mycelium covering the entire surface of the liquid media in approximately two weeks. The hydrocarbon fraction was extracted after three weeks using the solid-phase extraction process described above. A GC/MS analysis exhibited a large number of volatile products. A chromatographic comparison of the volatile compounds produced from each growth condition is illustrated in Figure 4, with tentative structural assignments and molecular weights listed in Table 2. The structures of the compounds listed in Table 2 are illustrated in Figure 5. In the case of BS15, both diets produced compounds 1, 2, 3, 5, and 7 but all the other products differed depending on the diet employed. Another notable difference in comparison with CI-4 is that BS15

on the degraded cellulose diet produced approximately the same amounts of volatile products as the potato dextrose diet.

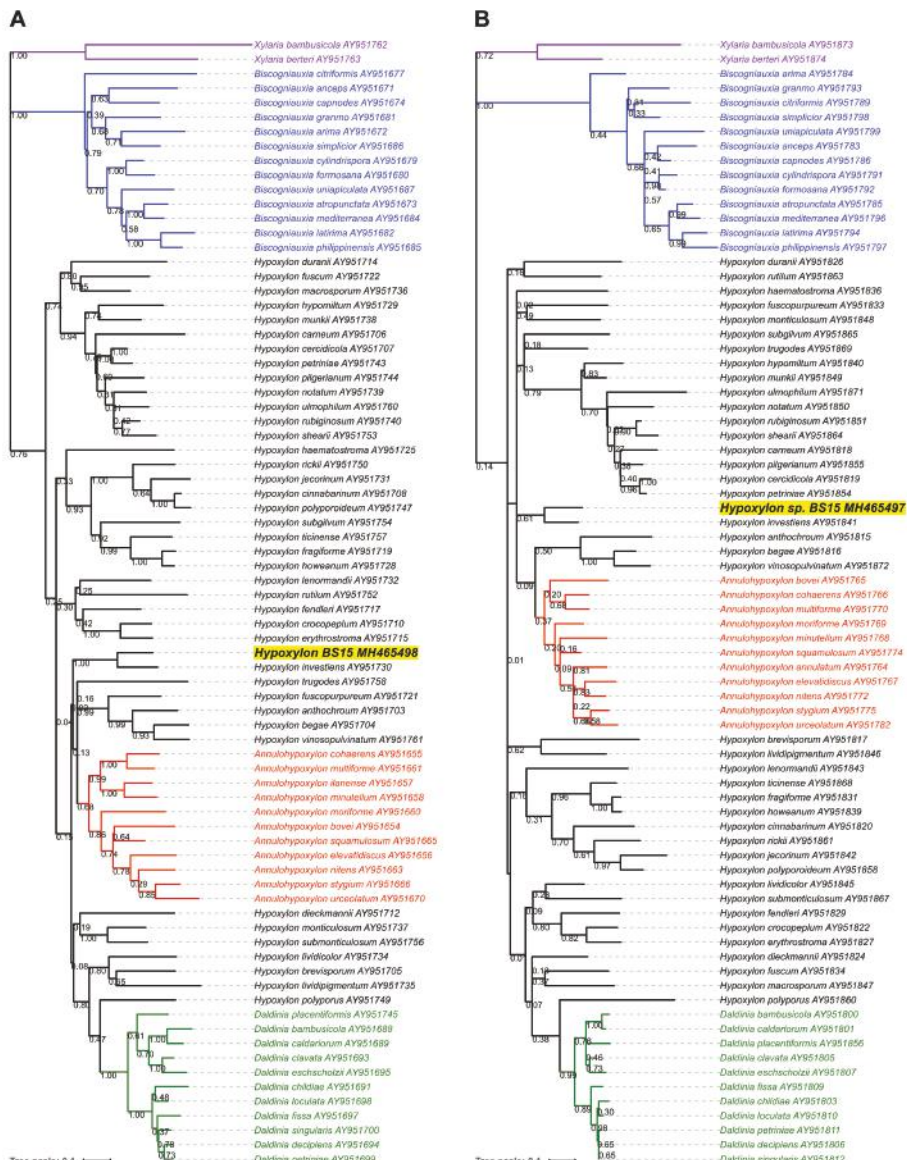


Figure 3. Phylogenetic reconstruction of *Hypoxylon* sp. BS15 and related organisms generated from maximum likelihood clustering of MUSCLE-aligned protein-coding gene sequences. A total of 78 sequences were analyzed for α -actin (A) and β -tubulin (B). The branch lengths are shown to scale, representing the average number of nucleotide substitutions per site between sequences. The bootstrap values at the nodes are from 1000 bootstrap iterations.

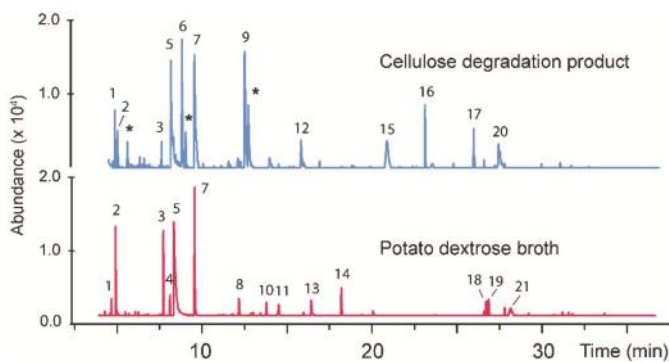


Figure 4. A gas chromatogram showing the volatile organic products produced by the fungus BS15 growth on a conventional media (**bottom**) and on carbohydrates produced from the degraded cellulose (**top**). The asterisks (*) denote volatile contaminants occurring in the degraded cellulose media as determined by analyzing a control sample with no BS15 added. The tentative identification and nominal masses of the individual peaks numbered in the chromatogram are given in Table 2.

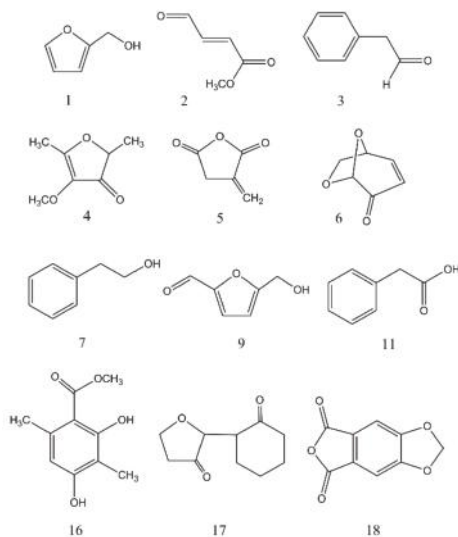


Figure 5. Structures of the volatile compounds tentatively identified from BS15 by comparison with the mass spectra in the NIST database. The compound numbers correspond to the peak numbers listed in Table 2 and in Figure 4.

In order to verify that the volatile compounds resulting from growth of BS15 on the degraded cellulose were being generated from simple sugars rather than from residual cellulose, a control was prepared containing non-degraded cellulose as described above for CI-4. After two weeks of growth on this cellulose medium, no growth was observed.

3.4. Comparing Solid-Phase and Liquid–Liquid Extraction Methods

All extractions of the volatile compounds in this study were performed using a solid-phase extraction with a C-18 stationary phase. This method is widely viewed as a “green chemistry”

alternative to liquid–liquid extractions that require two–three orders of magnitude less solvent. However, it comes with the risk of potentially extracting fewer or different compounds from the growth media. In order to verify that the solid-phase extraction was effectively extracting the growth media, a direct comparison was made versus a liquid–liquid extraction using ethyl acetate/water. The comparisons were made for CI-4 and for BS15. For CI-4, Figure 6 illustrates that the solid-phase extraction gave very similar results, while using approximately 100 times less solvent. The differences in the relative amounts of certain products are notable. For example, the liquid–liquid extraction includes 1,8-cineole, while this product is missing from the solid-phase extraction. At present, the cause of this difference is unknown and further study is needed. Overall, however, the majority of the compounds extracted are the same using either method, and each technique extracts similar amounts as judged by the similar peak areas for the signals having the same retention times.

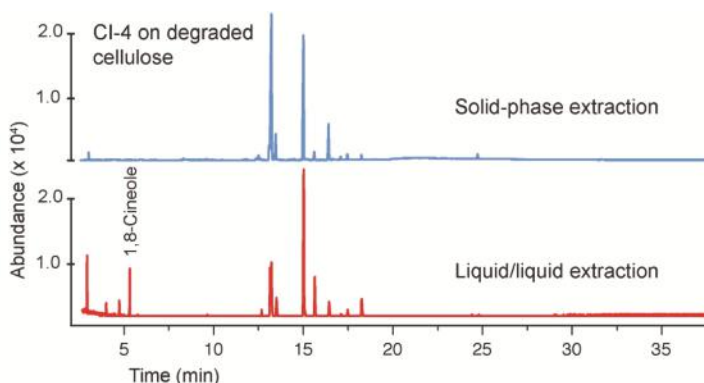


Figure 6. A comparison of the solid-phase extraction versus the liquid–liquid extraction (ethyl acetate/water). The growth media contained the degraded cellulose as a carbon source. A notable difference is the presence of 1,8-cineole in the liquid–liquid extraction. A comparison of the potato dextrose broth gave very similar results and, therefore, is not shown. The identity of 1,8-cineole was verified by comparison to an authentic standard.

A similar comparison of the effectiveness of the liquid–liquid extraction versus the solid-phase extraction was performed using BS15 grown on the degraded cellulose (Figure 7). Here, the differences between the methods were more pronounced with the solid-phase process extracting a more diverse range of products than did the liquid–liquid extraction. The absence of cineole in Figure 7 is a notable omission.

The solid-phase extraction process used herein differs from the vast majority of prior studies on volatile products from fungi, which have utilized solid-phase micro-extraction (SPME). This choice was made because SPME preferentially measures compounds having significant populations in the vapor phase and thus biases analysis against materials having low vapor pressures. Because compounds having potential use as fuels may have low vapor pressure, a methodology was employed here that includes these compounds. Admittedly, the use of a solid-phase extraction (SPE) represents a significant deviation from the common practice in the analysis of volatiles, and future work is needed to directly compare SPE and SPME to clearly identify the advantages and limitations.

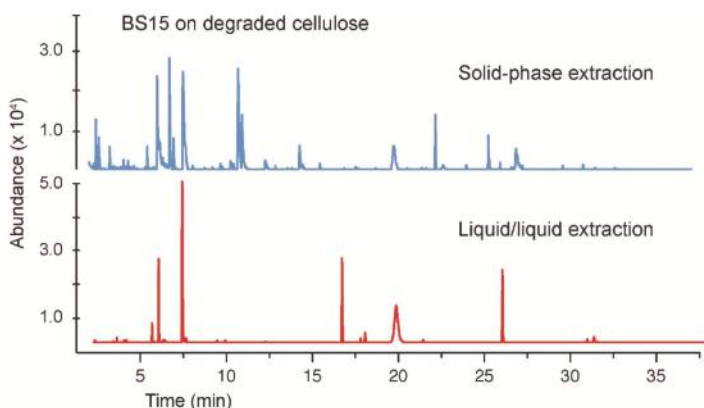


Figure 7. A comparison of the solid-phase extraction versus the liquid–liquid extraction (ethyl acetate/water). The growth media contained BS15 grown on the degraded cellulose as a carbon source.

4. Conclusions

The work described herein establishes the ability of certain endophytic fungi to convert a mechanochemically degraded cellulose product into volatile organic products with potential relevance as fuels. This process has been shown to be feasible with two different *Hypoxyton* sp. to demonstrate that the results are not limited to a single organism. One of the fungi employed (BS15) is described here for the first time, and the phylogenetic analysis demonstrated that it is substantially divergent from any other named species. BS15 produced a range of volatile products that differed significantly from those previously described from CI-4, emphasizing the importance of intra-generic species variation in metabolic studies. Here, the measurement of the total concentration of volatile products is not reported, because the quantification of the individual peaks in the chromatograms is not possible without standards and, at present, several products remain unknown. Our future work will focus on a more complete characterization of the individual compounds and the measurement of the total production.

Author Contributions: Conceptualization, T.M. and R.G.B.; Data curation, M.R.; Formal analysis, T.M., Y.W., and A.H.K.; Methodology, A.H.K. and S.D.M.; Supervision, S.D.M. and J.K.H.; Writing of original draft, J.K.H.

Funding: This research received no external funding.

Acknowledgments: The authors would like to thank Gary Strobel (Montana State University) for providing a sample of the fungus CI-4 for the work described herein. Domenic Valenti is acknowledged for initially isolating the fungus BS15 from an area along the Saint Johns River and near Orange City, FL, USA.

Conflicts of Interest: The authors declare no conflicts of interest.

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Article

Antiplasmodial Properties and Cytotoxicity of Endophytic Fungi from *Sympmania globulifera* (Clusiaceae)

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Received: 27 April 2018; Accepted: 8 June 2018; Published: 12 June 2018

Abstract: There is continuing need for new and improved drugs to tackle malaria, which remains a major public health problem, especially in tropical and subtropical regions of the world. Natural products represent credible sources of new antiparasitic agents for antimalarial drug development. Endophytes that widely colonize healthy tissues of plants have been shown to synthesize a great variety of secondary metabolites that might possess antiparasitic benefits. The present study was carried out to evaluate the antiparasitic potential of extracts from endophytic fungi isolated from *Sympmania globulifera* against a chloroquine-resistant strain of *Plasmodium falciparum* (*PfINDO*). Sixty-one fungal isolates with infection frequency of 67.77% were obtained from the bark of *S. globulifera*. Twelve selected isolates were classified into six different genera including *Fusarium*, *Paecilomyces*, *Penicillium*, *Aspergillus*, *Mucor*, and *Bipolaris*. Extracts from the 12 isolates were tested against *PfINDO*, and nine showed good activity ($IC_{50} < 10 \mu\text{g}\cdot\text{mL}^{-1}$) with three fungi including *Paecilomyces lilacinus* ($IC_{50} = 0.44 \mu\text{g}\cdot\text{mL}^{-1}$), *Penicillium janthinellum* ($IC_{50} = 0.2 \mu\text{g}\cdot\text{mL}^{-1}$), and *Paecilomyces* sp. ($IC_{50} = 0.55 \mu\text{g}\cdot\text{mL}^{-1}$) showing the highest promise. These three isolates were found to be less cytotoxic against the HEK293T cell line with selectivity indices ranging from 24.52 to 70.56. Results from this study indicate that endophytic fungi from *Sympmania globulifera* are promising sources of hit compounds that might be further investigated as novel drugs against malaria. The chemical investigation of active extracts is ongoing.

Keywords: malaria; chloroquine-resistant *Plasmodium falciparum*; *Sympmania globulifera*; endophytic fungi; antiparasitic IC_{50} ; selectivity index

1. Introduction

Malaria remains a major cause of morbidity and mortality with more than 3.3 billion people living worldwide in countries with ongoing transmission at risk [1]. In 2016, 91 countries reported a total of 216 million cases of malaria and 445,000 deaths. The most affected populations were located in tropical and subtropical regions of the world, particularly in sub-Saharan Africa and Southeast Asia, where almost 80% of malaria cases are caused by *Plasmodium falciparum* [1]. The widespread appearance of drug-resistant malaria parasites, even to newly developed second and third generation therapeutics such as artemisinin and its derivatives, makes the development of novel antimalarial drug treatments all the more urgent [2].

Natural products remain a consistent source of antimalarial drug leads, with the best examples being quinine and artemisinin that have acted as gifts of medicinal plants for victims of malaria. Therefore, investigating natural resources for antiplasmodial drug discovery continues to be one of the best scientific approaches that can lead to the identification of novel lead compounds against malaria [3]. Moreover, in comparison to other natural sources like plants, microorganisms are highly diverse but narrowly explored. Indeed, microbes often coexist with plants and animals making it difficult to assess if the real hosts of the promising metabolites identified are from higher organisms themselves or the tiny microbes residing in them. However, studies based on estimation of microbial populations have revealed that only about 1% of bacteria and 5% of fungi have been characterized and the rest remain unexplored for their potential to offer novel drugs against diverse diseases [4]. One of these groups being explored in recent times for their drug discovery potential is endophytic fungi from medicinal plants.

In fact, during the last 20 years it has been observed that much of the wealth of microbial biodiversity with novel biochemistry and secondary metabolite production resides in endophytic association with plant tissues [5]. Interest in such microorganisms, termed endophytes, increased immensely with the discovery of an endophytic fungus, from *Taxus brevifolia*, producing the billion dollar anti-cancer drug taxol [6]. Numerous bioactive molecules have been isolated from endophytic fungi since this groundbreaking discovery [7,8]. Endophytes are metabolically more active than their free counterparts due to their specific functions in nature and activation of various metabolic pathways needed to survive in the host tissues [4,9,10]. Therefore, investigation of endophytic fungi from medicinal plants used to treat malaria can lead to new antimalarial drug discovery.

Sympomia globulifera L. f., an evergreen tree of the Clusiaceae family, is widely distributed in Central and South America, and in tropical Africa, from Sierra Leone to Uganda, Zambia, and Angola. In traditional medicine, a decoction of leaves from this plant is used to manage malaria and several other diseases, including diabetes and skin diseases [11]. In addition, decoctions and extracts of bark are used as diuretics and antiparasitics, and to treat river blindness, chest complaints, cough in children, gonorrhea, scabies, intestinal worms, and prehepatitis jaundice [12,13]. Therefore, the present study was designed to investigate for the first time antiplasmodial activity of endophytic fungi isolated from the bark of *S. globulifera*.

2. Materials and Methods

2.1. Collection of Plant Material

Healthy and mature stem bark of *Sympomia globulifera* L. f. was collected at Nomayos in the Centre Region of Cameroon in January 2015, and was identified at the National Herbarium of Cameroon, where a voucher specimen was deposited under the number 32192/HNC. Plant material was directly brought to the laboratory in sterile bags and processed within a few hours after sampling.

2.2. Isolation of Endophytic Fungi

Stem bark was rinsed with tap water and cut into small pieces, approximately 5 mm length. Sterilization was assessed by means of a 5 min rinse with ethanol, followed by treatment with a 1% active chlorine solution for 15 min, 2 min in ethanol, and a final rinse in sterile water [14].

After surface sterilization, six plates of potato dextrose agar (PDA) supplemented with chloramphenicol ($200 \text{ mg} \cdot \text{L}^{-1}$), each containing 15 pieces of individual plant material, were prepared and kept in the dark at room temperature ($22\text{--}26^\circ\text{C}$). Fragments of mycelium emerging from plant pieces were transferred to new PDA plates without chloramphenicol to obtain pure cultures for identification. To assess whether disinfection methods were effective in eliminating surface fungi, imprints of treated fragments were made by pressing them against the surface of fresh PDA plates that were incubated without plant parts. These plates were checked for fungi emerging from the prints [14].

2.3. Identification of Endophytic Fungi

Fungal cultures were maintained at room temperature (i.e., $22\text{--}26^\circ\text{C}$) under a natural photoperiod for 10–21 days and then examined visually for macroscopic (i.e., morphology, size, and coloration of the mycelium and agar media) and microscopic (i.e., presence of spores or other reproductive structures) characteristics. Colonies were analyzed with respect to their average diameter, sporulation, and the size and coloration of the conidia. Isolates having similar culture characteristics were grouped into morphotype groups. From morphotype groups, only one isolate was processed for further identification.

Identification of endophytic isolates representative of each morphotype was based on the nucleotide sequence of the ITS1-5.8S rRNA-ITS2 region. DNA was extracted from samples of mycelium from the fungal cultures using a commercial kit (RedExtract-N-Amp Plant PCR, Sigma Aldrich, USA). The ITS1-5.8S rRNA-ITS2 region was amplified in a polymerase chain reaction (PCR) using primers ITS4 and ITS5 and the protocol described by White et al. [15]. Amplicons were purified by filtration (MSB Spin PCRapace, Invitrek, Germany) and sequenced.

The FASTA algorithm was used to find sequences similar to those obtained from fungal isolates. The criteria for identification of isolates were based on the similarity of their sequences to those of reliable reference isolates included in public nucleotide databases. To visualize the diverse fungal taxa identified sequences, a dendrogram was made with the ITS1-5.8S rRNA-ITS2 nucleotide sequences of the isolates and those of reference strains deposited in CBS (Centraalbureau voor Schimmelcultures), ATCC (American Type Culture Collection), or other fungal collections, as well as some used in published works of fungal taxonomy. Sequences were aligned using the Clustal X 2.1, and the dendrogram was made with MEGA 6.06 software using the neighbor-joining method with Kimura 2-parameter distances. Groups of sequences at close proximity within the same branch of the dendrogram were individually aligned with Clustal X 2.1 to determine their percentage of similarity. Sequences with a similarity greater than 99% were considered to belong to the same species [14].

2.4. Fermentation and Extraction

Each fungus was cultivated on 2 kg of rice by placing agar blocks of actively growing pure culture (3 mm in diameter) in a 1000 mL Erlenmeyer flask. Each flask was incubated at $25 \pm 2^\circ\text{C}$ for 30 days. After incubation, moldy rice was macerated with ethyl acetate and filtered through three layers of muslin cloth. The organic phase was collected and the solvent was then removed by evaporation under reduced pressure at 40°C using a rotary vacuum evaporator. The dry solid residues were examined for their antiplasmodial activity and cytotoxicity.

2.5. Biological Assays

2.5.1. In Vitro Cultivation of *Plasmodium falciparum*

A chloroquine-resistant *PfINDO* strain of *P. falciparum* was maintained in a continuous culture following the method of Trager & Jensen, 1976 [16] with minor modifications. Cultures were maintained in fresh O positive human erythrocytes suspended at 4% (*v/v*) haematocrit in complete medium (16.2 g·L⁻¹ RPMI 1640 (Sigma) containing 25 mM HEPES, 11.11 mM glucose, 0.2% sodium bicarbonate (Sigma), 0.5% Albumax I (Gibco), 45 mg·L⁻¹ hypoxanthine (Sigma), and 50 mg·L⁻¹ gentamicin (Gibco)) and incubated at 37 °C in a gas mixture consisting of 5% O₂, 5% CO₂, and 90% N₂. The spent medium was replaced with fresh complete medium every day to propagate the culture. Giemsa-stained blood smears were examined microscopically to monitor cell cycle transitions and parasitemia.

2.5.2. Preparation of Stock Solution of Chloroquine and Fungal Extracts

Stock solutions of fungal extracts were prepared in Dimethyl sulfoxide (DMSO) at 25 mg·mL⁻¹ and that of chloroquine phosphate (CQ) (Sigma) was prepared at 1 mM in Milli-Q grade water. The required drug concentrations were achieved by diluting the stocks with incomplete RPMI (Roswell Park Memorial Institute) 1640 medium. The solutions of drugs and extracts were placed in 96-well flat-bottom tissue culture grade plates (Corning). For each extract, the concentrations (μg·mL⁻¹) tested were 0.195, 0.39, 0.78, 1.562, 3.125, 6.25, 12.5, 25, 50, and 100.

2.5.3. In Vitro Antiplasmodial Assay

Before each experiment, synchronized ring stage parasites were obtained by 5% (*w/v*) sorbitol treatment [17]. It is important to note that use of synchronized cultures over mixed-stage cultures can enable the test molecules to interact with all three stages (i.e., ring, trophozoite, and schizont) of the 48 h long life cycle of *P. falciparum* in culture. Moreover, starting the experiment with synchronized ring stage culture provides the distinct advantage of observing growth inhibitory effects without a rise in parasitemia during the ring-trophozoite-schizont transitions.

For drug screening, the SYBR green I based fluorescence assay was used [18]. The ability of SYBR green to give strong fluorescence only in the presence of DNA forms the basis for its use to assess cell proliferation. The absence of a nucleus in human red blood cells where the malaria parasite proliferates allows the use of SYBR green for the specific monitoring of the growth of malarial parasite.

Sorbitol-synchronized ring stage parasites (haematocrit: 2%, parasitemia: 1%, 96 μL) under normal culture conditions were incubated in the presence or absence of increasing concentrations of the extracts. Four μL CQ (1 mM) was used as a positive control and 0.4% DMSO (*v/v*), which was found to be non-toxic to the parasite, was used as vehicle control. After 48 h of incubation, 100 μL of SYBR Green I buffer {0.2 μL of 10,000 × SYBR Green I (Invitrogen) per mL of lysis buffer (Tris (20 mM; pH 7.5), EDTA (5 mM), saponin (0.008%; *w/v*), and Triton X-100 (0.08%; *v/v*)} was added to each well, mixed twice gently with multi-channel pipette and incubated in the dark at 37 °C for 1 h. Fluorescence was measured using a Victor fluorescence multi-well plate reader (Perkin Elmer) with excitation and emission at 485 and 530 nm, respectively. Fluorescence counts for CQ representing zero growth were deducted from counts in each well. A dose-response curve was constructed by plotting fluorescence counts against the drug concentration and IC₅₀ (dose of a drug required to retard the growth of a cell population by 50%) was determined using IC Estimator-version 1.2 (<http://www.antimalarial-icestimator.net/MethodIntro.htm>). In this experiment, no drug (control) corresponds to 100% growth while 40 μM chloroquine (sufficient to cause total arrest of growth) corresponds to 0% growth. Giemsa-stained smears of extract-treated parasite cultures were visualized microscopically to validate the results from fluorescence-based assay. Experiments were done in triplicate and means with standard deviation were calculated. Extracts were classified according to criteria of the antiplasmodial activity based upon good (IC₅₀ < 10 μg·mL⁻¹), moderate (IC₅₀ > 10 to < 25 μg·mL⁻¹), and inactive (IC₅₀ > 25 μg·mL⁻¹) activity as proposed by Bagavan et al. [19].

2.5.4. Cytotoxicity Assay

The cytotoxic effects of the most potent crude extracts were determined by a functional assay [20] using the human embryonic kidney HEK239T cells cultured in complete medium containing 16.2 g·L⁻¹ DMEM, 10% fetal bovine serum, 0.2% sodium bicarbonate (*w/v*) (Sigma), and 50 µg·mL⁻¹ gentamycin. Cells (5 × 10³ cells/200 µL/well) were seeded into 96-well flat-bottom tissue culture plates in complete medium. After 24 h of cells seeding, test extracts were added and cells incubated for 48 h in a humidified atmosphere at 37 °C and 5% CO₂. DMSO (as positive inhibitor) was added at 10% *v/v*. Twenty microliters of a stock solution of MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] (5 mg·mL⁻¹ in 1× phosphate buffered saline) was added to each well, gently mixed, and incubated for another 4 h. The supernatant was thereafter removed and 100 µL DMSO (quench agent) was added to the cell pellet. Formazan formation was measured using a microtiter plate reader (VersaMax tunable multiwell plate reader) at 570 nm. The 50% cytotoxic concentration (CC₅₀) of drugs was determined by analysis of dose–response curves. Selectivity index (CC₅₀/IC₅₀) was calculated for each extract (tested in triplicate).

3. Results

3.1. Isolation and Identification of Fungi

From a total of 90 fragments of the stem bark of *Sympmania globulifera*, 61 isolates were obtained with an infection frequency of 67.77%. They were subsequently grouped into 14 morphotypes and one isolate from each morphotype was identified by sequence analysis of the Internal Transcribed Spacer (ITS) region. Twelve analyzed isolates were classified into six different genera including, *Fusarium*, *Paecilomyces*, *Penicillium*, *Aspergillus*, *Mucor*, and *Bipolaris* (Table 1; Figure 1). The most representative genera were *Fusarium* and *Penicillium* (Table 1).

Table 1. Identification of endophyte isolates and percentage homology of sequences.

Endophyte Isolates	% Sequence Homology	Organism with the Highest Sequence Identity, GenBank Acc. No.
<i>Fusarium decemcellulaire</i>	98	<i>Fusarium decemcellulaire</i> KM231809
<i>Paecilomyces</i> sp.	100	<i>Paecilomyces</i> sp. FJ196762
<i>Fusarium oxysporum</i>	100	<i>Fusarium oxysporum</i> KT794176
<i>Paecilomyces lilacinus</i>	100	<i>Paecilomyces lilacinus</i> GU980023
<i>Penicillium janthinellum</i>	99	<i>Penicillium janthinellum</i> AY373921
<i>Aspergillus tamarri</i>	100	<i>Aspergillus tamarri</i> HQ340111
<i>Penicillium</i> sp. (1)	99	<i>Penicillium</i> sp. JN021538
<i>Bipolaris sorokiniana</i>	94	<i>Bipolaris sorokiniana</i> KU194490
<i>Fusarium lateritium</i>	94	<i>Fusarium lateritium</i> KC453998
<i>Penicillium</i> sp. (2)	98	<i>Penicillium</i> sp. JN021538
<i>Aspergillus aculeatus</i>	100	<i>Aspergillus aculeatus</i> AY585556
<i>Mucor falcatus</i>	89	<i>Mucor falcatus</i> NR103647

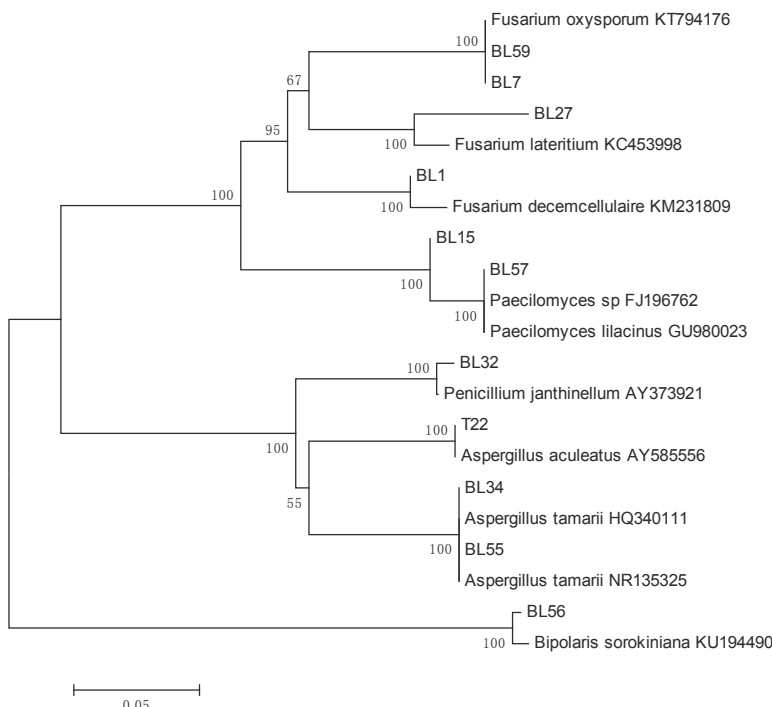


Figure 1. Classification of fungal endophytes isolated from *Symphonia globulifera* based on a neighbor-joining phylogenetic tree made with ITS1-5.8S rDNA-ITS2 nucleotide sequences. Taxa with accession numbers correspond to reference strains. Numbers at branch nodes are bootstrap values based on 500 replications.

3.2. In Vitro Antiplasmodial Activity of Endophytic Extracts

Twelve endophytic fungi were cultured in a rice medium at $25 \pm 2^{\circ}\text{C}$ for 30 days and subsequently extracted using ethyl acetate as a solvent. The extraction yields ranged from 6.30 g to 21.70 g with *A. tamarii* producing the higher yield.

The IC₅₀ of the tested extracts against the chloroquine-resistant *Plasmodium falciparum* INDO (*PfINDO*) strain ranged from 0.2 to $>100 \mu\text{g}\cdot\text{mL}^{-1}$ (Table 2). Out of the 12 tested extracts, nine showed good activity (IC₅₀ $< 10 \mu\text{g}\cdot\text{mL}^{-1}$). Extracts from *Paecilomyces lilacinus*, *Penicillium janthinellum*, and *Paecilomyces* sp. exerted highly potent activities with IC₅₀ $< 1 \mu\text{g}/\text{mL}$. Extracts from *Mucor falcatus* and *Aspergillus aculeatus* showed moderate potency with IC₅₀ between 10 and 25 $\mu\text{g}\cdot\text{mL}^{-1}$, while the extract from *Aspergillus tamarii* with IC₅₀ $> 100 \mu\text{g}\cdot\text{mL}^{-1}$ was considered as inactive.

3.3. Cytotoxicity of Extracts

The cytotoxicity of the extracts was tested against HEK239T cells. As shown in Table 2, CC₅₀ values ($\mu\text{g mL}^{-1}$) ranged from 3.06 to 70.91 leading selectivity indices to lie between 0.5 and 70.56. *Penicillium* sp. (2) (SI = 0.69), *Bipolaris sorokiniana* (SI = 0.504), and *Penicillium* sp. (1) (SI = 0.842) were found to be highly cytotoxic against HEK239T cells. Conversely, the most potent fungal extracts from *Paecilomyces lilacinus* (CC₅₀ = 10.79 $\mu\text{g mL}^{-1}$; SI = 24.52), *Penicillium janthinellum* (CC₅₀ = 9.14 $\mu\text{g mL}^{-1}$; SI = 45.7), and *Paecilomyces* sp. (CC₅₀ = 38.88 $\mu\text{g mL}^{-1}$; SI = 70.56) were less cytotoxic.

Table 2. Antiplasmodial and cytotoxic activities of extracts of endophytic fungi from the bark of *Sympmania globulifera*.

Sample	Dry Mass of Extract (g)	IC ₅₀ against <i>Pf</i> INDO Strain ($\mu\text{g}\cdot\text{mL}^{-1}$ \pm SD)	CC ₅₀ against HEK239T Cells ($\mu\text{g}\cdot\text{mL}^{-1}$ \pm SD)	SI (CC ₅₀ /IC ₅₀)
<i>Fusarium decemcellulare</i>	10.81	2.19 \pm 0.07	70.91 \pm 0.18	32.37
<i>Paecilomyces</i> sp.	14.66	0.55 \pm 0.01	38.81 \pm 0.19	70.56
<i>Fusarium oxysporum</i>	8.23	1.70 \pm 0.22	18.24 \pm 0.11	10.729
<i>Paecilomyces lilacinus</i>	6.20	0.44 \pm 0.03	10.79 \pm 0.2	24.52
<i>Penicillium janthinellum</i>	6.30	0.20 \pm 0.01	9.14 \pm 0.12	45.7
<i>Aspergillus tamarii</i>	21.70	>100	***	NA
<i>Penicillium</i> sp.	14.57	3.63 \pm 0.02	3.06 \pm 0.36	0.842
<i>Bipolaris sorokiniana</i>	9.89	6.10 \pm 0.37	3.08 \pm 0.19	0.504
<i>Fusarium lateritium</i>	11.67	6.61 \pm 0.01	2.82 \pm 0.23	0.426
<i>Penicillium</i> sp.(2)	16.32	9.08 \pm 0.13	6.28 \pm 1.23	0.69
<i>Aspergillus aculeatus</i>	8.74	22.08 \pm 0.43	***	NA
<i>Mucor falcatus</i>	9.22	17.51 \pm 0.19	***	NA
Chloroquine	NA	*400	***	NA

Data are presented as mean values \pm standard deviation of triplicate experiments; SD: standard deviation; ***: not tested; NA: not applicable. SI: Selectivity index (ratio of CC₅₀ to IC₅₀); * IC₅₀ of chloroquine given in nM.

4. Discussion

Endophytic fungi have been claimed to be responsible for the medicinal properties of several medicinal plants [21]. Therefore, in this study we have explored the antiplasmodial potential of endophytic fungi associated with *Sympmania globulifera* used in several African countries for treatment of malaria. More specifically, towards discovery of new drugs against malaria, this study was done to perform antiplasmodial screening of endophytic fungi obtained from the bark of this plant.

A high prevalence of endophyte infection of 67.77% was found in bark fragments of *Sympmania globulifera*. This high level of endophyte colonization is corroborated by previous findings which indicated that endophyte prevalence in some tissues of Cameroonian medicinal plants is quite high [14]. Moreover, Arnold and Lutzoni [22] reported that the prevalence of endophyte species in plant tissues of tropical regions is greater than in plant species of temperate and boreal forests. The identification based on sequence analysis indicated that all of the analyzed endophytic isolates obtained from bark of *S. globulifera* belong to six different genera including *Fusarium*, *Paecilomyces*, *Penicillium*, *Aspergillus*, *Mucor*, and *Bipolaris*. All these genera have also been reported in previous endophyte surveys in different plant species [14,23–26].

Although only 12 endophyte isolates were screened in the present study, as many as nine of them produced antiplasmodial compounds [27,28]. The finding that as high as 75% gave good activity (IC₅₀ < 10 $\mu\text{g}\cdot\text{mL}^{-1}$) with 33.3% (3) of this portion showing highly potent antiplasmodial activity (IC₅₀ < 1 $\mu\text{g}\cdot\text{mL}^{-1}$) suggests that deeper exploration of endophytic fungi may reveal a rich repertoire of antiplasmodial molecules. Previous studies reported that endophyte fungi of *Penicillium* and *Paecilomyces* species are important sources of antiplasmodial secondary metabolites [28]. In fact, β -Resorcyclic acid lactones, cyclodepsipeptides, and tropane derivatives with potent antiplasmodial activities have earlier been isolated from *Paecilomyces* [29–31] and *Penicillium* [27] species.

Our findings have indicated that endophytic fungi belonging to the genera *Fusarium*, *Paecilomyces*, *Penicillium*, and *Bipolaris* isolated from this plant species should be studied intensely for novel antiplasmodial compounds. Therefore, further studies on the antiplasmodial activity guided isolation and characterization of active metabolites produced by promising fungi necessary for the discovery of new drugs against malaria are ongoing.

5. Conclusions

This study is the first report on the antiplasmodial potential of endophytic fungi from the bark of *Symponia globulifera*. In this investigation, 9 out of 12 fungi showed promising antiplasmodial potency with *Paecilomyces lilacinus*, *Penicillium janthinellum*, and *Paecilomyces* sp. being the most promising. These results obtained in this study highlight endophytic fungi from *S. globulifera* as a promising source of novel lead antimarial compounds.

Author Contributions: B.N.L., E.T., and F.F.B. conceived and designed the experiments; J.E.T.A., R.M.K.T., B.M.M., A.F.A., S.G., and B.N.L. performed the experiments; E.R.-F. analyzed the data; N.S., D.S., and B.N.L. contributed reagents/materials/analysis tools; J.E.T.A. wrote the paper.

Acknowledgments: The authors wish to acknowledge the Alexander von Humboldt Foundation for providing a fellowship to Bruno N. Lenta. The authors are also grateful to the TWAS-CNPq (The World Academy of Science-Conselho Nacional de Desenvolvimento Científico e Tecnológico) for the fellowships awarded to Brice M. Mbanning (Award No. 190644/2015-0).

Conflicts of Interest: The authors declare no conflict of interest.

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Article

Restoring Waning Production of Volatile Organic Compounds in the Endophytic Fungus *Hypoxyylon* sp. (BS15)

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Received: 9 May 2018; Accepted: 6 June 2018; Published: 12 June 2018

Abstract: Certain endophytic fungi belonging to the *Hypoxyylon* genus have recently been found to produce volatile organic compounds (VOCs) that have potential relevance as hydrocarbon fuels. Here, a recently discovered *Hypoxyylon* sp. (BS15) was demonstrated to also produce VOCs, but with diminished VOC production after an extended period of in vitro growth. Restoring VOC production was partially achieved by growing BS15 in growth media containing finely ground woody tissue from the original host plant (*Taxodium distichum*). In an effort to isolate VOC production modulators, extracts from this woody tissue were made by sequentially extracting with dichloromethane, methanol, and water. Both the dichloromethane and water extracts were found to modulate VOC production, while the methanol extract had no effect. Surprisingly, the woody tissue remaining after exhaustive extraction was also shown to act as a VOC production modulator when included in the growth media, with changes observed in the production of four compounds. This woody tissue also induced production of two compounds not observed in the original BS15 extract. Filter paper had the same modulating effect as exhaustively extracted woody tissue, suggesting the modulation was perhaps due to cellulose degradation products. Overall, this study demonstrated that VOC production in BS15 can be influenced by multiple compounds in the woody tissue rather than a single modulator.

Keywords: endophytic fungi; volatile organic compounds; *Hypoxyylon*; production modulators

1. Introduction

Endophytes are microorganisms, usually fungi and bacteria, that live inside the host plant without showing signs of their presence or causing apparent disease symptoms. The relationship between endophytes and their host plants varies from symbiotic to pathogenic [1]. Typically, tropical areas and rainforests are presumed to have the greatest diversity and abundance of endophytes due to their vast plant diversity [2]. Investigations involving endophytes are of considerable interest, in part due to their production of a remarkable variety of natural products [3–6]. Although there has been significant focus on endophytes, they remain relatively understudied.

Recently some work has focused on fungi that produce volatile organic compounds (VOCs), including some with potential usefulness as fuels or antimicrobials. For the purposes of this manuscript, VOCs are defined as compounds having sufficient volatility to be separable/mobile on gas chromatography. Table 1 summarizes the presently known fungi that can produce VOCs. Currently, only fungi producing components similar to fossil fuels have been considered to have fuel potential. These compounds include branched alkanes and their derivatives, substituted cyclohexanes, benzenes, alkyl alcohols, aldehydes, and polycyclic aromatic hydrocarbons [7]. For instance, *Gliocladium roseum* produces more than 40 VOCs with fuel potential, such as pentyl, hexyl, heptyl, and octyl alcohols,

3,3,5-trimethyldecane, and other branched hydrocarbons [7]. Among all of the VOCs with fuel potential characterized and reported, 1,8-cineole is a compound of special interest because a 70/30 (*v/v*) mixture of petrol/1,8-cineole has performance characteristics similar to petrol with less carbon monoxide emissions [8–10]. In 2010, a *Hypoxyylon* sp. designated CI-4 was reported as the first non-plant source to produce 1,8-cineole (hereinafter referred to as cineole). More recently, other *Hypoxyylon* spp. have also been found to produce cineole [11,12].

Table 1. A list of fungi presently known to produce volatile organic compounds.

Species	Site Isolation	Extraction Method ^a	No. of VOCs Detected	Reference
<i>Aspergillus fumigatus</i>	New Jersey, USA	SPME	>10	[13]
<i>A. niger</i>	New Jersey, USA	SPME		[13]
<i>A. tubingensis</i>	New Jersey, USA	SPME		[13]
<i>A. niger</i>	Malaysia	LLE	>295	[14]
<i>Fusarium armeniacum</i>	New Jersey, USA	SPME		[13]
<i>F. graminearum</i>	New Jersey, USA	SPME	>10	[13]
<i>F. oxysporum</i>	New Jersey, USA	SPME		[13]
<i>F. proliferatum</i>	New Jersey, USA	SPME		[13]
<i>F. culmorum</i>	Belgium	SPME	>10	[15]
<i>F. langsethiae</i>	Russia	SPME		[16]
<i>F. sibiricum</i>	Russia	SPME	>40	[16]
<i>F. poae</i>	Russia	SPME		[16]
<i>F. sporotrichioides</i>	Russia	SPME		[16]
<i>Metarrhizium anisopliae</i>	New Jersey, USA	SPME	>5	[13]
<i>Mucor racemosus</i>	New Jersey, USA	SPME	>10	[13]
<i>Penicillium chrysogenum</i>	New Jersey, USA	SPME		[13]
<i>P. citreonigrum</i>	New Jersey, USA	SPME	>10	[13]
<i>P. commune</i>	New Jersey, USA	SPME		[13]
<i>P. corylophilum</i>	New Jersey, USA	SPME		[13]
<i>P. crustosum</i>	New Jersey, USA	SPME		[13]
<i>P. glabrum</i>	New Jersey, USA	SPME		[13]
<i>P. pinophilum</i>	New Jersey, USA	SPME		[13]
<i>P. polonicum</i>	New Jersey, USA	SPME		[13]
<i>P. sclerotiorum</i>	New Jersey, USA	SPME		[13]
<i>P. steckii</i>	New Jersey, USA	SPME		[13]
<i>P. sumatrense</i>	New Jersey, USA	SPME		[13]
<i>Nodulisporium</i>	Canary Islands, Ecuador, Thailand, Nicaragua, South Australia, Colombia, and Wetlands of Florida	SPME	>40	[17]
<i>Muscodorus albus</i>	Honduras, Thailand, and Ecuador	SPME	>20	[18]
<i>M. crispans</i>	Bolivian Amazon basin	SPME	>15	[19]
<i>M. kashayum</i>	India	SPME	>20	[20]
<i>M. stroblii</i>	India	SPME	>14	[21]
<i>M. darjeelingensis</i>	India	SPME	>20	[22]
<i>M. tigerii</i>	India	SPME	>20	[23]

Table 1. Cont.

Species	Site Isolation	Extraction Method ^a	No. of VOCs Detected	Reference
<i>M. suthepensis</i>	Thailand	SPME	>25	[24]
<i>M. musae</i>	Thailand	SPME	>15	[24]
<i>M. oryzae</i>	Thailand	SPME	>15	[24]
<i>M. equiseti</i>	Thailand	SPME	>15	[24]
<i>M. sutura</i>	Colombia	SPME	>20	[25]
<i>M. fengyangensis</i>	China	SPME	>20	[26]
<i>Myrothecium inundatum</i>	India	SPME	>30	[27]
<i>Bionectria ochroleuca</i>	India	LLE	>5	[28]
<i>Ampelomyces</i>	Japan	SPME	>5	[29]
<i>Phoma</i>	Japan	SPME	>5	[29]
<i>Cladosporium</i>	Japan	SPME	<5	[29]
<i>Phomopsis</i>	Ecuador	SPME	>10	[30]
<i>Gliocladium roseum</i>	Northern Patagonia	SPME	>40	[7]
<i>Beauveria bassiana</i>	Montana, USA	SPME	6	[31]
<i>Ascocoryne sarcoides</i>	Northern Patagonia, UK, Germany, France, Norway, and Canada	SPME	>100	[32]
<i>A. cylindrum</i>	Norway, Switzerland	SPME		[32]
<i>A. solitaria</i>	Netherlands	SPME		[32]
<i>Schizophyllum commune</i>	Chile	SPME	10	[33]
<i>Hypoxyylon</i>	Thailand, Spain	SPME	>15	[34]

^a The abbreviations LLE and SPME denote liquid-liquid extraction and solid phase micro-extraction, respectively.

An unexpected challenge involving cineole production in CI-4 was the observation that the production gradually decreased over a period of months when the organism was removed from the plant host. This decrease suggested the presence of one or more cineole production modulator compounds in the host plant. Nigg et al. isolated and characterized a modulator in an endophytic *Nodulisporium* species, the imperfect stage of *Hypoxyylon* [35]. This modulator was able to restore cineole biosynthesis, and it is likely that similar outcomes can be obtained in other endophytic fungi where production of valuable products decrease over time. In related work, Hassan et al. reported that the treatment of *Hypoxyylon* sp. with known epigenetic modulators not only cause phenotypic changes, but also modifies the VOCs production and the bioactivity [34]. All of these prior studies provide new insight into why such a diverse range of VOCs are found in different isolates of *Hypoxyylon* spp.

Recently, a *Hypoxyylon* sp. not corresponding to any named species was isolated from a bald cypress tree (*Taxodium distichum*) near Orange City, Florida, USA. A complete phylogenetic characterization of this fungus, designated BS15, will be given elsewhere. BS15 produces a variety of VOCs with possible relevance as fuels or antimicrobials. As with other *Hypoxyylon* spp., BS15 was found to exhibit a significant decrease in VOC production over time and, motivated by the work of Nigg et al. [35], techniques for restoring VOC production were evaluated. This manuscript describes a process in which extracts from woody tissue of the plant from which BS15 was originally isolated were added to the growth media in an effort to restore VOC production. Serial extractions of the woody tissue were performed with dichloromethane (DCM), methanol, and water. In the following, we describe the changes from each extract and show that the DCM and water extracts, as well as the exhaustively extracted wood tissue, induce production of compounds. These changes were found to be inheritable, and three of the products are shown to differ from those originally produced by BS15, suggesting that the modifications represent epigenetic changes.

2. Materials and Methods

Dichloromethane (DCM) and methanol were purchased from Fisher Scientific (Salt Lake City, UT, USA). SiliaPrep C-18 columns were purchased from Silicycle (Quebec City, Canada). Potato dextrose broth (PDB) and agar were purchased from Microtech Scientific (Vista, USA). All reagents were used as received.

The *Hypoxyylon* sp. BS15 was isolated from a bald cypress tree (*Taxodium distichum*) near the Saint Johns river near Orange City, Florida, USA. Initially, isolation of the fungus followed the procedures of Tomsheck et al. [11]. This involved treating branches with 70% ethanol, further sterilizing the wood in a flame, and then drying in a sterile laminar-flow hood. Outer tissue was cut away using a sterile knife blade and a square section of inner tissue was then placed on water agar. Any fungal hyphae growing out from the sample were transferred onto different plates of potato dextrose agar (PDA). One such sampling resulted in isolation of BS15.

Growth of BS15 samples without added modulators (i.e., extracts from *T. distichum*) was accomplished in PDB prepared by adding 2.4 g of potato dextrose broth to 100 mL purified water in a 500 mL Erlenmeyer flask. The flask was sealed with aluminum foil and autoclaved for 15 min to sterilize. A culture of BS15 growing on PDA was then added to the sterile broth, and it was resealed with aluminum foil and left to grow for 30 days in the lab at room temperature without stirring. The resulting broth was then vacuum filtered twice with Whatman Grade 4 filter paper to remove all particulates.

Extractions of woody tissue involved taking approximately 210 g of shredded cypress wood and extracting with 400 mL DCM for 90 min with stirring. This process was repeated two times. The DCM was then removed on a rotatory evaporator (Buchi Rotavapor R-205, Buchi, New Castle, DE, USA) at reduced pressure. The cypress wood was further extracted with methanol, and finally with water, using 400 mL of each solvent. In order to simulate the extractable components in a wood matrix, all three fractions were dripped onto filter papers, and the filter paper was allowed to completely dry. This process was repeated several times until the filter paper contained the desired mass of the extracted components. These filter papers were cut into pieces and added to 100 mL of the PDB medium in a 500 mL Erlenmeyer flask, and this flask was inoculated with BS15. As a control, filter paper containing no extractables was added to the PDB/BS15 media. As a final test, the exhaustively extracted cypress wood was also added to PDB. All flasks were then sealed with aluminum foil and autoclaved for 15 min. After cooling, BS15 was added to the sterile broth, and the resulting solution was allowed to grow for 30 days at room temperature without stirring.

Media used to evaluate the influence of the DCM, methanol, and water extracts on BS15 were prepared by removing a sample of the fungi grown in a solution containing PDB plus *T. distichum* extracts after 30 days of growth and transferring it onto a petri dish. Serial weekly transfers onto PDA were then performed over a period of 4 weeks to ensure that all changes in VOC production ultimately observed were epigenetic changes, and that exogenous contaminants from the extractable components were rigorously removed.

Isolation of VOCs involved solid phase extraction of the growth media on a C-18 stationary phase (500 mg). First, a C-18 cartridge (particle size: 40–63 µm) was washed with 5 mL of methanol, and then with 5 mL of water (three times). A total of 100 mL of filtered fungal broth was then passed through the column under vacuum. The column was washed with 15 mL of water to remove polar components (e.g., salts), and the column was dried by drawing air through the column for 30 min. The column was then eluted by passing 1.5 mL of methanol through the column to yield a clear brown solution. The eluent was filtered using a 0.22 µm syringe filter prior to gas chromatography/mass spectrometry (GC/MS) analysis. This solid-phase extraction methodology differs from the solid-phase microextraction (SPME) methodology usually employed when evaluating fungal VOCs. The methodology was employed in order to more efficiently retain compounds having low vapor pressure, and which may be missed by SPME.

The GC/MS method used was similar to that of Strobel et al. [36]. The instrument used was a Finnigan TraceGC Ultra with Trace DSQ detector (Thermo Scientific) and a Restek Rtx-225 capillary column (cyanopropyl-methyl/phenyl-methyl polysiloxane, 50/50, 30 m × 0.25 mm, film thickness 0.25 µm) (Restek, Bellefonte, PA, USA). The carrier gas was ultra-high purity helium with a 1.5 cm³/min constant flow rate and an initial column head pressure of 77 kPa. The injector was set to 250 °C with a 1 µL injection volume using splitless injection mode. The column oven temperature was initially 45 °C and held for one minute, followed by a 10 °C/min ramp to 100 °C, where the temperature was held for 5 min. Finally, the temperature was increased by 5 °C/min to 200 °C and held for 7 min. The detector was set at 280 °C and set to scan 50–650 m/z. Data acquisition and processing were performed on Xcalibur software. Identification of compounds was made via library comparison using National Institute of Standards and Technology (NIST) database. In all the GC/MS analyses described herein, quantities of individual compounds detected are not reported because many of the compounds are unknowns. This ambiguity prevents the construction of calibration curves required for quantitation.

3. Results and Discussion

The endophytic fungus BS15 was selected for study based on the observation that some of the compounds produced had a distinctive odor, which indicated production of volatile compounds. A GC/MS analysis of the original BS15 revealed a number of VOCs (Table 2). Unfortunately, the production of several compounds decreased with time in the absence of the host plant (Figure 1). The nominal masses of all compounds were obtained and five compounds were tentatively identified. A more complete characterization of the compounds present will be given elsewhere. When BS15 with diminished VOC production was transferred back to PDA containing woody tissue from the host plant (i.e., finely ground *Taxodium distichum* tissue), production of most VOCs was restored, albeit to varying degrees (Figure 1, top plot). The ability to restore VOC production in BS15 suggested the presence of a modulator compound or multiple modulators in the host plant.

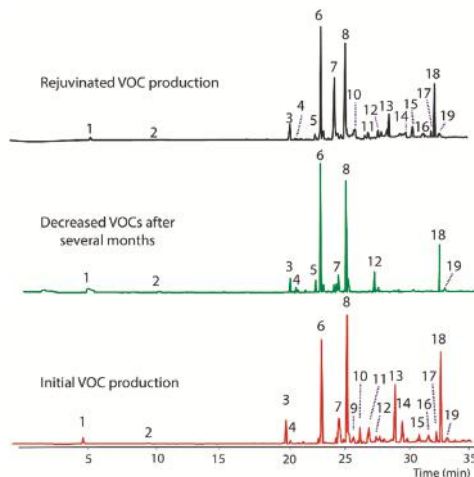


Figure 1. A gas chromatogram illustrating VOCs produced by BS15 showing the original production of VOCs immediately after isolation of BS15 (**bottom**) and decreased production after growing in the lab for several months (**middle**). Nominal mass of each peak and tentative identities are listed in Table 2. Production of VOCs was restored to varying extents (**top**) by growing BS15 on PDA containing finely ground woody tissue from the *Taxodium distichum* the fungus was originally isolated from.

Table 2. A GC/MS analysis of the VOCs produced by BS15.

Peak	R.T. (min)	Tentative Identify	Mol. Mass
1	4.75	unknown	70
2 ^a	9.41	1,8-cineole	154
3	19.15	unknown	142
4	19.78	unknown	120
5	21.84	unknown	126
6 ^a	22.35	Phenyl ethyl alcohol	122
7 ^a	23.85	2,3-naphthalenediamine	158
8	24.51	unknown	182
9	24.75	Unknown	184
10	25.87	unknown	220
11 ^a	26.58	Phenylacetic acid	136
12	27.62	Unknown	298
13 ^a	28.72	Diethyl phthalate	222
14	29.52	unknown	297
15	30.72	unknown	213
16	31.47	unknown	334
17	32.15	unknown	213
18	32.63	unknown	192
19	33.27	unknown	314

^a Assignment confidence for peaks 2, 6, 7, 11, and 13 are, respectively, 86%, 89.3%, 82.7%, 76.4%, and 94%. No other peaks correspond to compounds in the NIST database.

4. Decreased VOC Production in BS15 after Extended In Vitro Growth

In order to investigate modulators from the host plant that restores production of VOCs in BS15, serial organic solvent extraction of woody tissue was made using DCM followed by methanol and then water. Each extract was then tested for its ability to restore VOC production in BS15. Since some of the extracted compounds were insoluble in the growth media, the extracted solutions were dripped onto filter paper and then air dried. Filter paper was employed to simulate the woody matrix of the original tissue. This process was repeated until the desired mass of extract had been loaded onto the filter paper (see Section 2). Growth media (PDB) was then prepared, and the filter paper impregnated with extractable compounds was included in the media. The filter paper was cut into strips of approximately $1'' \times \frac{1}{4}''$ to give a uniform distribution in solution. In each case, a control was also prepared containing filter paper with no extract added. The exhaustively extracted wood was also evaluated by including it in the growth media. The impact of each extract on production of volatiles is discussed below.

5. Assessing the Influence of DCM Extract/Filter Paper on VOC Production

A culture of BS15 grown in a PBD medium containing DCM extract/filter paper was found to alter the VOCs produced by inducing the production of three new compounds. Specifically, the peaks labeled 20, 21, and 22 in Figure 2, with respective nominal masses of 112, 216, and 154, were observed only after addition of the DCM extract, and thus appear to represent an epigenetic change to BS15. Surprisingly, the control containing only filter paper also induced production of compounds 21 and 22. In both cases, these changes in VOC production were inheritable, and persist over several generations. Indeed, Figure 2 represents BS15 VOCs obtained from tissue removed from the DCM/filter paper media, then plated onto PDA, followed by weekly transfers onto PDA for one month, and finally regrown in PDB. In other words, Figure 2 represents a BS15 culture that was three generations removed from the initial DCM/filter paper treatment. Careful inspection of the chromatograms showed that the DCM extracts also increased production of peak 18 and decreased production of 6. Overall, it appeared that DCM contained a modulator that altered production of peaks 6 and 18, and created the ability to produce 20. Remarkably, the filter paper appeared to be solely responsible for the production of

compounds **21** and **22**, as discussed below. Isolation of individual modulator compounds from the DCM extract was not performed due to insufficient mass of DCM extract.

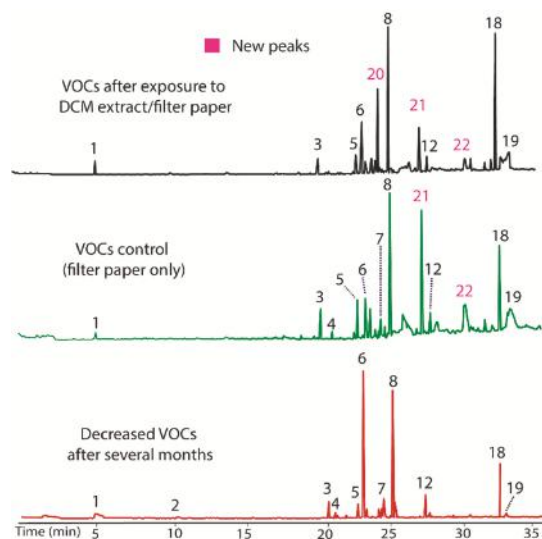


Figure 2. Gas chromatograms showing decreased VOCs production by BS15 after growing in vitro for several months (**bottom**). Adding BS15 to growth media (PDB) containing DCM extract/filter paper induced production of new compounds **20**, **21**, and **22** (**top**), and altered the production of compounds **6** and **18**. The **middle** plot demonstrates that filter paper containing no DCM extract also induces production of compounds **21** and **22** when included in growth media. Nominal mass of each peak and their tentative identities are listed on Table 3.

Table 3. Compounds detected by GC/MS from BS15 modified by treatment with DCM extract/filter paper.

Peak	R.T. (min)	Tentative Identify	Mol. Mass
1	4.75	Unknown	70
2	9.41	1,8-Cineole	154
3	19.15	Unknown	142
4	19.78	Unknown	120
5	21.84	Unknown	126
6	22.35	Phenyl ethyl alcohol	122
7	23.85	2,3-Naphthalenediamine	158
8	24.51	Unknown	182
9	24.75	Unknown	184
10	25.87	Unknown	220
11	26.58	Phenylacetic acid	136
12	27.62	Unknown	298
13	28.72	Diethyl phthalate	222
14	29.52	Unknown	297
15	30.72	Unknown	213
16	31.47	Unknown	334
17	32.15	Unknown	213
18	32.63	Unknown	192
19	33.27	Unknown	314
20 ^a	23.76	3-methyl-2,5-furandione	112
21	26.85	Unknown	216
22 ^a	29.82	4,4'-thiobis-benzeneamine	154

^a Assignment confidence for peaks **20** and **22** are, respectively, 67% and 71%.

6. Evaluating the Influence of Methanol and Water Extracts on VOC Production in BS15

The influence of both the methanol and water extracts from *T. distichum* on BS15 were also evaluated using the process described above for the DCM extract. The methanol extract/filter paper produced almost no change, with the exception that peaks 21 and 22 were again observed. Chromatograms illustrating VOC production before and after addition of methanol extract/filter paper are shown in Figure 3. In this case, the control containing only filter paper also induced production of 21 and 22. The production of these compounds also occurred in the DCM extract and their occurrence is thus attributed to the filter paper as a VOC production modifier rather than any compounds extracted by methanol.

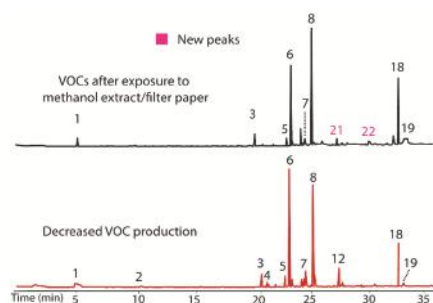


Figure 3. Chromatograms showing the negligible influence of the methanol extract/filter paper on VOC production in BS15. The new peaks (21 and 22) observed upon treatment (**top**) also occur in the control containing only filter paper. Their occurrence is therefore attributed to a change from the filter paper rather than the presence of VOC production modifiers extracted by methanol. The **bottom** plot illustrates BS15 after prolonged *in vitro* growth.

The water extract/filter paper was also evaluated (Figure 4) using the process described above. This extract increased production of compounds 7, 12, and 18, while decreasing the quantity of 6. The water extract thus likely contained a VOC production modulator. As in the other extracts, peaks 21 and 22 were again observed in both the extract/filter paper and in the control, strengthening the conclusion that filter paper induces their production. All changes from the water extract were inheritable.

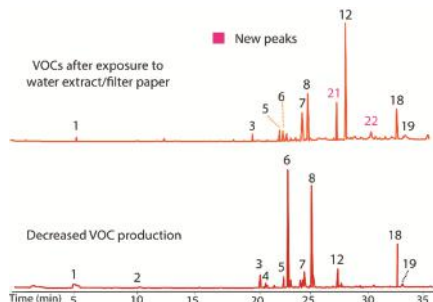


Figure 4. Chromatograms showing the influence of the water extract/filter paper of VOC production of BS15. Peaks 7, 12, and 18 increased upon exposure to the water extract while 6 decreased (**top**) versus BS15 with diminished VOC production (**bottom**). The water extract thus appeared to contain a modulator of VOC production. Peaks 21 and 22 were again observed to occur in both the water extract and in the control containing only filter paper.

7. The Influence of Exhaustively Extracted *T. distichum* Wood on VOC Production in BS15

As a final test of potential modulators of VOC production in BS15, the finely ground extracted woody tissue of *T. distichum* was evaluated. This unusual step was taken because the filter paper was repeatedly found to induce production of peaks 20 and 21 in prior extracts, and it was of interest to see if other cellulose contain materials could have the same effect. The extracted wood was observed to decrease production of peaks 7 and 12, while it increased 5 and 18 (Figure 5). Of greatest interest was the observation that peaks 21 and 22 appeared as prominent peaks, supporting the contention that these peaks were induced by cellulose containing materials. A comparison of the woody tissue to the filter paper controls from each extract is illustrated in Figure 6. The remarkable similarity in the changes induced by filter paper and those induced by exhaustively extracted wood from *T. distichum* indicated that a common production modulator is involved.

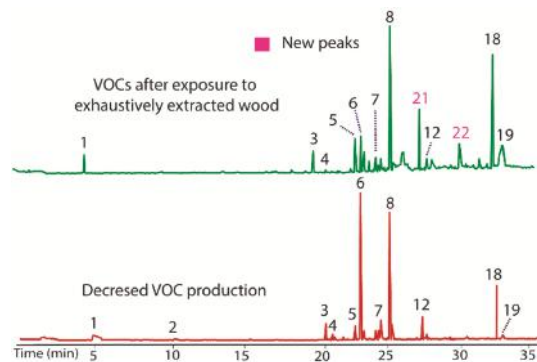


Figure 5. Chromatograms showing the influence on VOC production (**top**) of the addition of exhaustively extracted woody tissue of *T. distichum* to growth media (PDB). Peaks 5 and 18 increased, while 7 and 12 decreased versus BS15 after prolonged in vitro growth (**bottom**). Peaks 21 and 22 appeared as prominent components in the top plot, supporting the contention that these peaks were induced by cellulose containing materials.

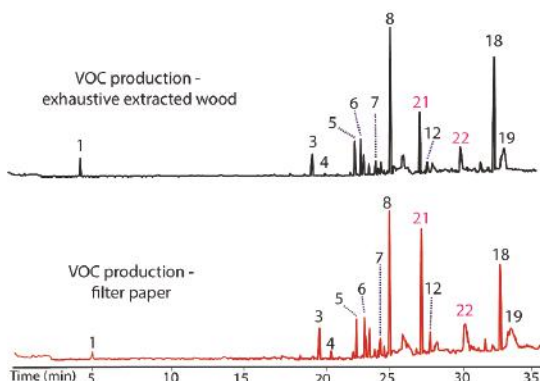


Figure 6. Chromatograms illustrating the changes in the production of VOCs by BS15 following exposure to filter paper (**bottom**) and to exhaustively extracted woody tissue from branches of *T. distichum* (**top**).

A direct comparison of each extract and its influence on VOC production is summarized in Table 4, and shows that the influence of DCM, water, and extracted wood (or equivalently, filter paper) clearly differed. This result indicated that multiple modulators were involved in VOC production as opposed to a single modulator.

Table 4. A summary of compounds influenced by different extracts of *T. disticum* ^a.

Extract	Increased Production	Decreased Production	New Compounds Produced
DCM	18	6	20, 21, 22
Methanol	None	None	21, 22
Water	7, 12, 18	6	21, 22
Wood, extracted	5, 18	7, 12	21, 22
Filter paper	5, 18	7, 12	21, 22

^a All analyses of VOC production were made on fungi from the fourth generation after initial exposure to the extract or woody tissue/filter paper.

An important question regarding the influence of modulators on VOC production is how long the fungus with compromised VOC production should be exposed to media containing ground plant tissue or extractable compounds in order to restore VOC production. In the case of BS15, we observed that continuous exposure to the woody tissue or extracts/filter paper for three generations was sufficient to restore production. For the purposes of this study, a transfer of the fungus was made each week to new media containing modulators (i.e., extracts). Thus, three generations correspond to three weeks. In all cases, the fungus was removed from the media containing extractables or wood and grown for a week to maturity before transfer to PDB to ensure that all exogenous contaminations were removed. Similarly, it was important to evaluate how long the fungus with restored production could grow in vitro before VOC production began to decrease. In BS15, a significant decrease in VOC production was observed after eight generations (8 weeks).

8. Conclusions

The research demonstrated that waning production of volatile organic compounds in an endophytic *Hypoxyylon* sp. (BS15) can be partially restored by re-exposure to chemical constituents contained in the DCM and water extracts of the woody tissue of *T. distichum*. Surprisingly, the exhaustively extracted woody tissue also induced changes in VOC production from BS15 by causing production of two compounds never observed in the original growth media, and in altering production of four other compounds. Filter paper alone can also produce this change, suggesting that the differences are epigenetic changes, and that cellulose or its degradation products were the active component in altered VOC production rather than other compounds (e.g., lignans). The cellulose-based modification of production may have been caused by hydrolysis of the woody tissue or filter paper, which could create water soluble carbohydrates, and further study of this effect is needed. It is notable that carbohydrates are known to alter gene expression in some bacteria [37], thus there is a precedent for this change in BS15 from cellulose containing materials. An interesting parallel to this observation of activity in cellulose, a material that is essentially insoluble, is a prior study [35] reporting that breakdown products from lignin (i.e., ferulic acid and vanillin) also act as VOC production modulators in a closely related endophytic fungus.

It is notable that the large solubility difference between the water and DCM extracts suggests that the production modulator in DCM differed from that found in water. Thus, it is likely that more than one modulator effectively influenced VOC production in BS15. At present, insufficient amounts of the DCM and water extracts were available to allow isolation of individual compounds, and future work will focus on identifying compounds in these extracts involved in VOC modulation.

Solid-phase extraction was used in this study, rather than the more commonly employed approach of solid-phase microextraction, in order to more effectively include compounds having lower vapor

pressures. One possible limitation to this approach is that compounds having high vapor pressures may be underrepresented. Further study is underway to directly compare these two techniques.

Author Contributions: Y.W. performed the growth of all BS15 cultures, performed all gas chromatography/mass spectrometry analyses, provided a comprehensive literature review of prior work on endophytic fungi producing volatile organic compounds, and assisted in manuscript preparation. The idea for this work was conceived of by J.K.H., and he wrote much of the manuscript.

Acknowledgments: Domenic Valenti is acknowledged for initially isolating the fungi BS15 from an area along the Saint Johns River and near Orange City, FL, USA.

Conflicts of Interest: The authors declare no conflict of interest.

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Article

Antifungal Activities of Volatile Secondary Metabolites of Four *Diaporthe* Strains Isolated from *Catharanthus roseus*

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Received: 1 May 2018; Accepted: 29 May 2018; Published: 30 May 2018

Abstract: Four endophytic fungi were isolated from the medicinal plant, *Catharanthus roseus*, and were identified as *Diaporthe* spp. with partial translation elongation factor 1-alpha (*TEF1*), beta-tubulin (*TUB*), histone H3 (*HIS*), calmodulin (*CAL*) genes, and rDNA internal transcribed spacer (ITS) region (*TEF1-TUB-HIS-CAL-ITS*) multigene phylogeny suggested for species delimitation in the *Diaporthe* genus. Each fungus produces a unique mixture of volatile organic compounds (VOCs) with an abundant mixture of terpenoids analyzed by headspace solid-phase microextraction (SPME) fiber-GC/MS. These tentatively-detected terpenes included α -muurolene, β -phellandrene, γ -terpinene, and α -thujene, as well as other minor terpenoids, including caryophyllene, patchoulene, cedrene, 2-carene, and thujone. The volatile metabolites of each isolate showed antifungal properties against a wide range of plant pathogenic test fungi and oomycetes, including *Alternaria alternata*, *Botrytis cinerea*, *Colletotrichum gloeosporioides*, *Fusarium graminearum*, and *Phytophthora cinnamomi*. The growth inhibition of the pathogens varied between 10% and 60% within 72 h of exposure. To our knowledge, the endophytic *Diaporthe*-like strains are first reported from *Catharanthus roseus*. VOCs produced by each strain of the endophytic *Diaporthe* fungi were unique components with dominant monoterpenes comparing to known *Diaporthe* fungal VOCs. A discussion is presented on the inhibitive bioactivities of secondary metabolites among endophytic *Diaporthe* fungi and this medicinal plant.

Keywords: endophytic fungi; *Diaporthe* spp.; *Catharanthus roseus*; volatile organic compounds (VOCs); antifungal bioactivity; inhibition; terpene; pathogens

1. Introduction

Many plants remain unexplored for their endophytic fungi and the potentially important products that they may produce [1]. *Catharanthus roseus* is known as a pharmaceutical plant containing rich anticancer alkaloids. The extracts of many organs of this plant also exhibit antimicrobial effects [2–6]. It turns out that *Catharanthus roseus* is host to a diverse group of endophytic fungi [7–10]. Some endophytic fungi were found to produce several metabolites biosynthesized by the host *Catharanthus roseus*. The endophytic fungi *Curvularia* sp. CATDLF5 and *Choanephora infundibulifera* CATDLF6 isolated from leaf issues were able to enhance leaf vindoline production content of *C. roseus* cv.

prabal by 2.29–4.03 times through root inoculation [8]. Endophytic *Fusarium* spp. from stem issues seemed to facilitate the host plant to produce secondary metabolites [9]. Additionally, some endophytic fungi from the plant produced antimicrobial compounds. For example, the compounds hydroxyemodin, citreoisocoumarin, citreoisocoumarinol, and cladosporin from endophytic fungi of leaves were effective in inhibiting fungal pathogens [10]. *Diaporthe* are commonly found as endophytes in a wide range of plants around the world [11–15]. These endophytes are prolific producers of antimicrobial metabolites [15,16]. *D. endophytica* and *D. terebinthifolii*, isolated from the medicinal plants *Maytenus ilicifolia* and *Schinus terebinthifolius*, had an inhibitory effect against *Pseudomonas citricarpa* in vitro and in detached fruits [12,13]. The crude extracts of *Diaporthe* sp. MFLUCC16-0682 and *Diaporthe* sp. MFLUCC16-0693 exhibited notable antibacterial and antioxidant activities [14]. An endophytic *Phomopsis* (asexual state of *Diaporthe*) fungus isolated from the stems of *Ficus pumila*, exhibited broad-spectrum antimicrobial activity against Gram-positive and Gram-negative human and phytopathogenic bacteria and fungi [15]. Thus, the genus *Diaporthe* is a potential source of metabolites that can be used in a variety of applications [14]. However, endophytic *Diaporthe* fungi have not been recorded from *Catharanthus roseus* to the present.

Volatile organic compounds (VOCs) have noted biofumigative effects especially from the endophytic fungus—*Muscodor albus* [17]. These observations opened a unique venue for the application of endophytic microorganisms to the ecological-friendly biocontrol of pests [17]. The inhibitive bioactive compounds were also found in a few isolates of endophytic *Diaporthe* [18]. An endophytic *Phomopsis* isolate of *Odontoglossum* sp. in Northern Ecuador was reported to produce a unique mixture of volatile organic compounds (VOCs) with sabinene, 1-butanol, 3-methyl; benzeneethanol; 1-propanol, 2-methyl, and 2-propanone. The VOCs showed antifungal bioactivities on a wide range of plant pathogenic fungi, such as *Sclerotinia*, *Rhizoctonia*, *Fusarium*, *Botrytis*, *Verticillium*, *Colletotrichum* and oomycetes *Pythium*, and *Phytophthora* [18]. The PR4 strain of an endophytic *Phomopsis* obtained from the medicinal plant *Picrorhiza kurroa* also produced a unique set of bioactive VOCs inhibitive to plant pathogenic fungi growth. The dominant compounds in VOCs of the PR4 strain were reported as menthol, phenylethyl alcohol, isomenthol, β -phellandrene, β -bisabolene, limonene, 3-pentanone and 1-pentanol [19]. In view of the antimicrobial properties of the extracts from the medicinal plant *Cantharatus roseus*, and limited knowledge on endophytic *Diaporthe* species in this host, we conducted an investigation on the antifungal bioactivity of VOCs from four endophytic *Diaporthe* strains isolated from wild *Catharanthus roseus* in China. The combined sequences of five loci, elongation factor 1-alpha (*TEF1*), beta-tubulin (*TUB*), histone H3 (*HIS*), calmodulin (*CAL*) genes, and the rDNA internal transcribed spacer (ITS) region were used for the strains' phylogenetic analyses within genus *Diaprothe*. Inhibitory bioactivity executed volatile organic compounds from the strains were observed on growths of tested plant pathogens in co-culture. Active components of VOCs were analyzed and inferred using headspace solid-phase microextraction (SPME) fiber-GC/MS and based on their reported properties.

2. Materials and Methods

2.1. Endophytic Fungal Isolation

The four endophytic fungi were isolated from wild plants, *Catharanthus roseus*, growing in the National Natural Conservation Area of TongGu Mountain, located in Wenchang city of Hainan Province. Several stem segments (5–10 cm in length) were collected for the eventual isolation of endophytes. Retrieving endophytic fungi followed a previously described procedure [20]. Briefly, the external tissues of segments were cleaned with tap water and scrubbed with 70% ethanol prior to excision of internal tissues. Then the segments were excised into smaller fragments about 0.2–0.5 cm in length. The fragments were thoroughly exposed to 75% ethanol for 60 s, 3% NaClO for 90 s, and sterile water for 60 s by agitation. The fragments at the last step were drained on sterile filter papers and put on water agar in Petri plates for growing endophytes. Further, pure isolates were obtained in potato dextrose agar media and stored on sterilized, inoculated barley seeds at 4 °C and –80 °C.

The four fungi of interest were assigned with our laboratory acquisition number-ID FPYF3053-3056 and deposited in China Forestry Culture Collection Center assigned IDs of CFCC 52704-52707.

2.2. DNA Extraction, PCR, and Sequencing

Fungal genomic DNA was extracted from colonies growing on PDA for one week with the CTAB procedure [20]. The extracted DNA was further purified through Mini Purification kit (Tiangen Biotech (Beijing) Co., Ltd., Beijing, China) following the manufacturer's protocols. The DNA quality and concentration were determined with a NanoDrop 2000 (Thermo Fisher Scientific Inc., Waltham, MA, USA) after the DNA was checked with Genegreen nucleic acid dye (Tiangen Biotech (Beijing) Co., Ltd.) in an electrophoresis on 1% agarose gel stained under ultraviolet light. The extracted DNA was used as a template for the further PCR amplification ITS sequence and *TEF1*, *CAL*, *TUB*, and *HIS* genes regions. The primers were used to amplify the ITS targets, namely, the ITS1 and ITS4 [21], *TEF1* with EF1-688F/EF1-1251R [22], *CAL* with CL1F/CL2A or CAL563F/CL2A [23], *TUB* with T1/Bt-2b or Bt2a/Bt-2b [24,25], and *HIS* with HISdiaF/HISdiaR, sequences that were 5'-GGCTCCCCGTYAACGAGCTCGCCTCC-3 and 5'-ATYCCGACTGGATGGTCACACGCTTGG-3, respectively. All PCR reaction mixtures and conditions were followed as per the Taq PCR MasterMix kits (Tiangen Biotech (Beijing) Co., Ltd.) according to the manufacturer's protocol. A PCR reaction system consisted of 0.5 µL of each primer (10 µM), 3 µL (15–80 ng) of DNA template, 12.5 µL of 2 × Taq PCR MasterMix (Tiangen Biotech (Beijing) Co., Ltd.), and 8.5 µL of double distilled water in total of 25 µL. The ITS thermal cycling program was as follows: 94 °C for 5 min, followed by 35 amplification cycles of 94 °C for 60 s, 55 °C for 30 s and 72 °C for 1 min, and a final extension step of 72 °C for 5 min. The annealing temperature at 55 °C for 45 s was changed in this program for *CAL*, *β-tubulin* and *TEF* amplification. For amplification of *HIS*, the program was changed with a cycling program of 32 cycles and an annealing temperature at 55 °C for 60 s. PCR products were visualized on 1.5% agarose gels mixed with Genegreen Nucleic Acid Dye and purified with a quick Midi Purification kit (Tiangen Biotech (Beijing) Co., Ltd.) according to the manufacturer's instructions. Sequencing PCR products were cycle-sequenced the BigDye® Terminator Cycle Sequencing Kit v. 3.1 (Applied Biosystems, Foster City, CA, USA) in an ABI Prism 3730 DNA Sequencer (Applied Biosystems, Foster City, CA, USA) at Biomed Company in Beijing. Then sequence data collected by ABI 3730 Data collection v. 3.0 (Applied Biosystems, Foster City, CA, USA) and ABI Peak Scanner Software v. 1.0 (Applied Biosystems, Foster City, CA, USA), were assembled with forward and reverse sequences by BioEdit. The gene sequences were submitted and awarded access numbers in GenBank of NCBI (Table 1).

Table 1. Access numbers for ITS, translation elongation factor 1-alpha (*TEF1*), beta-tubulin (*TUB*), histone H3 (*HIS*), calmodulin (*CAL*) genes region sequences of the four endophytic *Diaporthe* fungi in the GenBank of NCBI.

Isolate	ITS	<i>TEF1</i>	<i>Tublin</i>	<i>CAL</i>	<i>HIS</i>
FPYF3053	MH203054.1	MH220826.1	MH220836.1	MH220831.1	MH220839.1
FPYF3054	MH203055.1	MH220827.1	MH220833.1	MH220832.1	MH220840.1
FPYF3055	MH203056.1	MH220828.1	MH220834.1	MH220829.1	MH220837.1
FPYF3056	MH203057.1	MH220825.1	MH220835.1	MH220830.1	MH220838.1

2.3. Sequence Alignment and Phylogenetic Analysis

In order to determine the phylogenetic locations of the four isolates within the *Diaporthe* genus, 143 reference taxa [26] (Table 2) together with the four isolates were used for building a phylogenetic tree with *Diaporthella corylina* as a root outgroup species [23]. The evolutionary relationships were taken on a five-gene concatenated alignment of ITS, *TEF1*, *CAL*, *HIS*, and *TUB* regions by maximum likelihood (ML) and maximum parsimony (MP) phylogenetic analyses. Sequences were aligned using the MAFFTv.7 online program with default parameters [27]. A partition homogeneity test implemented in PAUP* v.4.0 (Sinauer Associates, Sunderland, MA, USA) was applied to determine if

the five sequence data could be combined. The best evolutionary model for the partitioning analysis was performed on the concatenated sequences by PartitionFinder 2.1.1 [28]. A concatenated alignment for the five gene regions was made from SequenceMatrix [29]. The inference methods of maximum likelihood and maximum parsimony in Mega 6.0 [30] were applied to estimate phylogeny for the concatenated sequences, with the evolutionary models GTR and AIC for ML and MP, respectively, with a bootstrap support of 1000 replicates. Evidence on the trees were visualized and edited by TreeGraph 2 [31].

Table 2. Reference sequences of *Diaporthe* strains with NCBI access numbers for phylogenetic analysis.

Source	ITS	TEF1	TUB	CAL	HIS
<i>Diaporthe acaciigena</i> _CBS 129521	KC343005.1	KC343731.1	KC343973.1	KC343247.1	KC343489.1
<i>Diaporthe acerina</i> _CBS 137.27	KC343006.1	KC343732.1	KC343974.1	KC343248.1	KC343490.1
<i>Diaporthe alleghaniensis</i> _CBS 495.72	KC343007.1	KC343733.1	KC343975.1	KC343249.1	KC343491.1
<i>Diaporthe alnea</i> _CBS 146.46	KC343008.1	KC343734.1	KC343976.1	KC343250.1	KC343492.1
<i>Diaporthe alnea</i> _CBS 159.47	KC343009.1	KC343735.1	KC343977.1	KC343251.1	KC343493.1
<i>Diaporthe ambigua</i> _CBS 114015	KC343010.1	KC343736.1	KC343978.1	KC343252.1	KC343494.1
<i>Diaporthe ambigua</i> _CBS 117167	KC343011.1	KC343737.1	KC343979.1	KC343253.1	KC343495.1
<i>Diaporthe amygdali</i> _CBS 126679	KC343022.1	KC343742.1	KC343984.1	KC343258.1	KC343506.1
<i>Diaporthe ampelina</i> _CBS 111888	KC343016.1	KC343748.1	KC343990.1	KC343264.1	KC343500.1
<i>Diaporthe amygdali</i> _CBS 111811	KC343019.1	KC343745.1	KC343987.1	KC343261.1	KC343503.1
<i>Diaporthe anacardii</i> _CBS 720.97	KC343024.1	KC343750.1	KC343992.1	KC343266.1	KC343508.1
<i>Diaporthe angelicae</i> _CBS 111592	KC343027.1	KC343753.1	KC343995.1	KC343269.1	KC343511.1
<i>Diaporthe angelicae</i> _CBS 123215	KC343028.1	KC343754.1	KC343996.1	KC343270.1	KC343512.1
<i>Diaporthe cucurbitae</i> _CBS 136.25	KC343031.1	KC343757.1	KC343999.1	KC343273.1	KC343515.1
<i>Diaporthe arecae</i> _CBS 161.64	KC343032.1	KC343758.1	KC344000.1	KC343274.1	KC343516.1
<i>Diaporthe arecae</i> _CBS 535.75	KC343033.1	KC343759.1	KC344001.1	KC343275.1	KC343517.1
<i>Diaporthe arengae</i> _CBS 114979	KC343034.1	KC343760.1	KC344002.1	KC343276.1	KC343518.1
<i>Diaporthe aspalathi</i> _CBS 117169	KC343036.1	KC343762.1	KC344004.1	KC343278.1	KC343520.1
<i>Diaporthe aspalathi</i> _CBS 117168	KC343035.1	KC343761.1	KC344003.1	KC343277.1	KC343519.1
<i>Diaporthe australaficana</i> _CBS 111886	KC343038.1	KC343764.1	KC344006.1	KC343280.1	KC343522.1
<i>Diaporthe australaficana</i> _CBS 113487	KC343039.1	KC343765.1	KC344007.1	KC343281.1	KC343523.1
<i>Diaporthe batatas</i> _CBS 122.21	KC343040.1	KC343766.1	KC344008.1	KC343282.1	KC343524.1
<i>Diaporthe beckhausii</i> _CBS 138.27	KC343041.1	KC343767.1	KC344009.1	KC343283.1	KC343525.1
<i>Diaporthe bicincta</i> _CBS 121004	KC343134.1	KC343860.1	KC344102.1	KC343376.1	KC343618.1
<i>Diaporthe brasiliensis</i> _CBS 133183	KC343042.1	KC343768.1	KC344010.1	KC343284.1	KC343526.1
<i>Diaporthe brasiliensis</i> _LGMF926	KC343043.1	KC343769.1	KC344011.1	KC343285.1	KC343527.1
<i>Diaporthe carpini</i> _CBS 114437	KC343044.1	KC343770.1	KC344012.1	KC343286.1	KC343528.1
<i>Diaporthe caulivora</i> _CBS 127268	KC343045.1	KC343771.1	KC344013.1	KC343287.1	KC343529.1
<i>Diaporthe caulivora</i> _CBS 178.55	KC343046.1	KC343772.1	KC344014.1	KC343288.1	KC343530.1
<i>Diaporthe cestrelaria</i> _CBS 139.27	KC343047.1	KC343773.1	KC344015.1	KC343289.1	KC343531.1
<i>Diaporthe chamaeropis</i> _CBS 454.81	KC343048.1	KC343774.1	KC344016.1	KC343290.1	KC343532.1
<i>Diaporthe chamaeropis</i> _CBS 753.70	KC343049.1	KC343775.1	KC344017.1	KC343291.1	KC343533.1
<i>Diaporthe cinerascens</i> _CBS 719.96	KC343050.1	KC343776.1	KC344018.1	KC343292.1	KC343534.1
<i>Diaporthe citri</i> _CBS 199.39	KC343051.1	KC343777.1	KC344019.1	KC343293.1	KC343535.1
<i>Diaporthe citri</i> _CBS 230.52	KC343052.1	KC343778.1	KC344020.1	KC343294.1	KC343536.1
<i>Diaporthe convolvuli</i> _CBS 124654	KC343054.1	KC343780.1	KC344022.1	KC343296.1	KC343538.1
<i>Diaporthe crataegi</i> _CBS 114435	KC343055.1	KC343781.1	KC344023.1	KC343297.1	KC343539.1
<i>Diaporthe crotalariae</i> _CBS 162.33	KC343056.1	KC343782.1	KC344024.1	KC343298.1	KC343540.1
<i>Diaporthe cuppedae</i> _CBS 117499	KC343057.1	KC343783.1	KC344025.1	KC343299.1	KC343541.1
<i>Diaporthe cynaroidis</i> _CBS 122676	KC343058.1	KC343784.1	KC344026.1	KC343300.1	KC343542.1
<i>Diaporthe decadens</i> _CBS 109772	KC343059.1	KC343785.1	KC344027.1	KC343301.1	KC343543.1
<i>Diaporthe decadens</i> _CBS 114281	KC343060.1	KC343786.1	KC344028.1	KC343302.1	KC343544.1
<i>Diaporthe detrusa</i> _CBS 109770	KC343061.1	KC343787.1	KC344029.1	KC343303.1	KC343545.1
<i>Diaporthe detrusa</i> _CBS 114652	KC343062.1	KC343788.1	KC344030.1	KC343304.1	KC343546.1
<i>Diaporthe elaeagni</i> _CBS 504.72	KC343064.1	KC343790.1	KC344032.1	KC343306.1	KC343548.1
<i>Diaporthe endophytica</i> _CBS 133811	KC343065.1	KC343791.1	KC344033.1	KC343307.1	KC343549.1
<i>Diaporthe endophytica</i> _LGMF928	KC343068.1	KC343794.1	KC344036.1	KC343310.1	KC343552.1
<i>Diaporthe eres</i> _CBS 439.82	KC343090.1	KC343816.1	KC344058.1	KC343332.1	KC343574.1
<i>Diaporthe eres</i> _CBS 101742	KC343073.1	KC343799.1	KC344041.1	KC343315.1	KC343557.1
<i>Diaporthe eres</i> _CBS 109767	KC343075.1	KC343801.1	KC344043.1	KC343317.1	KC343559.1
<i>Diaporthe cf. nobilis</i> RG-2013_CBS 113470	KC343146.1	KC343872.1	KC344114.1	KC343388.1	KC343630.1
<i>Diaporthe cf. nobilis</i> RG-2013_CBS 116953	KC343147.1	KC343873.1	KC344115.1	KC343389.1	KC343631.1

Table 2. Cont.

Source	ITS	TEF1	TUB	CAL	HIS
<i>Diaporthe cf. nobilis</i> RG-2013_CBS 200.39	KC343151.1	KC343877.1	KC344119.1	KC343393.1	KC343635.1
<i>Diaporthe eugeniae</i> _CBS 444.82	KC343098.1	KC343824.1	KC344066.1	KC343340.1	KC343582.1
<i>Diaporthe fibrosa</i> _CBS 109751	KC343099.1	KC343825.1	KC344067.1	KC343341.1	KC343583.1
<i>Diaporthe fibrosa</i> _CBS 113830	KC343100.1	KC343826.1	KC344068.1	KC343342.1	KC343584.1
<i>Diaporthe foeniculacea</i> _CBS 123208	KC343104.1	KC343830.1	KC344072.1	KC343346.1	KC343588.1
<i>Diaporthe foeniculacea</i> _CBS 111553	KC343101.1	KC343827.1	KC344069.1	KC343349.1	KC343585.1
<i>Diaporthe foeniculacea</i> _CBS 187.27	KC343107.1	KC343833.1	KC344075.1	KC343343.1	KC343591.1
<i>Diaporthe ganjae</i> _CBS 180.91	KC343112.1	KC343838.1	KC344080.1	KC343354.1	KC343596.1
<i>Diaporthe gardeniae</i> _CBS 288.56	KC343113.1	KC343839.1	KC344081.1	KC343355.1	KC343597.1
<i>Diaporthe helianthi</i> _CBS 592.81	KC343115.1	KC343841.1	KC344083.1	KC343357.1	KC343599.1
<i>Diaporthe helianthi</i> _CBS 344.94	KC343114.1	KC343840.1	KC344082.1	KC343356.1	KC343598.1
<i>Diaporthe hickoriae</i> _CBS 145.26	KC343118.1	KC343844.1	KC344086.1	KC343360.1	KC343602.1
<i>Diaporthe hongkongensis</i> _CBS 115448	KC343119.1	KC343845.1	KC344087.1	KC343361.1	KC343603.1
<i>Diaporthe hordei</i> _CBS 481.92	KC343120.1	KC343846.1	KC344088.1	KC343362.1	KC343604.1
<i>Diaporthe impulsa</i> _CBS 114434	KC343121.1	KC343847.1	KC344089.1	KC343363.1	KC343605.1
<i>Diaporthe impulsa</i> _CBS 141.27	KC343122.1	KC343848.1	KC344090.1	KC343364.1	KC343606.1
<i>Diaporthe inconspicua</i> _LGMF922	KC343124.1	KC343849.1	KC344091.1	KC343365.1	KC343607.1
<i>Diaporthe inconspicua</i> _CBS 133813	KC343123.1	KC343850.1	KC344092.1	KC343366.1	KC343608.1
<i>Diaporthe infecunda</i> _CBS 133812	KC343126.1	KC343852.1	KC344094.1	KC343368.1	KC343610.1
<i>Diaporthe infecunda</i> _LGMF933	KC343132.1	KC343858.1	KC344100.1	KC343374.1	KC343616.1
<i>Diaporthe longispora</i> _CBS 194.36	KC343135.1	KC343861.1	KC344103.1	KC343377.1	KC343619.1
<i>Diaporthe lusitanicae</i> _CBS 123212	KC343136.1	KC343862.1	KC344104.1	KC343378.1	KC343620.1
<i>Diaporthe lusitanicae</i> _CBS 123213	KC343137.1	KC343863.1	KC344105.1	KC343379.1	KC343621.1
<i>Diaporthe manihotia</i> _CBS 505.76	KC343138.1	KC343864.1	KC344106.1	KC343380.1	KC343622.1
<i>Diaporthe mayteni</i> _CBS 133185	KC343139.1	KC343865.1	KC344107.1	KC343381.1	KC343623.1
<i>Diaporthe megalospora</i> _CBS 143.27	KC343140.1	KC343866.1	KC344108.1	KC343383.1	KC343624.1
<i>Diaporthe melonis</i> _CBS 507.78	KC343142.1	KC343868.1	KC344110.1	KC343384.1	KC343626.1
<i>Diaporthe melonis</i> _CBS 435.87	KC343141.1	KC343867.1	KC344109.1	KC343382.1	KC343625.1
<i>Diaporthe musigena</i> _CBS 129519	KC343143.1	KC343869.1	KC344111.1	KC343385.1	KC343627.1
<i>Diaporthe neilliae</i> _CBS 144.27	KC343144.1	KC343870.1	KC344112.1	KC343386.1	KC343628.1
<i>Diaporthe neocarctii</i> _CBS 109490	KC343145.1	KC343871.1	KC344113.1	KC343387.1	KC343629.1
<i>Diaporthe nomurai</i> _CBS 157.29	KC343154.1	KC343880.1	KC344122.1	KC343396.1	KC343638.1
<i>Diaporthe novem</i> _CBS 127270	KC343156.1	KC343882.1	KC344124.1	KC343398.1	KC343640.1
<i>Diaporthe novem</i> _CBS 354.71	KC343158.1	KC343884.1	KC344126.1	KC343400.1	KC343642.1
<i>Diaporthe oncostoma</i> _CBS 109741	KC343161.1	KC343887.1	KC344129.1	KC343403.1	KC343645.1
<i>Diaporthe oncostoma</i> _CBS 100454	KC343160.1	KC343886.1	KC344128.1	KC343402.1	KC343644.1
<i>Diaporthe oxe</i> _CBS 133186	KC343164.1	KC343890.1	KC344132.1	KC343406.1	KC343648.1
<i>Diaporthe oxe</i> _CBS 133187	KC343165.1	KC343891.1	KC344133.1	KC343407.1	KC343649.1
<i>Diaporthe padi</i> var. <i>padi</i> _CBS 114200	KC343169.1	KC343895.1	KC344137.1	KC343411.1	KC343653.1
<i>Diaporthe padi</i> var. <i>padi</i> _CBS 114649	KC343170.1	KC343896.1	KC344138.1	KC343412.1	KC343654.1
<i>Diaporthe paranensis</i> _CBS 133184	KC343171.1	KC343897.1	KC344139.1	KC343413.1	KC343655.1
<i>Diaporthe perjuncta</i> _CBS 109745	KC343172.1	KC343898.1	KC344140.1	KC343414.1	KC343656.1
<i>Diaporthe perseae</i> _CBS 151.73	KC343173.1	KC343899.1	KC344141.1	KC343415.1	KC343657.1
<i>Diaporthe phaselorum</i> _CBS 116019	KC343175.1	KC343901.1	KC344143.1	KC343417.1	KC343659.1
<i>Diaporthe phaselorum</i> _CBS 116020	KC343176.1	KC343902.1	KC344144.1	KC343418.1	KC343660.1
<i>Diaporthe pseudomangiferae</i> _CBS 101339	KC343181.1	KC343907.1	KC344149.1	KC343423.1	KC343665.1
<i>Diaporthe pseudomangiferae</i> _CBS 388.89	KC343182.1	KC343908.1	KC344150.1	KC343424.1	KC343666.1
<i>Diaporthe pseudophoenicicola</i> _CBS 462.69	KC343184.1	KC343910.1	KC344152.1	KC343426.1	KC343668.1
<i>Diaporthe pseudophoenicicola</i> _CBS 176.77	KC343183.1	KC343909.1	KC344151.1	KC343425.1	KC343667.1
<i>Diaporthe pustulata</i> _CBS 109784	KC343187.1	KC343913.1	KC344155.1	KC343429.1	KC343671.1
<i>Diaporthe pustulata</i> _CBS 109742	KC343185.1	KC343911.1	KC344153.1	KC343427.1	KC343669.1
<i>Diaporthe raoniakayaporum</i> _CBS 133182	KC343188.1	KC343914.1	KC344156.1	KC343430.1	KC343672.1
<i>Diaporthe rhoina</i> _CBS 146.27	KC343189.1	KC343915.1	KC344157.1	KC343431.1	KC343673.1
<i>Diaporthe saccarata</i> _CBS 116311	KC343190.1	KC343916.1	KC344158.1	KC343432.1	KC343674.1
<i>Diaporthe schini</i> _CBS 133181	KC343191.1	KC343917.1	KC344159.1	KC343433.1	KC343675.1
<i>Diaporthe schini</i> _LGMF910	KC343192.1	KC343918.1	KC344160.1	KC343434.1	KC343676.1
<i>Diaporthe sclerotioides</i> _CBS 296.67	KC343193.1	KC343919.1	KC344161.1	KC343435.1	KC343677.1
<i>Diaporthe sclerotioides</i> _CBS 710.76	KC343194.1	KC343920.1	KC344162.1	KC343436.1	KC343678.1
<i>Diaporthe scobina</i> _CBS 251.38	KC343195.1	KC343921.1	KC344163.1	KC343437.1	KC343679.1
<i>Diaporthe sojae</i> _CBS 100.87	KC343196.1	KC343922.1	KC344164.1	KC343438.1	KC343680.1
<i>Diaporthe longicolla</i> isolate PL4	HM347700.1	HM347685.1	KC344167.1	KC343441.1	KC343683.1
<i>Diaporthe sojae</i> _CBS 116017	KC343197.1	KC343923.1	KC344165.1	KC343439.1	KC343681.1
<i>Diaporthe sojae</i> _CBS 180.55	KC343200.1	KC343926.1	KC344168.1	KC343442.1	KC343684.1
<i>Diaporthe subordinaria</i> _CBS 101711	KC343213.1	KC343938.1	KC344180.1	KC343454.1	KC343696.1
<i>Diaporthe subordinaria</i> _CBS 464.90	KC343214.1	KC343939.1	KC344181.1	KC343455.1	KC343697.1

Table 2. Cont.

Source	ITS	TEF1	TUB	CAL	HIS
<i>Diaporthe tecomae</i> _CBS 100547	KC343215.1	KC343940.1	KC344182.1	KC343456.1	KC343698.1
<i>Diaporthe terebinthifoliae</i> _CBS 133180	KC343216.1	KC343941.1	KC344184.1	KC343457.1	KC343699.1
<i>Diaporthe terebinthifoliae</i> _LGMF907	KC343217.1	KC343942.1	KC344183.1	KC343458.1	KC343700.1
<i>Diaporthe toxica</i> _CBS 534.93	KC343220.1	KC343943.1	KC344185.1	KC343459.1	KC343701.1
<i>Diaporthe toxica</i> _CBS 535.93	KC343221.1	KC343946.1	KC344188.1	KC343462.1	KC343704.1
<i>Diaporthe vaccinii</i> _CBS 160.32	KC343228.1	KC343947.1	KC344189.1	KC343463.1	KC343705.1
<i>Diaporthe vaccinii</i> _CBS 122112	KC343224.1	KC343954.1	KC344196.1	KC343470.1	KC343712.1
<i>Diaporthe vexans</i> _CBS 127.14	KC343229.1	KC343950.1	KC344192.1	KC343466.1	KC343708.1
<i>Diaporthe rufida</i> _CBS 113201	KC343234.1	KC343955.1	KC344197.1	KC343471.1	KC343713.1
<i>Diaporthe rufida</i> _CBS 109768	KC343233.1	KC343960.1	KC344202.1	KC343476.1	KC343718.1
<i>Diaporthe woodii</i> _CBS 558.93	KC343244.1	KC343959.1	KC344201.1	KC343475.1	KC343717.1
<i>Diaporthe woolworthii</i> _CBS 148.27	KC343245.1	KC343970.1	KC344212.1	KC343486.1	KC343728.1
<i>Diaporthe</i> cf. <i>heveae</i> 1 RG-2013_CBS 852.97	KC343116.1	KC343971.1	KC344213.1	KC343487.1	KC343729.1
<i>Diaporthe</i> cf. <i>heveae</i> 2 RG-2013_CBS 681.84	KC343117.1	KC343842.1	KC344084.1	KC343358.1	KC343600.1
<i>Diaporthe</i> sp. 1 RG-2013_CBS 119639	KC343202.1	KC343843.1	KC344085.1	KC343359.1	KC343601.1
<i>Diaporthe</i> sp. 1 RG-2013_LGMF947	KC343203.1	KC343928.1	KC344170.1	KC343444.1	KC343686.1
<i>Diaporthe</i> sp. 2 RG-2013_LGMF932	KC343204.1	KC343929.1	KC344171.1	KC343445.1	KC343687.1
<i>Diaporthe</i> sp. 3 RG-2013_CBS 287.29	KC343205.1	KC343930.1	KC344172.1	KC343446.1	KC343688.1
<i>Diaporthe</i> sp. 4 RG-2013_LGMF944	KC343206.1	KC343931.1	KC344173.1	KC343448.1	KC343689.1
<i>Diaporthe</i> sp. 5 RG-2013_CBS 125575	KC343207.1	KC343932.1	KC344174.1	KC343447.1	KC343690.1
<i>Diaporthe</i> sp. 6 RG-2013_CBS 115584	KC343208.1	KC343933.1	KC344175.1	KC343449.1	KC343691.1
<i>Diaporthe</i> sp. 6 RG-2013_CBS 115595	KC343209.1	KC343934.1	KC344176.1	KC343450.1	KC343692.1
<i>Diaporthe</i> sp. 7 RG-2013_CBS 458.78	KC343210.1	KC343935.1	KC344177.1	KC343451.1	KC343693.1
<i>Diaporthe</i> sp. 8 RG-2013_LGMF925	KC343211.1	KC343936.1	KC344178.1	KC343452.1	KC343694.1
<i>Diaporthella corylina</i> _CBS 121124	KC343004.1	KC343937.1	KC344179.1	KC343453.1	KC343695.1
<i>Diaporthe stictica</i> _CBS 370.54	KC343212.1	KC343730.1	KC343972.1	KC343246.1	KC343488.1

2.4. Antifungal Activity Tests for Fungal VOCs

The antifungal activity of the VOCs was determined by the methods previously described [17,18,20]. The four endophytic fungal strains of *Diaporthe* and targeted plant pathogenic microorganisms were paired opposite to each other in Petri plates containing PDA with a diameter of 90 mm. The agar was divided into two halves by removing a 2 cm wide strip in the center. An endophytic test fungus was inoculated onto one half-moon of the agar and incubated at 25 °C for five days for optimum production of volatile compounds before the antagonism bioassay. A test pathogen was inoculated onto the opposite half-moon part of the agar at the fifth day. The plates were then wrapped with parafilm and incubated at 25 °C in dark for 72 h. Growth of filamentous pathogenic fungi were quantitatively assessed after 24 h, 48 h, and 72 h based on multiple measurements of growth relative to controls, as described previously [17,18]. The colony diameter was measured in an average of four diameters on hours 24, 48, and 72 h, disregarding the initial inoculum size. Percentage of growth inhibition was calculated as the formula: $| (a - b) / b | \times 100$, a = mycelial colony diameter in control plate; b = mycelial colony diameter in the antagonism treatment plate. Statistical significance ($p < 0.01$) was evaluated by analysis of variance (ANOVA) followed by the Tukey 5% test. Antifungal activity of VOCs was tested against the plant pathogenic fungi *Alternaria alternata*, *Botryosphaeria dothidea*, *Botrytis cinerea*, *Cercospora* sp., *Colletotrichum gloeosporioides*, *Fusarium graminearum*, *Sphaeropsis sapinea*, and *Valsa sordida*, in addition to the oomycete *Phytophthora cinnamomi*. All tests were made in quintuplicate. Control cultures were obtained by growing each plant pathogen alone, under the same conditions.

2.5. Qualitative Analyses on Volatiles of the Four Endophytic Cultures

VOCs in the air space above the endophytic fungal colonies grown for five days at 25 ± 2 °C on PDA were analyzed using the solid phase microextraction (SPME) fiber technique according to previously described protocols [17,18,20]. Control PDA Petri plates not inoculated with the strain was used to subtract compounds contributed by the medium. All treatments and checks were done in triplicate. A fiber syringe of 50/30 divinylbenzene/carboxen on polydimethylsiloxane (Supelco, Bellefonte, PA, USA) was conditioned for 40 min at 200 °C, exposed to the vapor phase inside Petri

during 40 min through a small hole (0.5 mm in diameter) drilled on the sides of the Petri plate. The fiber was directly inserted into the TRACE DSQ inlet (Thermo Electron Corporation, Beverly, MA, USA), at 200 °C, splitless mode. The desorption time was 40 s and the desorbed compounds were separated on a 30.0 m × 0.25 mm × 0.25 µm, HP-5MS capillary column, using the following GC oven temperature program: 2 min at 35 °C up to 220 °C at 7 °C/min. Helium was used as the carrier gas at a flow rate of 1 mL/min. The electronic ionization energy was 70 eV and the mass range scanned was 41–560 um. The scan rate was 5 spec/s. Transfer line and ionization chamber temperatures were 250 °C and 200 °C respectively. Tentative identification of the volatile compounds produced by the four endophytic *Diaporthe* fungi was made via library comparison using the NIST database and all chemical compounds were described in this report following the NIST database chemical identity. Tentative compound identity was based on at least a 70% quality match with the NIST database information for each compound. Data acquisition and data processing were performed with the Hewlett Packard ChemStation software system (Version 2.0, Scientific Instrument Services, Inc., Ringoes, NJ, USA). Relative amounts of individual components of the treatments were determined and expressed as percentages of the peak area within the total peak area and as an average of the three replicates.

3. Results

3.1. The Identification on the Four Endophytic Isolates within the *Diaporthe* Genus

Each of the four isolates falling within the genus *Diaporthe* were further defined using molecular analyses as they appeared different, morphologically (Figure 1). For instance, strain FPYF3053 had compact mycelia with crenate margins, these colonies developed a brownish yellow pigmentation in the center on the underside having a growth rate of 18.3 mm day⁻¹ (Figure 1a). On the other hand, strain FPYF3054 had aerial mycelium forming concentric rings with grey and dark pigmentation at the center showing a growth rate of 30.97 mm day⁻¹ (Figure 1b). Strain FPYF3055 had vigorously-growing aerial hyphae near the margin, but loose hyphae scattered inside with aging, with a growth rate of 23 mm day⁻¹ (Figure 1c). Finally, strain FPYF3056 had a compact mycelium with a crenate margin, but no pigmentation with a growth rate of 21.7 mm day⁻¹ (Figure 1d).

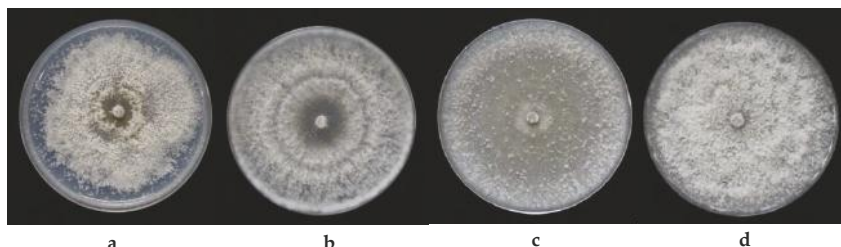


Figure 1. The colony cultures for the four endophytic *Diaporthe* fungi and their plant host. (a) FPYF3053; (b) FPYF3054; (c) FPYF3055; and (d) FPYF3056.

A combined alignment of five loci *ITS*, *TUB*, *TEF1*, *HIS*, and *CAL* was used for ML and MP phylogenetic analyses. Based on the multi-locus phylogeny (Figure 2), the four endophytic *Diaporthe* strains could not be placed in one species only because they are distinct from each other and from all reference species listed (Table 2, Figure 2). Strains FPYF3055 and FPYF3056 were clustered by giving a high bootstrap support (BS = 82) from MP inference (Figure S1) while both separated from each other in ML inference (Figure S2). The reference sequences used to construct the phylogenetic tree were listed in Table 2 with their Genbank accession numbers. The alignment was uploaded in Treebase assigned with SI 22757.

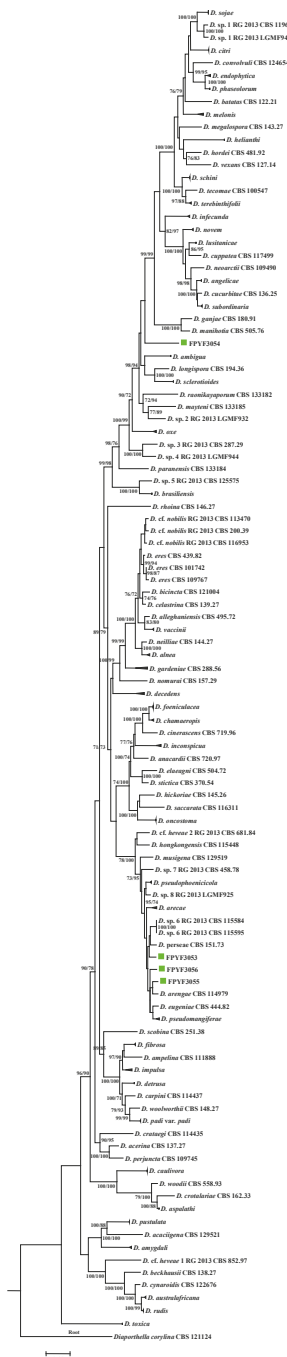


Figure 2. Phylogenetic tree based on combinedITS, CAL, TEF1, HIS, and TUB sequence alignment generated from a maximum parsimony and maximum likelihood analyses. Values near the branches represent parsimony/likelihood bootstrap support values (>70%), respectively. The tree is rooted with *Diaporthella corylina*. The four endophytic isolates were each named with strain ID marked green box. Compressed branches were used for saving space. The complete phylogenetic trees of MP and ML can be found in Figures S1 and S2, respectively. 46

3.2. The VOCs' Bioactivities of the Four *Diaporthe* Strains against Plant Fungal Pathogens

All of the four strains were observed to inhibit the growth of nine selected fungal pathogens by producing volatile compounds in the PDA medium (Table 3). The nine pathogens, *Alternaria alternata*, *Botryosphaeria dothidea*, *Botrytis cinerea*, *Cercospora asparagi*, *Colletotrichum gloeosporioides*, *Fusarium graminearum*, *Phytophthora cinnamomi*, *Sphaeropsis sapinea*, and *Valsa sordida*, are important causal agents to major trees, such as poplars and pines, or agricultural crops in China and elsewhere. All FPYF strains showed different inhibitory activities along the measurements, an exception was observed for the case of strain FPYF3053, which promoted the growth of *Phytophthora cinnamomi* (Table 3). Furthermore, all selected pathogens, except *V. sordid*, achieved obvious growth inhibition over around 10% during the testing period. After 24 h, *B. cinerea* was the most sensitive to VOCs emitted by all endophytic strains, reaching percent inhibitions of more than 55% when dual cultured with each strain. *B. dothidea* and *A. alternata* were highly sensitive to VOCs of all the endophytic strains, getting percent inhibitions of more than 30% with an exception to 28% of *A. alternata* in VOCs of the strain FPYF3053. *V. sordida* had the least sensitive or insensitive performance in VOCs from all the strains, showing percent inhibitions around 3% when dual cultured with FPYF3056. The inhibitive intensity of FPYF strains' VOCs on growth of pathogens decreased in times to most dual cultures. The maximum drop of the intensity was by 31% in percent inhibition on the pathogen *B. cinerea* dual culturing with strain FPYF3056. The obvious increase in intensity occurred in the pathogen *F. graminearum* dual culturing with FPYF3055 and FPYF3056, increasing by around 10% during 72 h. Some pathogens grew fast without percent inhibition records after 24 h (*V. sordida*) or 72 h (*B. dothidea* and *F. graminearum*).

Table 3. Growth inhibition percentage of plant pathogens by VOC bioassays of four *Diaporthe* strains.

Percent of inhibition is shown as the means of four measurements of diameters with standard deviation ($n = 4$).

Pathogen	Day	FPYF3053		FPYF3054		FPYF3055		FPYF3056	
		Percentage Inhibition	p-Value						
<i>Alternaria alternata</i>	24 h	28.77 ± 2.26	0.0003	41.41 ± 1.65	0.0001	37.68 ± 5.6	0.0078	34.51 ± 2.03	0.0002
	48 h	22.42 ± 2.34	0.0003	30.19 ± 1.56	0.0009	30.25 ± 5.12	0.0113	26.50 ± 3.42	0.0006
	72 h	15.25 ± 2.59	0.0019	23.43 ± 2.27	0.0039	22 ± 4.03	0.0148	16.34 ± 2.36	0.0010
<i>Botryosphaeria dothidea</i>	24 h	46.3 ± 4.30	0.0030	50.17 ± 2.43	0.0006	43.74 ± 2.15	0.0000	37.88 ± 3.80	0.0002
	48 h	45.28 ± 2.63	0.0000	45.14 ± 2.35	0.0000	42.78 ± 0.43	0.0000	38.99 ± 0.98	0.0000
	72 h	NE *		NE *		NE *		NE *	
<i>Botrytis cinerea</i>	24 h	64.47 ± 1.05	0.0000	55.26 ± 4.82	0.0000	60.72 ± 1.91	0.0001	55.26 ± 4.71	0.0000
	48 h	50.42 ± 1.79	0.0000	35.02 ± 1.22	0.0000	39.44 ± 3.70	0.0000	32.41 ± 3.75	0.0003
	72 h	36.55 ± 2.81	0.0000	24.27 ± 3.08	0.0001	30.10 ± 3.54	0.0005	24.25 ± 4.62	0.0025
<i>Cercospora asparagi</i>	24 h	31.46 ± 4.11	0.0003	22.64 ± 2.86	0.0074	23.21 ± 4.54	0.0202	18.57 ± 5.01	0.0086
	48 h	33.81 ± 2.97	0.0000	24.32 ± 2.21	0.0008	22.02 ± 2.96	0.0000	16.34 ± 1.53	0.0000
	72 h	23.32 ± 2.17	0.0007	19.09 ± 2.73	0.0031	11.94 ± 3.54	0.0083	4.56 ± 0.85	0.0032
<i>Colletotrichum gloeosporioides</i>	24 h	26.83 ± 4.78	0.0153	9.75 ± 2.33	0.0009	10.68 ± 1.14	0.0001	10.74 ± 2.38	0.0229
	48 h	20.94 ± 3.33	0.0051	11.76 ± 2.35	0.0006	8.91 ± 1.24	0.0001	9.78 ± 2.36	0.0017
	72 h	20.68 ± 1.56	0.0024	9.39 ± 2.78	0.0023	6.29 ± 0.80	0.0001	6.82 ± 2.01	0.0388
<i>Fusarium graminearum</i>	24 h	25.68 ± 1.13	0.0031	14.68 ± 2.05	0.0147	20.45 ± 3.62	0.0046	12.96 ± 1.42	0.0103
	48 h	29.99 ± 5.29	0.0086	12.9 ± 4.50	0.0284	31.12 ± 3.57	0.0203	21.59 ± 4.57	0.0425
	72 h	NE		NE		NE		NE	
<i>Phytophthora cinnamomi</i>	24 h	-5.01 ± 1.14 **	0.0029	19.21 ± 4.54	0.0036	31.02 ± 2.58	0.0001	8.38 ± 3.10	0.0154
	48 h	-15.65 ± 6.36	0.0186	12.19 ± 3.30	0.0500	25.21 ± 4.29	0.0050	11.32 ± 4.22	0.0302
	72 h	-19.70 ± 4.19	0.0153	11.91 ± 2.12	0.0209	21.03 ± 2.80	0.0031	8.94 ± 2.03	0.0013
<i>Sphaeropsis sapinea</i>	24 h	23.39 ± 4.25	0.0147	22.69 ± 5.23	0.0239	21.84 ± 7.61	0.0491	7.41 ± 2.68	0.0364
	48 h	20.93 ± 1.04	0.0009	23.85 ± 1.68	0.0023	18.33 ± 5.22	0.0367	9.53 ± 0.60	0.0024
	72 h	NE		NE		NE		NE	
<i>Valsa sordida</i>	24 h	5.96 ± 1.61	0.0115	9.73 ± 2.79	0.0014	5.14 ± 1.02	0.0153	3.15 ± 1.00	0.0177
	48 h	NE		NE		NE		NE	
	72 h	NE		NE		NE		NE	

* No data. ** Negative values mean growth stimulation.

Table 4. Chemical composition of volatiles obtained from mycelial cultures of the four endophytic *Diaporthe* fungi using solid-phase microextraction (SPME).

Retention Time (min)	Molecular Weight	Compound	Quality (%) ^a				Abundance (Relative) ^b
			FPTF3053	FPTF3054	FPTF3055	FPTF3056	
6.17	106	Ethylbenzene	91.8	75.5	77.9	0.42	0.80
7.67	136	α-Thujene	91.9	89.3	30.57	37.10	36.19
9.88	136	1,3-Cyclohexadiene, 1-methyl-4-(1-methyl ethyl)-2-Carene	84.7	80.2	6.26	5.92	1.87
9.89	136	α-Phellandrene	86	74.9	84.9	1.80	6.20
10.21	136	β-Phellandrene	88.4	90.9	75.6	0.74	5.15
10.22	136	γ-Terpinene	89.4	85.6	87.7	12.55	2.35
10.92	136	Cyclohexene, 1-methyl-4-(1-methylethylidene)-2-Cyclohexen-1-ol, 1-methyl-4-(1-methylethyl)-Unknown	81	81.1	88.7	21.15	18.70
11.63	136	Thujone	63.4	68.2	0.65	1.66	19.21
11.89	154	1-Menthone	90.4	71.5	0.29	1.24	1.24
12.29	152	Cyclohexanol, 5-methyl-2-(1-methylethyl)-3-Cyclohexen-1-ol, 4-methyl-1-(1-methylethyl)-(R)-2,4,4-Trimethyl-3-(3-methylbutyl)cyclohex-2-enone	88.5	89.7	85.6	0.44	0.65
12.3	154	Unknown	70.1	66.3	0.28	22.38	5.88
13.21	154	Unknown	64.9	64.4	0.28	27.91	10.29
13.68	156	Unknown	64.9	64.4	0.28	1.35	0.31
13.76	154	Biphenylene, 1,2,3,6,7,8,8a-octahydro-4,5-dimethyl-Caryphyllene	79	73.9	60.9	1.62	2.06
13.91	208	Unknown	73.6	68.5	72.2	0.46	6.81
14.12	352	Patelloiene	69.7	69.7	0.31	0.45	0.59
14.13	240	Unknown	76.8	63.1	1.11	2.51	0.31
14.14	170	Unknown	64.9	61.6	11.54	3.17	0.41
17.42	388	Unknown	64.9	64.4	0.28	2.26	2.26
18.41	188	Cedrene	78.2	81.1	81	0.41	0.41
18.92	204	α-Murolene	92.2	92.2	0.41	0.41	0.41
19.11	204	Unknown	64.9	64.4	0.28	2.26	2.26
19.74	204	Cedrene	78.2	81.1	81	0.41	0.41
20.01	222	Unknown	64.9	64.4	0.28	2.26	2.26
20.13	204	Patelloiene	76.8	63.1	1.11	2.51	0.31
20.46	204	Unknown	64.9	64.4	0.28	2.26	2.26

Notes: Data are averages of two cultures grown on the same medium with subtracting those from the control PDA plate. ^aThe quality match is the % likelihood that the compound is identical to that which is listed on the table based on the NIST database. Compounds assigned as unknown with lower than 70% quality match. ^bThe abundance figure presents the percentage amount of each compound in total area relative to all listed compounds detected for one strain.

3.3. The Qualification on VOCs of the Four Endophytic *Diaporthe* Strains

Each of the *Diaporthe* isolates showed a unique VOC profile as measured by SPME (Table 4). Nineteen VOC components from the four fungi were identified and seven compounds were unidentified according our set standard of a 70% quality match with the GC-MS. Generally, the terpenoids were the major components in the VOCs of each strain. The main terpenes included α -thujene, β -phellandrene, γ -terpinene, l-menthone, cyclohexanol, 5-methyl-2-(1-methylethyl)-, α -muurolene. The amounts of each component of these monoterpenes had a relative area over 10% of the total of its VOCs. There also existed other minor terpenoids at very low amounts, including carene, α -phellandrene, thujone, caryophyllene, patchoulene, etc. Two monoterpenes, β -phellandrene and α -muurolene, and a chemical biphenylene, 1,2,3,6,7,8,8a,8b-octahydro-4,5-dimethyl, which were detected in VOCs of all four strains. Four chemicals were common to VOCs from FPYF3053, FPYF3055, and FPYF3056, including α -thujene, 1,3-cyclohexadiene, 1-methyl-4-(1-methylethyl)-, γ -terpinene and 3-cyclohexen-1-ol, and 4-methyl-1-(1-methylethyl)-(R)-. However, each strain produced a unique mixture of volatile organic compounds. The strain FPYF3053 produced 15 volatile compounds with three prominent components, α -thujene, β -phellandrene, and α -muurolene. FPYF3054 was able to synthesize eight compounds with three prominent components of β -phellandrene, l-menthone, and cyclohexanol, 5-methyl-2-(1-methylethyl)- in VOC mixtures. Strains FPYF3055 and 3056 generated relatively close chemical compositions in amount and quality of VOCs compared to FPYF3053 and FPYF3054. However, FPYF3055 had three prominent components, α -thujene, γ -terpinene, and 3-cyclohexen-1-ol, 4-methyl-1-(1-methylethyl)-(R)-, in 12 compounds of the VOCs, while FPYF3056 had three prominent components—namely α -thujene, β -phellandrene, and γ -terpinene—of 13 compounds in its VOCs.

4. Discussion

4.1. Endophytic *Diaporthe* spp. from *Catharanthus roseus*

Four isolates of endophytes in the genus *Diaporthe* were obtained from the medicinal plant *Catharanthus roseus* growing in a conservation area of Southern China. In order to best distinguish these individual organisms they were subjected to a combined analysis of five-loci alignment of *TEF1-TUB-CAL-HIS-ITS* which gave a more robust isolate identification [23]. Adding our four endophytic isolates did not affect the congruency in each locus, partition homogeneity for the combination and the best evolutionary model for the five-locus concatenated alignment reported. *Diaporthe* fungi are one of the most common endophytic fungal communities found in plants [11]. However, the previous work on endophytic fungi from *C. roseus* [7,8,32–42] did not record strains of the *Diaporthe* genus. *Alternaria alternata* was determined as the dominant endophytic species in leaf tissue of *C. roseus* along with associated fungi from the following genera, *Aspergillus*, *Fusarium*, *Penicillium*, and *Helminthosporium* [33]. In addition the endophytes of root tissue appeared including *Colletotrichum* sp., *Macrophomina phaseolina*, *Nigrospora sphaerica*, and *Fusarium solani* [7]. Other isolated endophytic fungi from this plant included *Colletotrichum truncatum*, *Drechslera* sp., *Cladosporium* sp., and *Myrothecium* sp. [43]. To our four *Diaporthe* strains, no reproductive structures were obtained in the employed conditions. They were designated *Diaporthe* sp. strains (FPYF3053–3056) without spore characterization strictly using phylogenetic analysis. The strains seemed not to share a close phylogenetic relationship to any other species based on the five-locus alignment study (Figure 2, [12,23,26]). The robust inference on the strains will take place when fruit bodies appear combined with full species phylogeny in the genus *Diaporthe*.

4.2. VOCs Antifungal Effects of Endophytic *Diaporthe* spp. from *Catharanthus roseus*

Compounds extracted from *Catharanthus roseus* [4,5] and extracts from some endophytes of this plant [10,44] have been shown to have antimicrobial bioactivities to some human microbial pathogens and plant fungal pathogens, including *Staphylococcus aureus*, *Pseudomonas aeruginosa*, *Bacillus subtilis*,

Escherichia coli, *Aspergillus fumigatus*, *Candida albicans*, etc. However, the VOCs or essential oils from *Catharanthus roseus* in the literature is scarce results on antimicrobial activities [45,46]. The previous work on the other endophytic fungi of this host plant did not consider that VOCs of the endophyte may have antimicrobial activities [7,8,32–42]. However, this work shows that VOCs produced by four endophytic *Diaporthe* fungi from the plant are able to functionally inhibit the growth of a number of specifically-targeted fungal pathogens (Table 3).

In the past there have been three endophytic *Diaporthe* strains recorded with their VOCs [18,19,47]. Two of them were reported to be inhibitory to plant pathogens [18,19]. One strain PR4 was isolated from a medicinal plant growing in Kashmir, Himalayas [19]; the other strain EC-4 was isolated from *Odontoglossum* sp. in Northern Ecuador [18]. With our four strains, the volatile compounds from endophytic *Diaporthe* fungi varied in degrees of inhibition against selected pathogenic fungi and test timings depending on the endophytic strain (Tables 3 and 5). However, the maximal inhibition of fungal growth of *Diaporthe* was from strain PR4, which reduced growth of *Rhizoctonia solani* by 100%. FPYF strains' and EC-4 VOCs also appeared effective in the inhibition of growth of *Botrytis cinerea* by more than 30% with a maximal of $50.42 \pm 1.8\%$. During the test course of 72 h, to most cases, FPYF strains' VOCs showed strong bioactivities in the first day and then decreased inhibition on the pathogens in following two days (Table 3). PR4 VOCs were effective in reducing radial growth of *Pythium ultimum* by 13.3%; EC-4 VOCs were effective in reducing radial growth of *Pythium ultimum*, *Phytophthora cinnamomi*, and *Phytophthora palmivora* by $59.1 \pm 0.9\%$, $42.0 \pm 0.5\%$, and $5.6 \pm 0.5\%$, respectively. FPYF3054-3056's VOCs were effective against *Phytophthora cinnamomi* in a range of 25.21 ± 4.3 ~ $11.32 \pm 4.2\%$. The alcohol compounds such as 1-propanol, 2-methyl- and 1-butanol, 3-methyl- might make the oomycete *P. cinnamomi* more sensitive to EC-4's VOCs [18], which were lack in VOCs of all FPYF strains (Table 4). The two alcohol compounds had antimicrobial activities in VOCs of endophytic *Phomopsis* sp. strain EC-4 [18]. The sensitivity of the pathogen *F. graminearum* to VOCs from *Diaporthe* spp. might be analogous even though the VOCs components were not similar among *Diaporthe* strains. Two *Diaporthe* strains FPYF3053, 3055 (Table 2) and *Diaporthe* strain PR4 [19] had percent inhibition of *F. graminearum* growth of around 30% under their VOCs bioactivities. However, only beta-phellandrene was a common compound found in VOCs among them (Table 4, [19]). Contrast to cytochalasins as a predominantly common component in soluble secondary metabolites of *Diaporthe* strains [16], the genus-specific or predominant conserved components of fungal VOCs of genus *Diaporthe* should be proposed to illustrate further. The experimental data suggests that the VOCs of FPYF strains are both biologically active and biologically selective. Finally, isolate FPYF3053 were showed no effective inhibition of *Phytophthora cinnamomi* growth. In this study, we attempt to understand the VOCs inhibitory impacts from the endophytic *Diaporthe* strains without consideration of interaction between the strains and pathogens. Future research is proposed to investigate the dual interaction in the VOCs' levels and other molecules between fungal interactions [48].

The headspace analyses of the four *Diaporthe* strains in potato dextrose medium revealed that three monoterpenes— β -phellandrene, biphenylene, 1,2,3,6,7,8,8a,8b-octahydro-4,5-dimethyl and α -muurolene—seemed to be characteristic compounds of endophytic *Diaporthe* strains endophytic to *Catharanthus roseus*. However, among all monoterpenes mentioned above, only 1-menthone can be found in volatile compounds of *Catharanthus roseus* flowers, the essential oil of which is high in limonene and other monoterpenes [45,46]. Menthol and β -phellandrene were also found in VOCs of *Diaporthe* strain PR4 with very low relative amounts of less than 1.0% [19]. No chemicals were shared in VOCs between our FPYF strains and *Phomopsis* strain EC-4 (Table 4, [18]). Therefore, the antifungal VOCs from the four endophytic *Diaporthe* Chinese strains possesses unique VOC compositions compared with known *Diaporthe* VOCs. Although many fungi were reported to produce many terpene compounds in their VOCs [49], our *Diaporthe* fungi maybe of some interest as a source of some other monoterpenes, which often only have been thought to originate from specific plants. For instance, essential oils from many plants containing more or less such monoterpenes as α -thujene, β -phellandrene [50–52], γ -terpinene [53,54], l-menthone [55,56], cyclohexanol, α -muurolene, thujone, and caryophyllene have

some antifungal activities. For example, γ -terpinene, singly or in mixtures with sabinene in oil from coastal redwood leaves, has strong antifungal activity on some endophytic fungi [53]. Therefore, it could be rational to infer the terpenes in FPYF strains synergistically played a main role in their inhibition pathogenic fungi growths. In addition, the high content of monoterpenes in the *Diaporthe* VOCs does have potential for the biofuel industry [18,20,57].

Table 5. Comparison VOCs' inhibitive effect among *Diaporthe* strains.

Pathogens	Percent Growth Inhibition		
	<i>Phomopsis</i> sp. EC-4 [17] *	<i>Diaporthe</i> Strain PR4 [18]	FPYF3053-3056 **
<i>Aspergillus flavus</i>	/ ***	34.6	/
<i>Aspergillus fumigatus</i>	57.0 ± 0.5	/	/
<i>Alternaria alternata</i>	/	/	30.25 ± 5.1~22.42 ± 2.3
<i>Botryosphaeria dothidea</i>	/	/	45.14 ± 2.4~25.28 ± 2.6
<i>Botryotinia cinerea</i>	37.8 ± 0.5	/	50.42 ± 1.8~32.41 ± 3.8
<i>Ceratocystis fimbriata</i>	/	0.0	/
<i>Ceratocystis ulmi</i>	11.1 ± 1.5	/	/
<i>Cercospora asparagi</i>	/	/	33.81 ± 2.97~16.34 ± 1.5
<i>Cercospora beticola</i>	19.5 ± 0.5	/	/
<i>Colletotrichum</i> sp.	/	/	20.94 ± 3.3~8.91 ± 2.4
<i>Colletotrichum lagenarium</i>	0.0	/	/
<i>Fusarium oxysporum</i>	/	34.6	31.12 ± 3.6~12.9 ± 4.5
<i>Fusarium solani</i>	43.2 ± 0.00	16.6	/
<i>Geotrichum candidum</i> ,	45.3 ± 0.5	57.0	/
<i>Trichoderma viride</i>	0.0	/	/
<i>Rhizoctonia solani</i>	53.0 ± 1.0	100	/
<i>Sphaeroopsis sapinea</i>	/	/	23.85 ± 1.7~9.53 ± 0.6
<i>Sclerotinia sclerotiorum</i>	70.7 ± 1.1	/	/
<i>Valsa sordida</i>	/	/	9.73 ± 2.8~3.15 ± 1.00
<i>Verticillium dahliae</i>	19.4 ± 0.0	0.0	/
<i>Pythium ultimum</i>	59.1 ± 0.9	13.3	/
<i>Phytophthora cinnamomi</i>	42.0 ± 0.5	/	−19.70 ± 4.19~−5.01 ± 1.14, 25.21 ± 4.3~11.32 ± 4.2
<i>Phytophthora palmivora</i>	5.6 ± 0.5	/	/

* Data reference, ** the values listed as range for the four strains during 72 h, *** no data.

Supplementary Materials: The following are available online at <http://www.mdpi.com/2309-608X/4/2/65/s1>. Figure S1. MP phylogenetic tree of five-locus alignment for FPYF3053-3056 in genus *Diaporthe* with *Diaporthella corylina* as an outgroup; Figure S2. ML phylogenetic tree of five-locus alignment for FPYF3053-3056 in genus *Diaporthe* with *Diaporthella corylina* as an outgroup.

Author Contributions: D.-H.Y. conceived the grant and designed the experiments; D.-H.Y., and T.L. collected the plant samples and identified the plant; X.S. and H.L. performed the experiment; D.-H.Y. organized the manuscript; G.S. advised research and revised manuscript. And G.D. managed laboratory jobs for this work and processed the submission.

Acknowledgments: This work was supported by Fundamental Research Funds of CAF (CAFYBB2017MA010, to D.-H.Y.). We thank Han Xu (Research Institute of Tropical Forestry, Chinese Academy of Forestry) for kindly helping with host plant identification, and thank Kaiying Wang (Research Institute of Forest Ecology, Environment and Protection) for taking part in the work on sample collection and fungal isolation. The authors sincerely appreciate the three peer reviewers' valued comments on the paper.

Conflicts of Interest: The authors declare no conflict of interest.

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Article

Observations on the Early Establishment of Foliar Endophytic Fungi in Leaf Discs and Living Leaves of a Model Woody Angiosperm, *Populus trichocarpa* (Salicaceae)

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Received: 1 May 2018; Accepted: 13 May 2018; Published: 16 May 2018

Abstract: Fungal endophytes are diverse and widespread symbionts that occur in the living tissues of all lineages of plants without causing evidence of disease. Culture-based and culture-free studies indicate that they often are abundant in the leaves of woody angiosperms, but only a few studies have visualized endophytic fungi in leaf tissues, and the process through which most endophytes colonize leaves has not been studied thoroughly. We inoculated leaf discs and the living leaves of a model woody angiosperm, *Populus trichocarpa*, which has endophytes that represent three distantly-related genera (*Cladosporium*, *Penicillium*, and *Trichoderma*). We used scanning electron microscopy and light microscopy to evaluate the timeline and processes by which they colonize leaf tissue. Under laboratory conditions with high humidity, conidia germinated on leaf discs to yield hyphae that grew epiphytically and incidentally entered stomata, but did not grow in a directed fashion toward stomatal openings. No cuticular penetration was observed. The endophytes readily colonized the interiors of leaf discs that were detached from living leaves, and could be visualized within discs with light microscopy. Although they were difficult to visualize within the interior of living leaves following in vivo inoculations, standard methods for isolating foliar endophytes confirmed their presence.

Keywords: *Cladosporium*; colonization; microscopy; *Penicillium*; *Populus*; stomata; *Trichoderma*

1. Introduction

Plants in natural and agricultural ecosystems worldwide harbor fungal endophytes in their symptomless tissues, including apparently healthy leaves [1,2]. These endophytes often influence aspects of leaf physiology, gene expression, and function [3–5]. Their prevalence and importance prompt inquiries into the processes by which endophytic symbioses establish in living foliage. Despite their occurrence in all of the groups of land plants that have been examined to date [6], little is known regarding the early phases of endophyte colonization in leaves.

Most foliar endophytes are horizontally transmitted and form highly localized infections in leaf tissue (i.e., class 3 endophytes in [6]; hereafter, endophytes). These endophytes have been found in all of the plant species that have been surveyed thus far, and are highly diverse in wild and agroecosystems [6–11]. They typically colonize newly flushed leaves as spores or hyphal fragments [2].

After landing on leaf surfaces, most endophytes are thought to enter leaf tissues through stomata or wounds, or in some cases may enter directly through the cuticle [6,12]. Stone [13] reported that certain endophytes in the family Rhytismataceae, which occur in the leaves of Douglas fir, are limited mainly to a single epidermal cell until leaf senescence (ca. two to five years). At that time, formerly quiescent hyphae began to colonize surrounding tissue. Other studies have reported that endophytes occupy intercellular spaces near parenchyma cells [14], or grow slowly between epidermal and hypodermal cells instead of remaining quiescent [15]. These studies, while illuminating, have focused on the foliage of conifers, leaving aside the most diverse and prevalent clade of plants that host horizontally-transmitted endophytic fungi: the angiosperms.

Arnold and Herre [16] and Arnold et al. [17] demonstrated that the endophyte-free leaves of a woody angiosperm can be produced by planting surface-sterilized seeds in sterilized soil and protecting emergent leaves from surface wetting. Plants then can be inoculated by endophytes, providing a basis to study early interactions between endophytes and leaves, the processes of colonization, and—depending on the inoculum—competition, antibiosis, or other traits of the mixed endophyte communities in plant tissues [18].

The aim of this study was to examine how foliar endophytes that were isolated originally from a woody angiosperm establish endophytically within leaf tissues. We focused on inoculation experiments with *Populus trichocarpa* (Salicaceae), which is a model woody plant that can be cultivated readily from cuttings under greenhouse conditions. We examined three endophyte strains that represent three classes of Pezizomycotina (Dothideomycetes, Eurotiomycetes, and Sordariomycetes).

2. Materials and Methods

We selected three fast-growing fungal strains that were originally isolated from the leaves of *P. trichocarpa*. *Cladosporium tenuissimum* SK2012 and *Trichoderma* sp. NM2012 are endophytes that have been isolated from wild *P. trichocarpa* in Washington State [19]. They are of particular interest because they appear to reduce the severity of plant disease in *P. trichocarpa* [19]. *Penicillium citrinum* 0079 was isolated by Naupaka B. Zimmerman from *P. trichocarpa* leaves collected near Flathead Lake, Montana by Posy E. Busby. The three fungal strains readily produce conidia on V8 media within approximately seven days when incubated at room temperature (Figure 1).

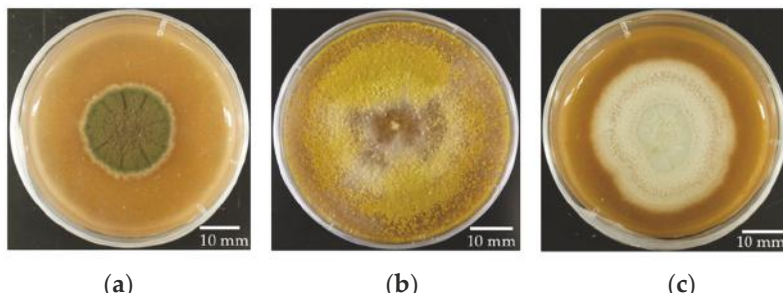


Figure 1. Selected strains of endophytic fungi. (a) *Cladosporium tenuissimum* SK2012; (b) *Trichoderma* sp. NM2012; (c) *Penicillium citrinum* 0079.

Cuttings of *P. trichocarpa* Nisqually-1 were provided by Steven Strauss (Department of Forest Ecosystems and Society, Oregon State University) from a common garden on the campus of Oregon State University. They were rooted as leafless cuttings in two-gallon pots of Sunshine Mix #4 without mycorrhizae, and grown in a greenhouse at the University of Arizona in Tucson, Arizona. Aside from experimental inoculations as described below, care was taken to avoid the surface-wetting of aboveground plant tissues throughout the experiment to avoid contamination by ambient endophytic

fungi [16,17]. Plants were surface-irrigated with a solution of Peter's Excel 15-5-15 Cal-Mag special fertilizer (Everris NA Inc., Dublin, CA, USA). The temperature in the greenhouse ranged from 7 °C to 35 °C, and humidity ranged from 20% to 90%. All of the plants had produced at least 10 leaves before the start of the experiment.

2.1. Leaf Disc Assays

We first conducted an in vitro inoculation experiment to characterize the timeline of endophyte colonization and establishment in leaf tissue. We used leaf discs that were detached from living leaves. Leaf discs were evaluated using light microscopy (LM) and scanning electron microscopy (SEM), as described below (for details regarding microscopy, see Sections 2.3 and 2.4).

Three leaves from leaf plastochron index (LPI) five to seven were collected from each of six plants on the same day. Each leaf was placed in a Ziploc bag immediately after collection and stored at 4 °C. Within 48 h of collection, leaf discs were cut from healthy leaves with sterile cork borers of the desired size, with large discs (1.2 cm in diameter) and small discs (6 mm in diameter) cut from each leaf for LM and SEM, respectively. Discs were washed by vortexing for one minute in sterile 0.1% Tween 20, and rinsed in double autoclaved water. Five 1% water agar plates in 150-mm Petri dishes were prepared for each leaf. Three plates received nine large discs each, and one plate received 24 small discs. One plate received six large discs as a control. Leaf discs were placed so that the abaxial epidermis was facing up. Agar plates with leaf discs were sealed with Parafilm and stored at 4 °C for up to two days before inoculation. Leaf discs were green and turgid at inoculation time.

After comparing fungal growth and conidia production on several types of common fungal media, we found that potato dextrose agar amended with 1% yeast extract (PDA + 1% YE) induced the most abundant conidia production across our three focal strains. Inoculum was prepared by culturing each fungal strain on PDA + 1% YE for 16 days at approximately 21 °C. Five mL of sterile 0.1% (*v/v*) Tween 20 was added to each plate, and conidia were gently dislodged with a sterile rubber policeman. The concentration of conidia was quantified with a hemacytometer and adjusted to 10⁶ conidia/mL [17]. A total of 30 µL and 5 µL of conidia solution was inoculated onto each large and small leaf disc, respectively. Plates were sealed by Parafilm and incubated in room temperature (ca. 21 °C) with approximately 12-h light/dark cycles.

We expected the initial stages of spore germination to be mainly on the leaf surface, which we evaluated by SEM within three days. Later entry was evaluated by LM for up to 21 days. After three days, samples were no longer of high enough quality for SEM, as water loss led to tissue shrinkage, which interfered with sample visualization.

For SEM, three small discs were collected at 0 h, 6 h, 12 h, 24 h, 48 h, and 72 h after inoculation and placed in 1% glutaraldehyde in 0.1-M phosphate buffer. For LM, three large discs were collected at 0 days, 1 day, 2 days, 3 days, 5 days, 7 days, 10 days, 14 days, and 21 days after inoculation, immersed in 1 M of sterile potassium hydroxide (KOH), and stored at room temperature. Samples for time point 0 for both LM and SEM were collected immediately following inoculation. We had two types of negative controls: three discs were inoculated with the standard amount of sterile 0.1% Tween 20 (but no conidia), and three discs were prepared without any inoculation solution. Both types of controls were collected at 72 h and 14 days after inoculation for SEM and LM, respectively (see below).

2.2. In Planta Assays

Since leaf discs are detached from living leaves, leaf disc assays may provide only partial insight into how endophytes interact with living tissue. Therefore, we introduced endophytes into the leaves of living plants under greenhouse conditions.

Sixteen plants were selected randomly for in vivo inoculation. The inoculum was prepared as described above, except that conidial suspensions were transferred to sterile hand-held spray bottles. Prior to use, bottles were soaked in dilute commercial bleach (0.5% sodium hypochlorite) for 10 min, and then rinsed multiple times with autoclaved water. The viability of conidia in each inoculation

solution was evaluated by spraying three times into a 50-mL sterile Falcon tube (mimicking foliar application), and then pouring the solution onto a plate with V8 agar. Fungal growth on the plate was confirmed by light microscopy three days later.

For each plant, seven leaves that ranged from LPI 5 to 11 were inoculated concurrently. The LPI of each leaf at the time of inoculation was recorded as the original LPI for later reference. Each leaf was covered with a plastic produce bag and inoculated by spraying the specified conidial suspension onto both leaf surfaces. Each bag was then sealed around the petiole to maintain high local humidity for 36 h. One leaf per plant was collected at 0 days, 1 day, 2 days, 3 days, 4 days, 7 days, and 14 days after inoculation. Leaves collected on day 0 were collected immediately after inoculation, and leaves collected on day 0 and 1 were collected before the plastic bags were removed. Leaves were collected in descending order of the original LPI, from LPI 11 to 5 (i.e., a leaf that was originally designated as LPI 11 was collected on day 0, a leaf that was originally designated as LPI 10 was collected on day 1, and so on). Each leaf was placed in a Ziploc bag and carried in a cooler to the lab for processing.

Leaves were processed immediately after collection. At each time point, 10 discs (6 mm in diameter) were cut from each leaf for SEM. An additional 10 pieces of leaf tissue (ca. 0.5 cm²) were cut from each leaf for sectioning by microtome. Leaf tissues for sectioning were placed in FAA (formalin:acetic acid:ethanol = 5:5:90), vacuumed to remove air bubbles, and stored at 4 °C.

For samples collected 0 and 14 days after inoculation, half of each leaf piece was processed for SEM and microtome sectioning immediately after collection. The other half was put in a Ziploc bag and stored at 4 °C overnight for culturing.

We used a culturing approach to determine whether inoculation was successful per traditional methods of endophyte isolation from leaf tissue. Thirty discs (6 mm in diameter) were cut from leaves collected on day 0 and day 14. Discs were washed by vortexing first in sterile 0.1% v/v Tween 20 for 30 s, and then in autoclaved water for 30 s to remove the fungi that was present on the leaf surface. The washed discs then were placed on V8 agar amended with CaCO₃ [20]. Fungi that emerged from the leaf discs were examined and identified based on morphology. For those fungi that could not be identified conclusively by morphology, genomic DNA was extracted, and the nuclear ribosomal internal transcribed spacers (ITS) and 5.8S gene, and an adjacent portion of the nuclear ribosomal large subunit (partial LSU) were amplified as a single fragment by PCR and sequenced following U'Ren et al. [21].

2.3. Sample Preparation for Light Microscopy

We used 1 M of KOH to clear leaf discs and remove cellular materials so that the endophytes could be visualized with LM on and within leaf tissues without interference by leaf pigments (modified from [13]). The KOH was changed several times until the solution and leaf discs became clear. Usually, the leaf discs were clear enough for microscopy in one month, but the time varied depending on leaf toughness and coloration.

Once the leaf discs were clear, they were transferred gently to a 35-mm Petri dish and rinsed three times with distilled water for 10 min each; then, they were stained by 0.05% trypan blue in lactoglycerol (lactic acid:glycerol:water = 1:1:1) for 2 h (modified from [22]). Discs then were destained in lactoglycerol for at least 30 min before examination by LM.

2.4. Sample Preparation for SEM

We used SEM to examine spore germination, the growth on leaf surfaces, and entry points into leaves by endophytes. Leaf discs stored in 1% glutaraldehyde were transferred to half-strength Karnovsky's fixative (Electron Microscopy Sciences, Hatfield, PA, USA) for 4 h and rinsed with 0.1 M of phosphate buffer three times for 20 min each, followed by an ethanol series dehydration (in sequence: 30%, 50%, 70%, 80%, 90%, 95%, and 100%; modified from [23]). Each dehydration step was 25 min. Discs then were immersed in 100% ethanol overnight before critical point drying using a Polaron Critical Point Drier (Polaron, Hertfordshire, UK). Leaf discs were mounted

on SEM stubs using carbon conductive tabs and then coated with platinum using a Hummer 6 Sputtering coater (Anatech, CA, USA, Hayward, CA, USA). Sample stubs were examined by Hitachi S-4800 Field-Emission (Hitachi, Tarrytown, NY, USA) SEM at the University of Arizona's University Spectroscopy and Imaging Facilities (USIF). We quickly examined each leaf disc under low magnification and then took several photos at random with a magnification of $300\times$ once patches of conidia or fungal growth were found. At this magnification, the area represented in each photo was approximately $422\text{ }\mu\text{m} \times 276\text{ }\mu\text{m}$. The number of conidia per photo was counted for reference.

2.5. Microtome Sectioning

We used microtome sectioning to obtain microsections of leaves from the *in vivo* inoculation experiment. From observations with SEM, we found that leaf discs inoculated with *Cladosporium* and *Penicillium* had fungal growth at 14 days after inoculation; as a result, we focused on these two fungi for the sectioning work. In particular, we selected samples from leaves with the highest reisolation frequency in the culturing test.

Prior to sectioning, we used Technovit 7100 (Electron Microscopy Sciences, Hatfield, PA, USA) as the embedding medium. The sample preparation protocol followed Tobe and Kadokawa [24]. Briefly, leaf tissue stored in FAA was transferred to 50% ethanol with a few drops of 1% safranin O and incubated for 4 h, followed by ethanol series dehydration (70%, 80%, 90%, 95%, and 100%). Each dehydration step was 4 h to 12 h, depending on the tissue size and thickness, and the final step was repeated to ensure complete dehydration. The plastic embedding medium was compatible with 100% ethanol, and five pre-embedding solutions in different concentrations of Technovit 7100 were prepared for filtration. Samples were immersed in each filtration solution for 12 h to 24 h; then, each piece of tissue was transferred to each block of HistoForm S for embedding and polymerization at $4\text{ }^{\circ}\text{C}$ overnight. The ratio of the embedding media Technovit 7100: hardner II: polyethylene glycol 400 (PEG400) was 15:1:0.6. Adding PEG400 can soften the embedding medium, which facilitates the cutting by disposable microtome knife [25]. Samples were suspended in blocks with Histobloc and Technovit 3040. Blocks were removed from the block mold, trimmed, and re-embedded for proper orientation before cutting by microtome. Microsections at a thickness of six μm were cut by an American Optical Spencer No. 815 (American Optical, Buffalo, NY, USA) with Tissue-Tek high-profile disposable microtome blades (Sakura Finetek USA Inc., Torrance, CA, USA). Five blocks were cut to represent each fungal strain. Forty microsections were aligned in order on one glass slide, and five slides were made for each embedded block. Slides were stained by cotton blue and safranin O [26].

3. Results

3.1. Leaf Disc Assays

Leaf discs inoculated in the *in vitro* experiment were robust to clearing by KOH prior to staining and LM, and the methods described above were also appropriate for fixing and preparing samples for SEM. Overall, we found that conidia were unevenly distributed near the central area of each leaf disc when examined by SEM. In general, conidia were visible at each time point. Most of the stomata were closed in the SEM samples, but most were open in the LM samples. The time and frequency of conidial germination varied among fungal strains, as described below.

Germinated conidia of *Cladosporium* were observed by SEM at 6 h after inoculation (HAI) (Figure 2). A small number of germinated conidia were observed at 12 HAI (Figure 2), and most conidia had germinated by 24 HAI (mean, 91%; Figure 2). Germinated conidia also were observed by LM. Extensive growth of fungal hyphae could be seen on leaf discs at two days after inoculation (DAI; Figure 3). SEM showed that fungal hyphae grew readily on the leaf surface, with growth apparently random rather than directed toward stomatal openings. However, we did not see direct evidence of hyphae entering plant tissues with SEM, and we found that some hyphae were rejected by stomata or turned away from them (Figure 3). However, examination by LM showed that conidia were usually in or

close to the stomata (Figures 2 and 3), and some hyphae were entering and growing inside leaf discs (Figure 4).

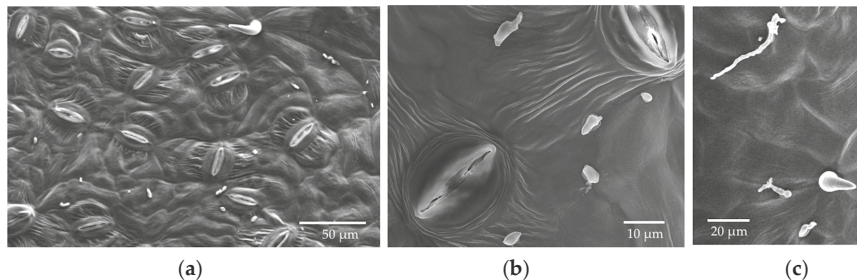


Figure 2. SEM photos from the in vitro inoculation of leaf discs of *P. trichocarpa* with an endophytic strain of *Cladosporium tenuissimum* SK2012 (a) Low magnification view of lower epidermis, six hours after inoculation (HAI); (b) Germinated and non-germinated conidia, 12 HAI; (c) Germinated conidia, 24 HAI.

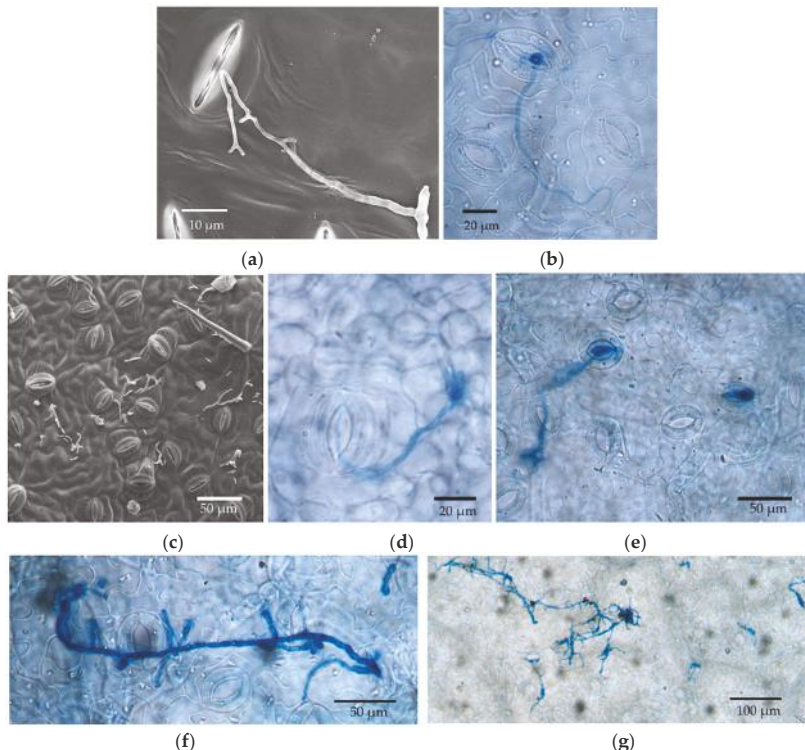


Figure 3. Photographs of hyphal growth from the in vitro inoculation of leaf discs of *P. trichocarpa* with *Cladosporium tenuissimum* SK2012. (a) Scanning electron microscopy (SEM), two days after inoculation (DAI); (b) Light microscopy (LM), 2 DAI; (c) SEM, 3 DAI; (d) LM, 3 DAI; (e) LM, 7 DAI; (f) LM, 10 DAI; (g) LM, 14 DAI.

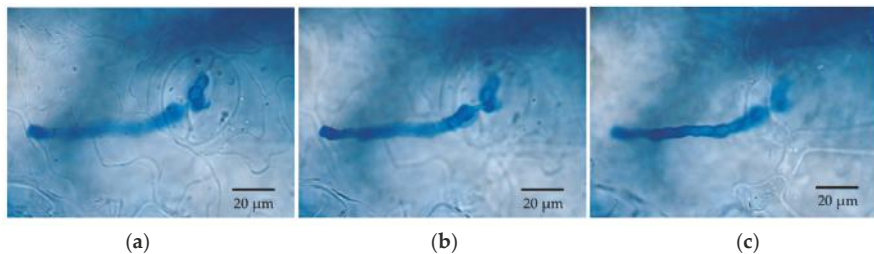


Figure 4. *Cladosporium* hypha entering a stoma of a leaf disc of *P. trichocarpa*. (a) Focused on the leaf epidermis; (b) Focused right below epidermis; (c) Focused within deeper leaf tissue.

No germinated conidia of *Penicillium* were found on leaf discs at 6, 12 and 24 HAI (Figure 5), but germinated conidia were observed by SEM at 48 HAI (Figure 5). The germination rate was less than 50% overall (Figure 5). Extensive growth of hyphae could be seen by SEM and LEM at 48 HAI or later (Figure 6). Conidia that germinated around stomata grew hyphae in random directions (Figure 6). Examination by LM showed that the margins of leaf discs were colonized extensively by fungal hyphae by 3 DAI, and some were producing conidia (Figure 6). Extensive hyphal growth across leaf surfaces was observed by LM at 3 DAI (Figure 6), but we were unable to distinguish the hyphae from inoculation/spore germination vs. growth from the disc margins.

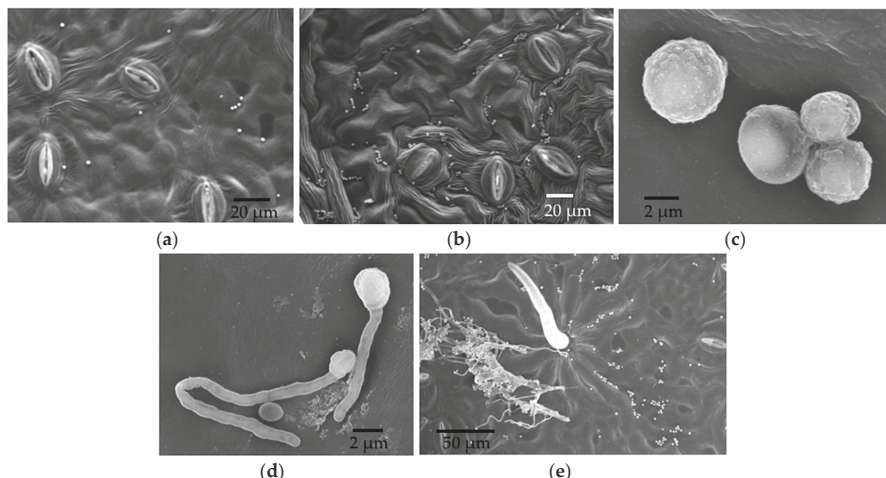


Figure 5. SEM photos from the in vitro inoculation of leaf discs of *P. trichocarpa* with an endophytic strain of *Penicillium citrinum* 0079. (a) Low magnification view of lower epidermis, six HAI; (b) Non-germinated conidia are dispersed on lower epidermis, 24 HAI; (c) Non-germinated conidia at high magnification, 24 HAI; (d) Germinated and non-germinated conidia at high magnification, 48 HAI; (e) Germinated conidia and hyphae, as well as non-germinated conidia, on lower epidermis, 48 HAI.

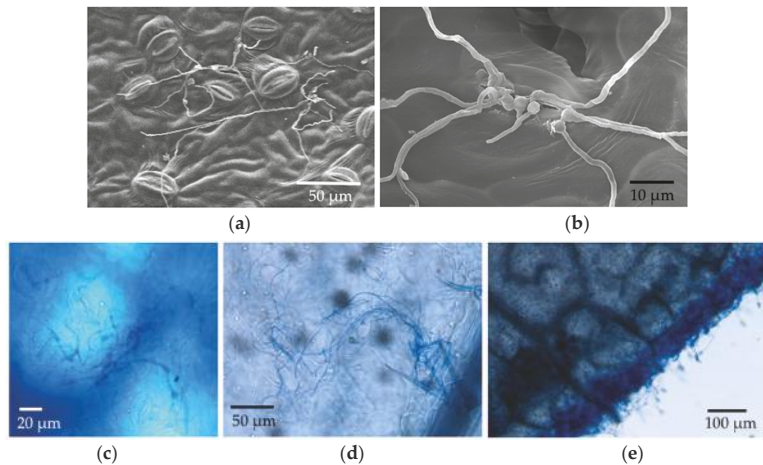


Figure 6. Hyphal growth on leaf discs of in vitro inoculation with *Penicillium citrinum* 0079. (a) SEM at low magnification, 3 DAI; (b) SEM at high magnification, 3 DAI; (c) LM, 7 DAI; (d) LM, 14 DAI; (e) LM at the margin of leaf disc, 3 DAI.

For *Trichoderma*, we examined SEM samples only at 72 HAI, as very few germinated conidia were found before this time (Figure 7). We observed hyphae from germinated conidia entering stomatal openings (Figure 7). LM at seven and 14 DAI showed a few non-germinated conidia near center of each leaf disc (Figure 8), but extensive fungal growth with conidia and chlamydospores were found at the margin of leaf discs as early as three DAI (Figure 8). We could not distinguish inoculated conidia from the conidia produced from the leaf margin. In addition, chlamydospores were produced at the leaf margin and dispersed on the leaf disc by three DAI (Figure 8). Conidia from the freshly prepared cultures were small and did not have thick walls compared with chlamydospores produced in older cultures (Figure 8).

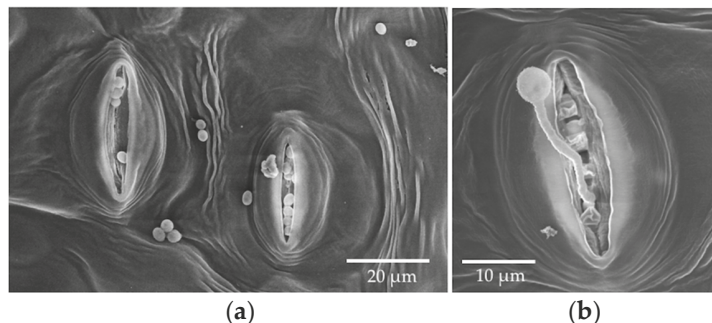


Figure 7. SEM photos from the in vitro inoculation of leaf discs of *P. trichocarpa* with an endophytic strain of *Trichoderma* sp. NM2012 at 72 HAI. (a) Non-germinated conidia; (b) A germinated conidium with hyphal penetration into the stomatal opening.

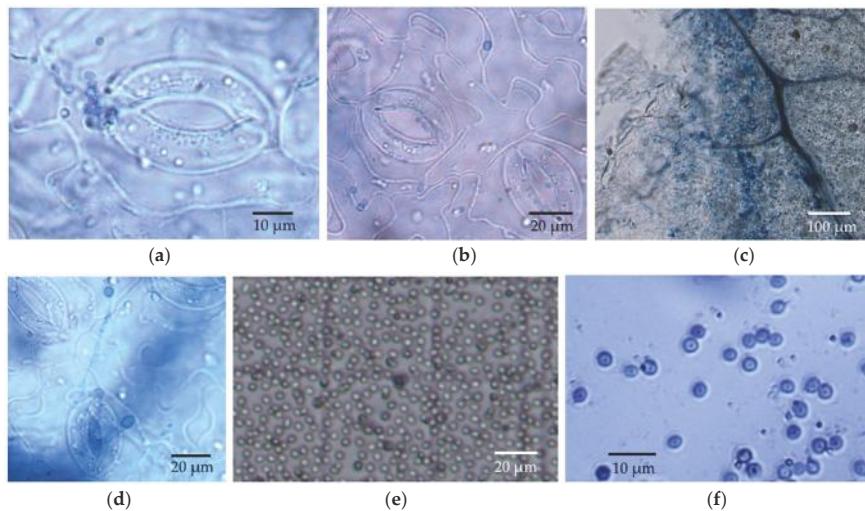


Figure 8. LM photos of in vitro inoculation with *Trichoderma* sp. NM2012. (a) Non-germinated conidia, 7 DAI; (b) Non-germinated conidia, 14 DAI; (c) Hyphal growth at the margin of leaf disc, 3 DAI; (d) Chlamydospores, 3 DAI; (e) Conidia from fresh cultures on hemocytometer; (f) Chlamydospores from a four-month culture.

3.2. In Planta Assays

Overall, 80–100% of the leaf discs that were collected immediately after leaf inoculation *in vivo* (i.e., on day 0) yielded fungal growth in culture. Non-inoculated fungi were present on 1% to 13% of leaf discs per set (Figure 9). Overall, 53% to 91% of leaf discs collected at 14 DAI yielded fungal growth in culture. At that time, non-inoculated fungi were present on 17% of discs (Figure 9). Controls consistently lacked the strains that were used in the inoculation trials, and fungi were isolated rarely from control leaves.

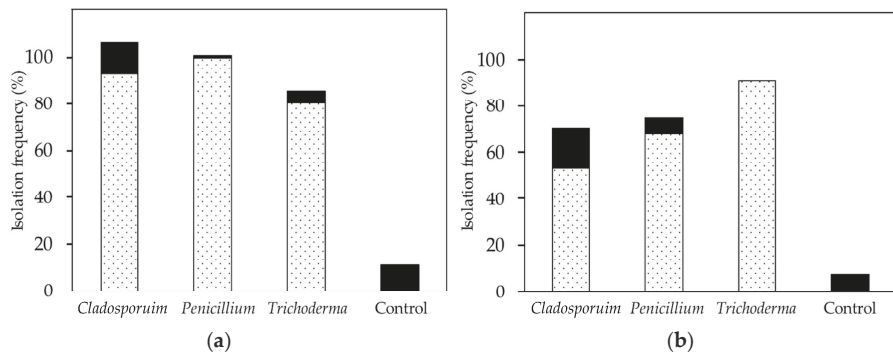


Figure 9. Results of culturing after *in vivo* inoculation. Dotted: reisolated fungi from inoculation trials, black: strains apparently present in the greenhouse. (a) Day of inoculation (i.e., day 0), isolation frequency higher than 100% because some discs yielded more than one isolate; (b) 14 DAI.

At 14 DAI, three leaf discs per leaf were examined by SEM. The greenhouse was not a sterile environment, so the leaves were not as clean as the leaf discs that were used in the in vitro inoculation. However, few bacterial cells and fungal spores were found on the control leaves, which was consistent with the culturing results presented in Figure 9. Unlike the stomata that were on the discs examined in the leaf disc assay, some stomata were open (Figure 10).

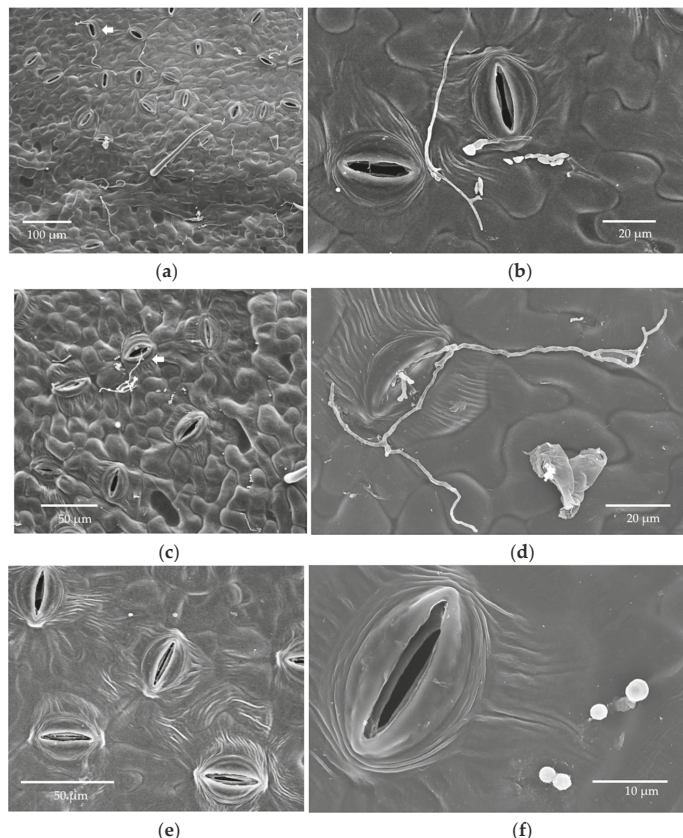


Figure 10. SEM photos of samples from in planta inoculations, 14 DAI. White arrow: hyphal entry through stomata. (a,b) *Cladosporium*; (c,d) *Penicillium*; (e,f) *Trichoderma*.

Germinated conidia and the extensive hyphal growth of *Cladosporium* and *Penicillium* were observed on the surfaces of inoculated leaves, and non-germinated conidia also were found around them (Figure 10). We did not see evidence that the hyphae of these species grew toward stomatal openings in all cases. Some grew on the leaf surface along the furrows between epidermal cells (Figure 10). Several appeared to enter the stomata (Figure 10, arrows). The conidia of *Trichoderma* were rarely found, and none had germinated (Figure 10). Microtome sectioning showed evidence of hyphae of *Cladosporium* on the upper epidermis and *Penicillium* on the lower epidermis (Figure 11), but no fungi were observed within leaf tissues.

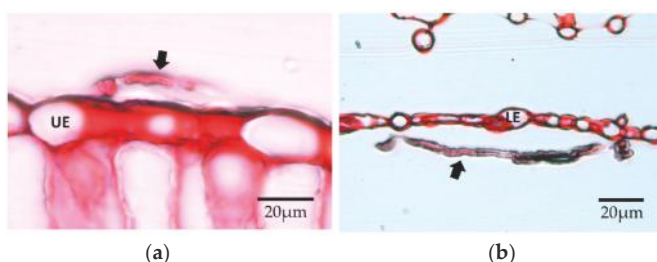


Figure 11. Sections of leaves from in planta inoculations, 14 DAI. Arrow: fungal hypha, UE: upper epidermis, LE: lower epidermis. (a) *Cladosporium*; (b) *Penicillium*.

4. Discussion

The aim of this project was to visualize the timeline and processes that were relevant to the establishment of endophytic symbioses in the leaves of a model plant. By combining leaf disc assays with in vivo inoculations of living leaves, and examining tissues with scanning electron microscopy and light microscopy, we were able to observe conidial germination, early hyphal growth, and entry into stomata by three endophytic fungi. The endophytes studied here differed in the timing and prevalence of conidial germination, grew in an undirected fashion across leaf surfaces when conditions were humid, engaged in saprotrophic growth on leaf discs, entered stomata (but not in high frequency), and were capable of growth within plant tissues. They were particularly difficult to visualize in leaves after in vivo inoculations, but could be isolated with standard methods from leaves that were inoculated in vivo.

The endophytes that were chosen for this work represent three classes of Ascomycota, and thus are phylogenetically diverse. At the same time, they appear to have converged on endophytic lifestyles, at least for part of their life cycles. These strains are different from many endophytes in that they grow rapidly and conidiate readily in culture, providing ample inoculum for these experiments.

The strains of *Cladosporium* and *Penicillium* examined here grew readily on the leaf surface and did so in an undirected manner with respect to stomatal openings. Similar growth has been reported in a previous study of pathogenic *Cladosporium* (*C. fulvum*) [27]. In that study, virulent and avirulent races of *C. fulvum* were evaluated. Neither race displayed hyphal growth toward stomata, but they differed in terms of callose deposition along the hyphae inside leaf tissues (a plant defense response [28]), and the avirulent strains did not sporulate [27]. We observed hyphae bypassing or being rejected by stomata, which is consistent with De Wit's observations [27]. However, it is possible that hyphal growth might be more directed under different conditions (e.g., under drier conditions, when stomatal openings might be targeted).

We observed via SEM that most stomata were closed in the leaf disc experiment, but frequently they were open in material collected from the in vivo experiment. Humidity, light, temperature, physiological shock, and chemical agents can alter the movement of guard cells around stomata [29,30], and the closing of stomata is a defensive response to the presence of some microbes [31,32]. The accumulation of potassium in guard cells is critical in controlling stomatal opening [33]. Open stomata in LM samples could reflect abundant potassium ions in the KOH, which was used for clearing leaf discs. These ions can flux into guard cells when the dead plant cells lose active potassium transporters in their cell membranes, thus changing the osmosis and shape of the guard cells to open the stomata. Although the humidity in the sealed agar plates should be higher than in the open air, leaf discs in the in vitro experiment did not have the nutrient and water supplies. Thus, it is possible that the stomata were closed due to this stressful environment, or in response to the inoculated fungi themselves. Whether endophyte hyphae are actively or passively rejected by the stomata remains an interesting avenue for further research.

4.1. Methods for Visualization of Fungal Endophytes in Leaves

Clearing and microsections of leaves for LM, SEM, and TEM have been used in the visualization of endophytes from tree leaves [13–15], but these have focused on gymnosperms (particularly Pinaceae), which are inhabited by fungi that include endophytic Rhytismataceae with the apparently rare habit of often occurring primarily within single epidermal cells. It is thought that the endophytes of angiosperms generally occur between cells in leaves, which is consistent with *Cladosporium* as depicted in [34]. Our visualization with LM was consistent with the intercellular growth in leaf discs by the endophytes considered here; however, we were not able to observe such growth using the microtome sections of leaves from the in vivo inoculation.

Leaf disc assays have been used extensively in plant pathology [35–38], but have been seldom applied in endophyte studies. One challenge they present is that leaf discs may encourage a saprotrophic growth phase that is not representative of endophytic growth: leaf discs have limited access not only to water and nutrients, but also to defense, when compared with living leaves. Most control (non-inoculated) leaf discs in our experiment were still green on day 14, but some became slightly discolored. Thus, leaf discs were likely under water stress, and may have had a reduced ability to respond to infection relative to leaves in planta. In general, water stress can cause plant cells to shrink, decrease leaf area, increase the synthesis of abscisic acid (ABA) and ethylene, decrease stomatal conductance, and reduce the rate of photosynthesis. Exploring the relevance of these factors for endophyte colonization is of interest for future work, as it is possible that one or more may have influenced the results of this study.

The three fungal genera used in this study are distributed widely, and are thought to have saprotrophic life phases in addition to their endophytic modes [19,39–41]. Saprotrrophic stages were observed in the in vitro inoculation, in which we found rapid growth of *Penicillium* and *Trichoderma* at the leaf disc margins. It is possible that the endophytes chosen for this work are biased somewhat toward saprotrophy and are only opportunistic as endophytes, warranting the exploration of other endophytic fungi of angiosperms with different life histories.

4.2. Visualizing Colonization in Living Leaves

Previous studies have reisolated endophytes from inoculated plants after 14 days, but report more successful isolations after 28 days or more [17]. We found that after 14 days, we frequently isolated the inoculated strains from living leaves, suggesting that *Cladosporium* and *Penicillium* did establish endophytic associations in vivo. If proliferation within tissues is relatively limited after 14 days, then perhaps a small amount of intercellular mycelium would be missed by our sectioning and visualization approach (Figure 10). Thus, a longer period between inoculation and evaluation may be useful: over time, more intercellular growth might yield more visible infections [17]. A challenge with our study system is that the individual leaves of *P. trichocarpa* were relatively short-lived in the greenhouse conditions used here, precluding long periods of incubation before harvesting. One useful way forward would be to apply these methods to mature leaves of *P. trichocarpa* in the field, wherein culturing studies suggest there exists a relatively high density of natural infections. We could then improve the methods to make them more sensitive and effective for detecting recent inoculations.

Fungal endophytes are highly diverse, even in a single leaf [6,7,42]. The three distantly related strains that were used in this study represent a small part of the vast diversity of endophytes at a global scale, and thus may not be universally representative of plant–endophyte interactions. Additionally, interactions such as competition and priority effects among fungal strains in living leaves are not captured in this report, and within-species variability among particular fungi or plant genotypes was not considered. Future studies could examine a range of isolates within focal clades of interest, seeking phylogenetically conserved modes of infection in diverse hosts. Despite these caveats, the interactions that we documented here contribute three case studies toward understanding foliar colonization by fungal endophytes in the context of woody angiosperms.

Author Contributions: Methodology, Y.-L.H. and N.B.Z.; Formal Analysis, Y.-L.H.; Writing-Original Draft Preparation, Y.-L.H.; Writing-Review & Editing, A.E.A. and N.B.Z.; Visualization, Y.-L.H.; Supervision, A.E.A.

Funding: N.B.Z. was supported by the Gordon and Betty Moore Foundation through Grant GBMF 2550.03 to the Life Sciences Research Foundation. Y.-L.H. was supported by the College of Agriculture and Life Science (CALS) and School of Plant Sciences at the University of Arizona. This study is a part of doctoral dissertation work of Y.-L.H.

Acknowledgments: We thank Posy E. Busby (Department of Botany and Plant Pathology, Oregon State University) for sharing two fungal strains. We thank Steven Strauss (Department of Forest Ecosystems and Society, Oregon State University) for providing *P. trichocarpa* cuttings. We thank Paul Wallace and the University Spectroscopy and Imaging Facility (USIF) at the University of Arizona for assistance with operation of the SEM. We appreciate Diana Wheeler (Department of Entomology, the University of Arizona) for providing the rotary microtome.

Conflicts of Interest: The authors declare no conflict of interest.

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Article

Fungi as Endophytes in *Artemisia thuscula*: Juxtaposed Elements of Diversity and Phylogeny

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Received: 25 October 2017; Accepted: 23 January 2018; Published: 27 January 2018

Abstract: *Artemisia* is a plant genus highly studied for its medicinal applications. The studies on the associated fungal endophytes are scarce. Ten plants specimens of *Artemisia thuscula* from Tenerife and La Palma were sampled to isolate the endophytic fungi. Identification of the endophytic fungi was based on morphology, Internal Transcribed Spacer (ITS) and Large Subunit (LSU) regions sequencing and indicates 37 fungal species affiliated to 25 fungal genera. Colonization rate varied among plants (CR = 25% to 92.11%). The most dominant colonizers found were *Alternaria alternata* (CF = 18.71%), *Neofusicoccum* sp. (CF = 8.39%) and *Preussia* sp. (CF = 3.23). Tendency for host specificity of most endophytic fungal species was observed. Sorenson–Dice index revealed that of 45 cases in the matrix, 27 of them were of zero similarity. Further, only one case was found to have 57% similarity (TF2 and TF7) and one case with 50% similarity (TF1 and TF4). The rest of the cases had values ranging between 11% and 40% similarity. Diversity indices like Brillouin, Margalef species richness, Simpson index of diversity and Fisher’s alpha, revealed plants from La Palma with higher values than plants from Tenerife. Three nutrient media (i.e., potato dextrose agar—PDA, lignocellulose agar—LCA, and tomato juice agar—V8) were used in a case study and revealed no differences in terms of colonization rate when data was averaged. Colonization frequency showed several species with preference for nutrient medium (63% of the species were isolated from only one nutrient medium). For the phylogenetic reconstruction using the Bayesian method, 54 endophytic fungal ITS sequences and associated GenBank sequences were analyzed. Ten orders (Diaporthales, Dothideales, Botryosphaeriales, Hypocreales, Trichosphaeriales, Amphisphaeriales, Xylariales, Capnodiales, Pleosporales and Eurotiales) were recognized. Several arrangements of genera draw the attention, like *Aureobasidium* (Dothideales) and *Aplosporella* (Botryosphaeriales) which are clustered with a recent ancestor (BS = 0.97).

Keywords: *Artemisia*; fungal endophytes; biodiversity; phylogeny

1. Introduction

Vascular plants species [1,2], aquatic plants and algae [3,4], mosses and ferns [5,6] examined to date are found to be hosts for endophytic bacteria and fungi [7]. Endophytic microorganisms have been isolated from different parts of plant-like scale primordia, meristem and resin ducts [8,9], leaf segments with midrib and roots, stem, bark, leaf blade, petiole [10], buds [11], and seeds [12]. Successful endophytic colonization is dependent on many factors including plant tissue type, plant genotype, the microbial taxon and strain type, and biotic and abiotic environmental conditions. Fungal endophytes aid plants to withstand and tolerate unfavorable environmental conditions [13,14] and also promote plant growth [15,16]. These inhabitants can produce the same or similar secondary metabolites [17–20] as their host and play vital roles in vivo such as signaling, defense, and regulation of the symbiosis [21]. Mainly investigations are based on their use as biochemical tools and the end products are to be used in pharmaceuticals, industry, and agriculture.

Artemisia is a plant highly evaluated for medicinal and biopesticide traits. A survey of the literature shows this plant genus to be in the hot spot among researchers with over 11,200 publications in Scopus library. Even though *Artemisia* is a large plant genus with species producing a variety of interesting and active compounds, its endophytic communities are under investigated. The identification of the fungal endophytes in *Artemisia* spp. is made mainly based on morphological characterization and molecular analysis using nuclear ribosomal DNA sequences, including both the internal transcribed spacers and the 5.8S gene region. To the best of our knowledge, there have been only four studies which investigate the phylogenetic analysis of the *Artemisia* spp. fungal endophytes [22–25]. In terms of diversity, the studies are also scarce but interesting facts are brought to light in terms of diversity and plant colonization. For instance, Yuan et al., 2011 [26] performed a comparative study related to infection frequency between cultivated plants and wild plants of *Artemisia annua*. The results revealed slightly higher infection frequency of the endophytic fungi in cultivated roots (20.9%) than in native roots (16.7%). Further, authors described that the naturally regenerated roots harbored richer fungal genotypes, which supports the hypothesis that wild plant species are predisposed to host rich and novel mycoflora [27]. It is worth mentioning that Qian et al., 2014 [27] reported the presence of *Rhodotorula* sp. and *Fusarium* sp. in *Artemisia argyi* for the first time. The endophytic fungi associated with *Artemisia nilagirica* were investigated and one strain of *Pythium intermedium* (Oomycota) and one strain of *Rhizopus oryzae* (Mucoromycota) were isolated among the majority clade of Ascomycota [28]. Huang et al., 2009 [24] classified 108 fungal isolates obtained from three medicinal plant species *Artemisia capillaris*, *Artemisia indica* and *Artemisia lactiflora* using morphological identification and among the three plant hosts, the highest endophytic colonization rate occurred in *Artemisia capillaris*, which exhibited highest fungal diversity. Five fungal isolates belonging to *Aureobasidium pullulans*, *Ephelis*, *Pestalotiopsis*, and *Pleosporaceae*, were only recovered from *Artemisia capillaris*. *Xylaria* species was reported to be dominant endophytic fungi in *Artemisia indica*. Seven *Artemisia* species were sampled in two locations (Qichun and Wuhan in China) and 21 fungal endophytic species belonging to: *Diaporthe*, *Colletotrichum*, *Nigrospora*, *Botryosphaeria*, *Aspergillus*, *Penicillium*, *Neofusicoccum*, *Cercospora*, *Rhizoctonia*, *Alternaria*, and *Curvularia* were found [23]. The highest incidences of colonization frequency per plant host revealed *Nigrospora sphaerica* in *Artemisia* sp., *Nigrospora oryzae* in *Artemisia argyi*, *Alternaria alternata* in *Artemisia subulata* and *Artemisia tangutica* and *Botryosphaeria dothidea* in *Artemisia lavandulifolia*. The authors report for the first time *Nigrospora*, *Neofusicoccum* and *Curvularia* species in *Artemisia* spp.

Artemisia thuscula is an endemic plant of Canary Islands and community of endophytes housed inside its plant tissues remains unexplored. With the idea of exploring endemic medicinal plants for useful and underexplored fungal endophytes, we strategically pinned down to *Artemisia thuscula* that has been harboring in western areas of islands i.e., Tenerife and La Palma, for ages. Elements of phylogeny and diversity were framed for the strains obtained from both islands with a case study of Tenerife where diversity was intended to be enhanced by using different nutrient media and stem ages. Questions on host specificity were explored, having one plant species and various collection locations.

2. Materials and Methods

2.1. Plants Sampling

Plants of *Artemisia thuscula* species were collected from Canary Islands (La Palma and Tenerife). 10 plants specimens were sampled in total. Three plants were sampled from La Palma and seven plants were sampled from Tenerife; GPS coordinates are mentioned in Table 1. In situ, plants were observed for their healthy appearance prior to the sampling, only those individuals that did not show symptoms of attack by pest or disease were selected. From each plant only stems segments were cut, labeled and kept in paper bags inside zip-locked bags at $T = 4\text{--}5\text{ }^{\circ}\text{C}$ until transported to the laboratory and then processed within 24 h. Identification of the plant species was performed using classical morphological examination. The plants were deposited at the University of La Laguna (ULL) herbarium (TFC).

Table 1. Details of collected *Artemisia* species plants.

Plant Species	Plant Code	Collection Place	Country/Island	Herbarium Type Details	GPS UTM Latitude	GPS UTM Longitude
<i>A. thuscula</i>	LP1	El Granel	La Palma	TFC. No. 52658	28°45'47.43" N	17°45'7.47" W
<i>A. thuscula</i>	LP2	San Bartolo	La Palma	TFC. No. 52659	28°46'1.08" N	17°45'26.07" W
<i>A. thuscula</i>	LP4	Tigalate	La Palma	TFC. No. 52661	28°32'35.45" N	17°48'41.29" W
<i>A. thuscula</i>	TF8	El Palmar	Tenerife	TFC. No. 52669	28°20'35.18" N	16°51'26.57" W
<i>A. thuscula</i>	TF7	Granadilla	Tenerife	TFC. No. 52668	28°06'54.19" N	16°34'51.14" W
<i>A. thuscula</i>	TF4	Caletillas	Tenerife	TFC. No. 52665	28°23'2.03" N	16°21'54.71" W
<i>A. thuscula</i>	TF1	Mesa Mota	Tenerife	TFC. No. 52662	28°30'38.75" N	16°19'20.55" W
<i>A. thuscula</i>	TF2	Mesa Mota	Tenerife	TFC. No. 52663	28°30'38.75" N	16°19'20.55" W
<i>A. thuscula</i>	TF5	San Andres	Tenerife	TFC. No. 52666	28°30'51.01" N	16°11'41.94" W
<i>A. thuscula</i>	TF3	Taborno	Tenerife	TFC. No. 52664	28°33'18.36" N	16°15'53.10" W

2.2. Fungal Endophyte Isolation

Surface sterilization method was used to suppress epiphytic microorganisms from the plant [23]. Thus, stem fragments were first washed with sterile water, then immersed in 70% ethanol for 1 min, followed by an immersion in 15% sodium hypochlorite for 1 min, again in 70% ethanol for 1 min and lastly were washed with sterile distilled water. To assure a successful sterilization, fragments were rolled on potato dextrose agar (PDA) medium and drops of last step sterilization water were poured on medium, as a control check for complete sterilization. After this process, plant material was dried on sterile blotting sheet, excised in pieces of 2 cm and cut longitudinally with a sterile scalpel. Segments were placed in PDA (Sigma-Aldrich, St. Louis, MI, USA) Petri plates amended with tetracycline (10 mg L⁻¹). Plates incubated with the plant segments were incubated at 25 °C in the dark for two weeks and observed daily for fungal growth. When fungal outgrowth from the plant tissues occurred observations on emerged fungi were made. Only the fungi with different morphological characteristics were subcultured. Eventually, when an endophyte was acquired in pure culture it was preserved in Czapek medium (Fluka Analytical, Sigma-Aldrich), T = 5 °C and in glycerol (\geq 99.5, Sigma-Aldrich) 20% in deionized H₂O, T = -32 °C and identified. To analyze the fungal diversity, each replicate of the distinct stem fragments was noted. To enhance bioprospection and diversity, variable nutritive media were utilized to incubate stem fragments (with ages less than one year and more than one year) of eight plants from Tenerife. Therefore, V8 tomato juice agar and lignocellulose agar (LCA) [29] media were additionally used. All the reagents were purchased from Sigma-Aldrich, except Agar Agar—GUINAMA (Valencia, Spain) and Potassium chloride—PanReac AppliChem (Barcelona, Spain).

2.3. Fungal Endophyte Collection and Maintenance

Every isolate and its plant origin were duly recorded for calculation of colonization rate from host, counting the same isolate identification only once if it emerges from the same plant segment. After purification of each isolate, it was subjected to microscopical observations followed by molecular analysis to identify at genus and/or species level. Isolates are presently maintained in three types of media: Czapek, T = 5 °C; mineral oil (Sigma-Aldrich), T = 5 °C and glycerol (Sigma-Aldrich) 20% diH₂O, T = -80 °C. For short term use, fungal isolates were maintained on PDA, 25 °C.

2.4. Morphological Identification

Prior to taxonomic identification, a preliminary classification was made to avoid the selection of identical strains arising from the same plant individual, separating isolates into morphotypes. Observations targeted characteristics related to the colony and medium as: colony shape, texture and colour; exudates, medium colour and growth rate. For the microscopic observations, a strain was inoculated onto a PDA Petri plate and a sterile cover slide was attached at two centimeters. Once the growth of the fungus partially covered the cover slide, the slide was removed, inverted on a slide with cotton blue (for the slightly coloured colonies) and observed under microscope.

2.5. Molecular Identification

Out of several procedures for genomic DNA extraction, the most efficient protocol, although time consuming, was the one described by Shu et al., 2014 [20] to which the following modifications were made. Samples were centrifuged for 15 min at 12,000 rpm; after the chloroform ($\geq 99.5\%$, Sigma-Aldrich) procedure the upper phase was mixed with 10% Sodium acetate (ReagentPlus[®], $\geq 99.0\%$, Sigma-Aldrich) and 60% Isopropyl alcohol (Aldrich $\geq 97.0\%$, Sigma-Aldrich), incubated for 10 min at -30°C and centrifuged (10 min, 12,000 rpm). Finally, the pellet was washed twice with 75% ethanol (before maintained at -20°C) and centrifuged (10 min, 12,000 rpm). The solvent was removed by evaporation, keeping the sample in the laminar flow cabinet. The purified DNA was suspended in 20 μL TE buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA); all reagents were purchased from Sigma-Aldrich. RNase A was added, and the sample was incubated for 1 hour at room temperature (long-term storage at -32°C).

The second protocol for DNA extraction involves no purification of DNA but acceptable results were garnered (around 50% samples succeeded). 20 μL of TE buffer was pipetted into a microtube and glass beads (diameter = 0.4–0.6 mm) were added to make up 3/4 of the reagent's volume. A small quantity of fungal mycelium was added (2–5 mm/2–3 mg) with a needle. Samples were homogenized using FastPrep 24TM 5 G (MP Bio, Santa Ana, California, USA) at 4 m/s, 20 s. Subsequently samples were centrifuged at 13,000 rpm for 1 min and maintained on ice. One μL of the supernatant was used for the PCRs.

The third and fourth protocol involved two genomic DNA extraction kits. First one used was E.Z.N.A. Fungal DNA Kit according to the manufacturer indications (OMEGA bio-tek, Norcross, Georgia, USA) with overall good results (around 80% of the samples succeeded). The second one tested was Fungi/Yeast Genomic DNA Isolation Kit, according to the manufacturer indications (NORGEN Bioteck, Thorold, ON, Canada) with overall good results also (approximately 70% of the samples succeeded).

The fourth protocol approaches nucleic acid extraction by application of silica coupled to magnetic particles, which is efficient and automated. Genomic fungal DNA was extracted using Maxwell 16 Mouse Tail DNA purification kit. The Promega kit is designed for automated DNA extraction from tissue samples using the MaxwellTM 16 platform (Promega BioSciences, San Luis Obispo, CA, USA). This protocol was performed at the University Institute of Tropical Diseases and Public Health of the Canary Islands, University of La Laguna.

Molecular identification of the fungal Dicarya strains was performed using ITS1 (5'-TCCGTAGGTGAAACCTGCGG-3') and ITS4 (5'-TCCTCCGCTTATTGATATGC-3') primer pair to amplify the 5.8S rDNA and the two internal transcribed spacers ITS1 and ITS2 [30] for the majority of the samples and NL-1 (5'-GCA TAT CAA TAA GCG GAG GAA AAG-3') and NL-4 (5'-GGT CCG TGT TTC AAG ACG G-3') primer pair to amplify the 5' end of 28S rDNA spanning domains D1 and D2) [31]. PCRs were performed in a total volume of 25 μL containing 10 ng genomic DNA, 0.5 μM primer, 200 μM dNTPs, 1X Buffer Taq, 0.0125U of Taq DNA Polymerase. For ITS sequences, PCR cycling parameters were carried out according to Shu et al. 2014 [20] with slight modifications: 94°C for 2.5 min; 40 cycles of 94°C for 30 s, 58°C for 30 s, and 72°C for 1 min; and a final extension at 72°C for 10 min. For 28S rDNA domain, the PCR conditions were denaturation for 4 min at 95°C followed by 45 s at 95°C and then annealing for 45 s at 58°C , 1 min at 72°C , followed by an extension at 72°C for 5 min. The final step was at 16°C for 5 min. A total of 40 cycles were performed. All PCR products were detected by agarose gel electrophoresis (110V, 35 min, on 2% agarose gels, 1X TAE Buffer) loading 5 μL PCR product, 1 μL Loading Buffer (6X) and 2 μL SYBR Green I (Sigma-Aldrich; dilution 1:10,000). PCR and electrophoresis reagents were purchased from Sigma-Aldrich. PCR products were purified using GenEluteTM PCR Clean-Up Kit (Sigma-Aldrich) and sequenced by Sequencing Services SEGAI (La Laguna, Spain). The sequences were run through the BLASTN search page using Megablast program (National Center for Biotechnology Information; Bethesda MD, USA) where the

most identical hits and their accession numbers were obtained. Further, only ITS sequences were used for the phylogenetic analysis, therefore details on 28S sequenced strains are listed in Table 2.

Table 2. *Artemisia* fungal endophytic strains: codes, identities as per morphology, 28S rDNA LSU sequences and their most similar hits from Genbank with accession numbers and values.

EF Code	Assigned Species/Species Complex	GenBank Identified Seq.	Max Score	Total Score	E Value	Max Identity (%)	Accession No.
HLP1	<i>A. alternata</i>	<i>A. alternata</i>	1107	1107	0	99	KX609781.1
HLP10	<i>A. alternata</i>	<i>A. alternata</i>	1093	1093	0	99	KX609781.1
HLP31	Fungus sp. 1	Uncultured fungus clone	135	724	2.00×10^{-27}	100	KP843503.1
HLP5	<i>A. alternata</i>	<i>A. alternata</i>	1052	1052	0	99	KF751621.1
HLP6	<i>Curternaria lunata</i>	<i>Cochliobolus lunatus</i>	1026	1026	0	99	KC616350.1
HLP8	<i>Neofusicoccum</i> sp. 1	<i>N. cryptoaustrale</i>	511	511	5.00×10^{-141}	92	KX464415.1
HLP9	<i>Preussia</i> sp. 1	<i>P. mimooides</i>	1000	1000	0	97	KF557659.1
HTF25	<i>Alternaria</i> sp. 6	<i>A. brassicicola</i>	289	289	5.00×10^{-74}	77	AF397222.1
HTF37	<i>A. alternata</i>	<i>A. alternata</i>	982	982	0	96	KX609781.1
HTF42	<i>Neofusicoccum australe</i>	<i>N. australe</i>	1036	1036	0	97	KF766367.1
HTF49	<i>Neofusicoccum australe</i>	<i>N. australe</i>	1058	1058	0	98	HM176550.1
HTF50	<i>A. alternata</i>	<i>A. alternata</i>	971	971	0	98	KF543048.1
HTF67	<i>Chaetomium</i> sp. 1	<i>C. coarctatum</i>	846	846	0	99	KX976729.1
HTF75	<i>Neofusicoccum australe</i>	<i>N. australe</i>	934	934	0	98	HM176550.1
HTF78	<i>A. alternata</i>	<i>A. alternata</i>	1051	1098	0	99	FJ839651.1
HTF80	<i>Camarosporium</i> sp. 1	<i>Camarosporium</i> sp.	1024	1024	0	97	KF733369.1

2.6. Phylogenetic Analysis

ITS sequences [i.e., endophytic fungi—Table 3, their most similar hits from GenBank (NCBI, Bethesda MD, USA) and type sequences of the selected taxa] were aligned with the multiple alignment program ClustalW [32] as implemented in Mega 6.0 (Molecular Evolutionary Genetics Analysis) [33] and indels corrected manually to minimize alignment gaps [34]. Designated outgroup was *Caloscypha fulgens* (GenBank Accession No. DQ491483). After the exclusion of non-overlapping leading/trailing gaps the length of the alignment was 603 bps. Because of the high number of indels, these were recoded as a binary matrix by means of the simple indel coding algorithm [35], appending the fragments to the nucleotide data as additional characters, as implemented in FastGap 1.21 (Department of Biosciences, Aarhus University, Denmark) [36]. This “indel matrix” was used in all Bayesian and maximum likelihood analyses. Formerly, Gblocks program (hosted at www.phylogeny.fr) was used to eliminate poorly aligned positions and divergent regions [37]. Best-fit models were compared in jModelTest 2 according to Bayesian Information Criterion (BIC) [38]. Best fit according to the BIC criterion model (K80 + G) was selected to reconstruct the Bayesian tree. Bayesian Inference analysis was conducted with MrBayes 3.2.3 (hosted by Mobyle SNAP Workbench, North Carolina State University) [39] and run for 1×10^7 generations with a sampling frequency of 100 generations. Of the resulting trees, the first 25,000 trees were discarded as burn-in and the following 75,001 were used to estimate topology and tree parameters. The percentage number of times a node occurred within these 75,001 was interpreted as the posterior probability of the node [40]. Convergence of the runs was indicated by an average standard deviation of split frequencies between duplicate runs of less than 0.01. The consensus trees were drawn using Treepgraph 2 software (Institute for Evolution and Biodiversity, University of Munster, Germany) [41] and edited with Adobe Illustrator CS3 (Adobe Systems Incorporated, San Jose, CA, USA).

Table 3. Endophytic fungi isolated from *A. thuscula* and used for the phylogenetic analysis: codes, identity and accession numbers of the ITS sequences.

Strain Code	Identity	Accession No.
HLP12	<i>Phoma</i> sp.	MG025848
HLP14	<i>Preussia</i> sp.	MG025849
HLP15	<i>Diaporthe phaseolorum</i>	MG025850
HLP19	<i>Alternaria alternata</i>	MG025851
HLP23	<i>Diaporthe novem</i>	MG025852
HLP24	<i>Nigrospora oryzae</i>	MG025853
HLP25	<i>Camarosporium brabeji</i>	MG025854
HLP27	<i>Coniothyrium</i> sp.	MG025855
HLP3	<i>Aspergillus flavus</i>	MG025856
HLP32	<i>Alternaria alternata</i>	MG025857
HLP37	<i>Diaporthe</i> sp.	MG025858
HLP40	<i>Alternaria alternata</i>	MG025859
HLP43	<i>Trematea</i> sp.	MG025860
HLP44	<i>Neoplatysporoides aloicola</i>	MG025861
HLP45	<i>Neofusicoccum parvum</i>	MG025862
HLP46	<i>Neofusicoccum parvum</i>	MG025863
HLP7	<i>Paraphoma chrysanthemicola</i>	MG025864
HTF23	<i>Nectria maurititicola</i>	MG025865
HTF26	<i>Stachybotrys longispora</i>	MG025866
HTF27	<i>Stemphylium solani</i>	MG025867
HTF30	<i>Aplospora prunicola</i>	MG025868
HTF31	<i>Stemphylium solani</i>	MG025869
HTF40	<i>Biscogniauxia mediterranea</i>	MG025870
HTF41	<i>Alternaria alternata</i>	MG025871
HTF44	<i>Alternaria alternata</i>	MG025872
HTF46	<i>Alternaria alternata</i>	MG025873
HTF48	<i>Phoma</i> sp.	MG025874
HTF52	<i>Alternaria alternata</i>	MG025875
HTF53	<i>Alternaria alternata</i>	MG025876
HTF62	<i>Neofusicoccum parvum</i>	MG025877
HTF64	<i>Pestalotiopsis</i> sp.	MG025878
HTF66	<i>Aureobasidium pullulans</i>	MG025879
HTF68	<i>Stemphylium solani</i>	MG025880
HTF70	<i>Cladosporium</i> sp.	MG025881
HTF74	<i>Preussia australis</i>	MG025882
HTF76	<i>Alternaria alternata</i>	MG025883
HTF79	<i>Phoma</i> sp.	MG025884
HTF81	<i>Preussia</i> sp.	MG025885
HTF82	<i>Biscogniauxia mediterranea</i>	MG025886
HTF83	<i>Stemphylium solani</i>	MG025887
HTF84	<i>Preussia</i> sp.	MG025888
HTF85	<i>Preussia</i> sp.	MG025889

2.7. Diversity Analysis

The colonization rate (CR%) was calculated as the total number of stem fragments in a sample (plant/nutritive medium) yielding at least one isolate divided by the total number of stem fragments in that sample. Colonization frequency (CF%) was calculated as the total number of fragments in a sample (plant/location) colonized by a species divided by the total number of fragments plated. For the diversity of endophytic fungi, the Margalef index, Brillouin index, Fisher's alpha index and Simpson's dominance index were used. Margalef index [42] measures species richness while Brillouin index combines richness and evenness. The Margalef index was calculated using formula $d = (S - 1) / \ln N$, where S is the number of species and N is the number of individuals in the sample. The Brillouin index [43,44] was calculated using formula: $HB = (\ln N! - S \ln n_i!) / N$, where N is the total number

of individuals, S is the number of taxa and n_i is the number of individuals belonging to i species. Fisher's logarithmic series model [45] is a species-abundant model and describes the relationship between the number of species and the number of individuals of those species. It was calculated using formula $S = a \times \ln(1 + n/a)$, where S is number of taxa, n is the number of individuals and a is the Fisher's alpha. The dominance of Simpson [46] was calculated according to the formula $D = 1 - \sum [n_i(n_i-1)/N(N-1)]$, where n_i is the number of individuals belonging to i species and N is the total number of individuals. The Sorenson–Dice coefficient of similarity [47,48] which expresses the beta diversity was employed to compare the similarity of endophytic fungi communities regarding species composition between two host plants, nutrient media and stem ages. The Sorenson–Dice coefficient is calculated with the formula $QS = 2C/(A + B)$ where A and B are the species numbers in samples A and B, respectively, and C is the number of species shared by the two samples. The Sorenson–Dice coefficient weighs more the joint occurrences than the mismatches and is expressed with values between 0 (no similarity) and 1 (absolute similarity). This index was used to assess host preference and spatial heterogeneity by describing the similarity of endophytic communities within ten host plants at distinct sampling sites. Nevertheless, as the coefficient analyses the presence/absence data, no judgments on abundance or rare taxa can be pursued. A binary matrix was produced and used to calculate the similarity matrix and to plot a dendrogram based on an unweighted paired group method of arithmetic average (UPGMA) cluster analysis. For the diversity indices, PAST software version 3.15 (copyright Hammer & Harper, Natural History Museum, University of Oslo, Norway) was used.

3. Results and Discussion

3.1. Fungal Endophytic Diversity in *Artemisia* Species

3.1.1. Colonization Rate and Colonization Frequency of Endophytic Fungi in *Artemisia thuscula*

In this study, the employed analyses indicate that 37 fungal species and 25 fungal genera were isolated from 10 plants of *Artemisia thuscula*. Colonization rate (further CR) shows how much a plant can be colonized within predetermined conditions. It is valuable information as different plants showed distinct values of this index; therefore, low values could express plants poor in endophytic fungi culturable in the given conditions.

To calculate the colonization frequency (CF) of fungal endophytes in *Artemisia* species plants, we have considered same fungal endophytic species isolated from two or more plant fragments as being a distinct isolate belonging to the same species. Therefore, if the same species was isolated twice from the same plant fragment, it was considered only one time. This issue is to be expected at isolation moment, when no precise differentiation between the isolates can be defined, and only once purified and further analyzed then only the strain received a final identification. The CF% gives a hint over the distribution and abundance of a certain fungal species in a sample (i.e., plant/location/region). To know the “area” of the distribution and abundance of a certain endophytic fungal species, we have analyzed the data per plant individual or plant location, plant species, and plant region. Regions were grouped here as: La Palma Island and Tenerife Island. This way we can have an overview on where certain fungal species are more abundant or rare, as well as if there is a relation between their distribution and plant-specific parameters.

In *Artemisia thuscula*, only one plant out of 10 had a colonization rate value over 90% (LP2). The lowest values (CR% = 25) were recorded for three plants (TF8, TF7 and TF3). Interestingly, as per variable geographical location there is a considerable variation between La Palma Island and Tenerife Island, with the former having the most colonized plant individuals (Table 4).

Table 4. Colonization rate (CR) of fungal endophytes in *Artemisia thuscula* collected in Canary Islands.

Collection Place	Region	Plant Number	Locality Code	CR%
El Granel	La Palma	*LP1	EG	48.28
El Palmar—Teno	Tenerife	**TF8	EP	25.00
Granadilla	Tenerife	TF7	GR	25.00
Igualte Caletillas	Tenerife	TF4	IC	50.00
Mesa Mota	Tenerife	TF1	MM	62.50
Mesa Mota	Tenerife	TF2	MM	62.50
San Andrés	Tenerife	TF5	SA	62.50
San Bartolo	La Palma	LP2	SB	92.11
Taborno	Tenerife	TF3	TA	25.00
Tigalate	La Palma	LP4	TIG	55.00
AVG				50.78
SD				16.13

*LP = La Palma; **TF = Tenerife; AVG = average; SD = standard deviation.

Artemisia thuscula cannot escape of the “omnipresence” of *Alternaria alternata*, this species was isolated from eight plants but with relevant differences in the frequency, CF% = 15–50%. A notable presence is remarked here, *Neofusicoccum australe*, isolated from three plants at relatively high values (CF% = 25) when considering that the maximum value is 50. Moreover, the *Neofusicoccum* genus, consisting here of three species was isolated from eight plants, one of which revealed a CF% of 34.21. Interestingly, around 70% of the fungal species in *Artemisia thuscula* were isolated from only one plant each (Table 5). This suggests a host specificity which was also exhibited by the low and moderate values of Sorenson–Dice coefficient when the similarity of the endophytic assemblages was analyzed (see further Diversity indices for endophytic fungi in *Artemisia thuscula*).

Table 5. Colonization frequency on potato dextrose agar (PDA) medium of fungal endophytic species in *Artemisia thuscula* plants.

Plant Code	Locality Code	EF Species	CF%	Plant Code	Locality Code	EF Species	CF%
LP1	EG	<i>Alternaria alternata</i>	24.14	LP2	SB	<i>Preussia</i> sp. 3	2.63
LP1	EG	<i>Alternaria</i> sp. 5	13.79	LP2	SB	<i>Trematelia</i> sp. 1	2.63
LP1	EG	<i>Aspergillus flavus</i>	3.45	LP4	TIG	<i>Alternaria alternata</i>	15.00
LP1	EG	<i>Aspergillus flavus</i>	6.90	LP4	TIG	<i>Curvularia lunata</i>	5.00
LP1	EG	<i>Diaporthe novem</i>	3.45	LP4	TIG	<i>Neofusicoccum</i> sp. 1	5.00
LP1	EG	Fungus sp. 1	3.45	LP4	TIG	<i>Paraphoma</i> cf. <i>chrysanthemicola</i>	5.00
LP1	EG	<i>Neofusicoccum parvum</i>	3.45	LP4	TIG	<i>Preussia</i> sp. 1	25.00
LP1	EG	<i>Nigrospora oryzae</i>	3.45	TF1	MM	<i>Alternaria alternata</i>	25.00
LP1	EG	<i>Penicillium viridicatum</i>	3.45	TF1	MM	<i>Thielavia</i> sp. 1	8.33
LP1	EG	<i>Phoma</i> sp. 3	3.45	TF2	MM	<i>Alternaria alternata</i>	50.00
LP1	EG	<i>Pleosporales</i> sp. 2	6.90	TF2	MM	<i>Biscogniauxia mediterranea</i>	12.50
LP1	EG	<i>Preussia</i> sp. 3	3.45	TF2	MM	<i>Neofusicoccum australe</i>	12.50
LP2	SB	<i>Alternaria alternata</i>	10.53	TF2	MM	<i>Phoma</i> sp. 1	12.50
LP2	SB	<i>Alternaria</i> sp. 5	2.63	TF3	TA	<i>Neofusicoccum australe</i>	25.00
LP2	SB	<i>Camarosporium bradgi</i>	2.63	TF4	IC	<i>Alternaria alternata</i>	37.50
LP2	SB	<i>Coniothyrium</i> sp. 1	2.63	TF4	IC	<i>Aureobasidium pullulans</i>	12.50
LP2	SB	<i>Diaporthe phaseolorum</i>	7.89	TF5	SA	<i>Alternaria alternata</i>	12.50
LP2	SB	<i>Diaporthe</i> sp. 1	5.26	TF5	SA	<i>Alternaria</i> sp. 6	12.50
LP2	SB	<i>Dothideomycetes</i> sp. 1	2.63	TF5	SA	<i>Macrophomina phaseolina</i>	25.00
LP2	SB	Fungus sp. 1	2.63	TF5	SA	<i>Stachybotrys longispora</i>	12.50
LP2	SB	<i>Neofusicoccum parvum</i>	13.16	TF7	GR	<i>Alternaria alternata</i>	25.00
LP2	SB	<i>Neofusicoccum</i> sp. 3	34.21	TF7	GR	<i>Neofusicoccum australe</i>	25.00
LP2	SB	<i>Neoplatyphoroides aloicola</i>	5.26	TF7	GR	<i>Stemphylium solani</i>	25.00
LP2	SB	<i>Nigrospora</i> sp. 2	2.63	TF8	EP	<i>Camarosporium</i> sp. 1	12.50
LP2	SB	<i>Pleosporales</i> sp. 3	2.63	TF8	EP	<i>Phoma</i> sp. 1	12.50

EG = El Granel; SB = San Bartolo; TIG = Tigalate; MM = Mesa Mota; TA = Taborno; IC = Igualte Caletillas; SA = San Andres; GR = Granadilla; EP = El Palmar.

34 endophytic fungal species were isolated from *Artemisia thuscula* (Table 6) and their frequency varied within a low range with two exceptions: *Alternaria alternata* (CF% = 18.71) and *Neofusicoccum* sp. 3 (CF% = 8.39).

Table 6. Colonization frequency of fungal endophytic species in *Artemisia thuscula* (overall CF%/plant species).

EF Species	CF%	EF Species	CF%
<i>Alternaria alternata</i>	18.71	<i>Neofusicoccum parvum</i>	3.87
<i>Alternaria</i> sp. 5	3.23	<i>Neofusicoccum</i> sp. 1	0.65
<i>Alternaria</i> sp. 6	0.65	<i>Neofusicoccum</i> sp. 3	8.39
<i>Aspergillus flavus</i>	1.94	<i>Neoplatyphoroides aloicola</i>	1.29
<i>Aureobasidium pullulans</i>	0.65	<i>Nigrospora oryzae</i>	0.65
<i>Biscogniauxia mediterranea</i>	0.65	<i>Nigrospora</i> sp. 2	0.65
<i>Camarosporium bradgi</i>	0.65	<i>Paraphoma chrysanthemica</i>	0.65
<i>Camarosporium</i> sp. 1	0.65	<i>Penicillium viride</i>	0.65
<i>Coniothyrium</i> sp. 1	0.65	<i>Phoma</i> sp. 1	1.29
<i>Curularia lunata</i>	0.65	<i>Phoma</i> sp. 3	0.65
<i>Diaporthe novem</i>	0.65	<i>Pleosporales</i> sp. 2	1.29
<i>Diaporthe phaseolorum</i>	1.94	<i>Pleosporales</i> sp. 3	0.65
<i>Diaporthe</i> sp. 1	1.29	<i>Preussia</i> sp. 1	3.23
<i>Dothideomycetes</i> sp. 1	0.65	<i>Preussia</i> sp. 3	1.29
Fungus sp. 1	1.29	<i>Stachybotrys longispora</i>	0.65
<i>Macrophomina phaseolina</i>	1.29	<i>Stemphylium solani</i>	0.65
<i>Neofusicoccum australe</i>	2.58	<i>Tremateia</i> sp. 1	1.94

Studies that are independent of fungal isolation and identification methods often revealed higher numbers of fungal species [49]. We purposely chose the culture method to further select endophytic fungi of high interest according to their biological activities. Our goal was to yield a large number of endophytes, and not to produce a complete species list of fungal endophytes in these *Artemisia* species. Nevertheless, the data obtained gave us an interesting fragment of knowledge about the communities of these microorganisms in their plant hosts.

In terms of endophytic fungal species CF%, the most isolated species was *Alternaria alternata* (CF = 18.71; eight of ten plants), as expected. It is a common saprobe found on various plants and other substrata worldwide [50,51] and has often been isolated as endophyte in previous studies [52–55]. Qian et al., 2014 [27] isolated endophytic fungi from *Artemisia argy* and found Pleosporales to be the most represented group, with three species of *Alternaria* present. It was found as the most predominant species in grasses [56] and various plants families, also [57]. Among dominant endophytic fungal species, we observed taxa like *Neofusicoccum* and *Preussia*. These genera of endophytic fungi were previously isolated from a wide range of host plants including *Artemisia* spp. [24,57–60].

Interestingly, it was observed a tendency on host specificity of most endophytic fungal species. In *Cirsium arvense* similarity in endophytic communities decreased with increasing intersite distance [61] while in *Holcus lanatus* the similarity between leaf and root myco-assemblages at the same location was lower than that observed in leaves at different locations [58]. Further, in leaf fungal communities the average number of species shared by any pair of location was 3.13 and in root assemblages was 1.73 out of an average of 12.2 species identified at each location [58].

Despite the dominant species, the rest of the endophytic fungal species reflect an unequal distribution of a certain endophytic species among plant individuals. This same issue was previously observed [58] but no definitive answer has been found. Some hypotheses were proposed like ubiquitous taxa with spatial dominance or selection of certain dependent on culture conditions [58]. In the case study on *Artemisia thuscula* (see Section 3.1.3) taxa such as *Preussia*, *Pestalotiopsis*, *Aplosporella*, *Chaetomium* and *Cladosporium* were isolated from only one nutrient medium out of the three media

tested. Nevertheless, this is not a unique parameter, which should account for the determination of an endophytic taxa preference for a nutrient medium. One of the major variables which we consider is the rest of the community involved and their role in the interaction when the isolation performed. That is, which are the other taxa living in the same “space” (i.e., plated plant fragment) and we must consider if there are (i) fast-growing taxa versus slow growing taxa; (ii) nutrient deficiency or promoting medium for certain taxa, as well as (iii) the interaction between the taxa (i.e. antagonism).

3.1.2. Diversity Indices for Endophytic Fungi in *Artemisia thuscula*

In the La Palma results of diversity, Margalef index revealed the highest value for species richness in San Bartolo (Margalef = 4.24) followed by El Granel (Margalef = 3.69). The Brillouin index agrees that the highest diversity is found in San Bartolo (Brillouin = 1.8) but Fisher’s alpha index shows a higher abundance of rare species in El Granel (Fisher’s alpha = 18.6) than in San Bartolo (Fisher’s alpha = 13.9). Diversity regarded as evenness was found to be similar in both localities (Simpson’s index: El Granel = 0.88 and San Bartolo = 0.87). In La Palma Island, San Bartolo locality was revealed as having the highest value for species richness and diversity. Yet, El Granel was shown as having a higher abundance of rare species (Fisher’s alpha: El Granel = 18.6 and San Bartolo = 13.9) and a higher value of evenness than San Bartolo (Simpson’s index: El Granel = 0.88 and San Bartolo = 0.87). In Tenerife, the locality San Andres showed by far the highest diversity in all previously mentioned terms and all the indices confirm it (Table 7).

Table 7. Diversity indices of fungal endophytic species per plant individual/locality.

Locality Code	Taxa No.	Strains No.	Simpson 1-D	Brillouin	Margalef	Fisher Alpha
LP1/EG	11	15	0.89	1.65	3.69	18.60
LP2/SB	15	27	0.87	1.85	4.25	13.90
LP4/TIG	5	5	0.80	0.96	2.49	0.00
TF1/MM	5	10	0.60	0.85	1.74	3.98
TF3/TA	1	2	0.00	0.00	0.00	0.80
TF4/IC	2	3	0.44	0.37	0.91	2.62
TF5/SA	4	5	0.72	0.82	1.86	9.28
TF7/GR	3	3	0.67	0.60	1.82	0.00
TF8/EP	2	2	0.50	0.35	1.44	0.00

LP1/EG = El Granel; LP2/SB = San Bartolo; LP4/TIG = Tigalate; TF1/MM = Mesa Mota; TF3/TA = Taborno; TF4/IC = Igreste Caletillas; TF5/SA = San Andres; TF7/GR = Granadilla; 14/EP = El Palmar.

Sorenson–Dice index revealed that of 45 cases in the matrix, 27 of them were of zero similarity. Further, only one case was found to have 57% similarity (TF2 versus TF7) and one case with 50% similarity (TF1 versus TF4). The rest of the cases had values ranging between 11% and 40% similarity. These different similarity values may be due to distance among hosts, soil composition and/or climatic conditions. When the distance was plotted (UPGMA), the Sorenson–Dice coefficient clustered plants LP1 and LP2 with maximum bootstrap support (BPP = 100), although these plants had only 38% similarity in between. Nevertheless, this is to be considered a high value of similarity in the given matrix and one of the reasons for obtaining it might be the proximity of the collection places (approx. 5 km) between the host plants and similar altitudes and climate. Further clusters were formed like LP4 and TF1; TF2 and TF3; TF4, TF5 and TF7 (Figure 1). As we expected (from CF and CR values) TF8 is the most different host plant, the backbone of the dendrogram divides into this branch and the other branches which form various clusters of similarity. Also, cluster LP1 and LP2 is a sister cluster of the other clusters which were exhibited as more related in terms of similarity.

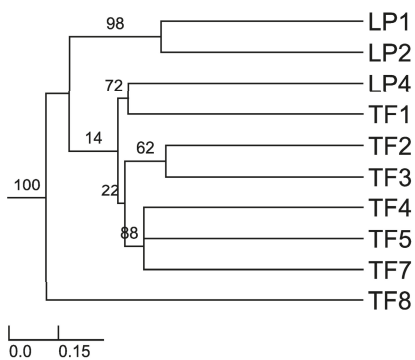


Figure 1. Sørensen's similarity coefficient for the endophytic fungi isolated from *A. thuscula*: unweighted paired group method of arithmetic average (UPGMA) dendrogram plot. The results were obtained with 95% of confidence and bootstrap values calculated from 1000 iterations.

3.1.3. Case Study: *Artemisia thuscula* of Tenerife, Endophytic Fungi Isolated from Two Types of Stems on Three Media: Colonization Frequency and Colonization Rate

In this study, we can observe throughout various individual plants from the same species (i.e., *Artemisia thuscula*) the relevance of nutrient media and the age of the stem as the selected organ to yield endophytic fungi. When averaged the colonization rates of the three nutrient media selected (PDA, LCA, and V8) do not show relevant differences (CR% = 33.93, 33.93 and 37.50, respectively). Neither do the differences of age in stems; stems with the age < 1 year have CR% = 30.95 and stems with age > 1 year have CR% = 36.90.

Differences may be observed (Table 8) when comparing different plants, as for instance TF3 and TF4 had the lowest colonization rates (CR% = 16.67) and no endophytic fungi was isolated from V8 or stems with age of more than 1 year for TF4 and TF3, respectively. In addition, there is no higher value than 58.33 of colonization rate, as observed in other individuals of *Artemisia*.

Table 8. *Artemisia thuscula* colonization rate (CR%) of endophytic fungi per plant, per medium and per stem age.

Plant Code	CR%/Plant	CR%/PDA	CR%/LCA	CR%/V8	CR%/Stem	
					Age < 1 Year	Age > 1 Year
TF3	16.67	25	12.5	12.5	33.33	0
TF4	16.67	37.5	12.5	0	8.33	25
TF5	58.33	62.5	75	37.5	58.33	41.67
TF7	45.83	25	50	62.5	50	41.67
TF8	37.5	25	37.5	50	8.33	66.67
AVG	35.12	33.93	33.93	37.50	30.95	36.90

If colonization frequency data is segregated into plants sampled (Table 9), we observe that *Alternaria alternata* is the major colonizer in three out of seven plants, namely TF2, TF4, and TF5. Plants had different yields considering number of endophytic fungal species, ranging between three (TF3) and eight (TF7).

Among the major colonizers we observed *Neofusicoccum austral* and *Neofusicoccum parvum* in TF3 (CF% = 8.33), *Chaetomium* sp. 1 and *Phoma* sp. 1 in TF7 and *Phoma* with two different species in TF8 (CF% = 16.67; Table 8). Myrchiang et al., 2014 [28] investigated the endophytic fungi associated with *Artemisia nilagirica* and comparing the colonization of three organs (i.e., root, stem and leaf), the authors obtained the highest diversity in the roots (i.e., 14 species), less in stem (i.e., 10 species) and

the smallest number in the leaves (i.e., 6 species). Similarly, in *Artemisia thuscula* Cosoveanu et al., 2012 [62] isolated 29 distinct morphotypes: 20 from roots, 7 from stem and 2 from leaves. In addition, Myrchiang et al., 2014 [28] observed that from all fungal endophytic species, only *Phoma eupyrena* was found to be a common occurrence in all plants sample, the other species having a certain preference for one or maximum two organs.

Table 9. *Artemisia thuscula* colonization frequency (CF%) of endophytic fungi species per plant, per medium and per stem age.

EF Species	Plant	CF%/Plant	CF%/Medium			CF%/Stem Age	
			PDA	LCA	V8	≤1 year old	>1 year old
<i>Alternaria alternata</i>	TF2	50	50	37.5	62.5	41.67	58.33
<i>Alternaria</i> sp. 4		4.17		12.5			8.33
<i>Biscogniauxia mediterranea</i>		4.17	12.5			8.33	
<i>Neofusicoccum australe</i>		16.67	12.5	12.5	12.5	16.67	8.33
<i>Pestalotiopsis</i> sp.		4.17		12.5		8.33	
<i>Phoma</i> sp. 1		4.17	12.5			8.33	
<i>Neofusicoccum australe</i>	TF3	8.33	25			16.67	
<i>Neofusicoccum parvum</i>		8.33		12.5	12.5	16.67	
<i>Pestalotiopsis</i> sp.		4.17		12.5		8.33	
<i>Alternaria alternata</i>	TF4	12.5	37.5			16.67	8.33
<i>Aureobasidium pullulans</i>		4.17	12.5			8.33	
<i>Nectria mauritica</i>		8.33		25		16.67	
<i>Alternaria alternata</i>	TF5	37.5	12.5	50	50	8.33	66.67
<i>Alternaria</i> sp. 6		4.17	12.5			8.33	
<i>Aplosporella prunicola</i>		4.17		12.5		8.33	
<i>Aureobasidium pullulans</i>		4.17		12.5		8.33	
<i>Macrophomina phaseolina</i>		8.33	25			16.67	
<i>Neofusicoccum australe</i>		4.17		12.5		8.33	
<i>Stachybotrys longispora</i>		4.17	12.5			8.33	
<i>Stemphylium solani</i>		8.33		12.5	12.5	16.67	
<i>Alternaria alternata</i>		8.33	8.33	12.5	12.5		16.67
<i>Aureobasidium pullulans</i>	TF7	12.5		12.5		8.33	
<i>Chaetomium</i> sp. 1		16.67			25	16.67	
<i>Cladosporium</i> sp. 1		12.5			25	16.67	
<i>Neofusicoccum australe</i>		8.33			12.5	8.33	
<i>Phoma</i> sp. 1		16.67		50		33.33	
<i>Preussia australis</i>		8.33	12.5				8.33
<i>Stemphylium solani</i>		4.17	12.5			8.33	
<i>Biscogniauxia mediterranea</i>	TF8	8.33			12.5		8.33
<i>Camarosporium</i> sp. 1		8.33	12.5			8.33	
<i>Phoma</i> sp. 1		16.67	12.5				
<i>Preussia</i> sp. 2		16.67			37.5	25	
<i>Preussia</i> sp. 3		12.5		12.5			8.33
<i>Preussia</i> sp. 5		12.5		25		16.67	
<i>Stemphylium solani</i>		4.17		12.5		8.33	

Comparing different plant individuals of the same species and observing the distribution of fungal endophytes provides insights to determine the occurrence of a certain species. For instance, in TF2 four fungal species were isolated only from one nutrient medium, namely *Biscogniauxia mediterranea* in PDA, *Alternaria* sp. on LCA, *Phoma* sp. on PDA and *Pestalotiopsis* sp. on LCA (Table 9). Furthermore, we may observe that the same species of *Phoma* sp. 1 was also isolated from TF8 on PDA, similar to *Pestalotiopsis* isolated from TF3 on LCA while *Biscogniauxia mediterranea* was isolated on V8 from TF8.

When the distribution of endophytic fungi species is observed in terms of colonization frequency per total number of the studied plants (Figure 2), data showed several species like *Aplosporella prunicola*, *Camarosporium* sp., *Chaetomium* sp., *Cladosporium* sp., *Nectria mauritica* and others with certain “preference” for nutrient medium. It is well known that fungi have specific carbon and nitrogen requirements for sporulation [63–65]. However, the requirements for fungal growth are less stringent but not less important when isolation is pursued. Nutrient - rich media result in selective isolation for fast-growing fungi, overlooking slow growing species if present [66]. Osono and Takeda [29] stated

that LCA due to its low glucose content suppresses the overgrowth of fast-growing species. 22 species of fungal endophytes were isolated from all *Artemisia thuscula* plants in this case study and 14 species (63%) were isolated only from one nutrient medium. Additionally, 12 fungal species were isolated from stems older than 1 year and seven were isolated from stems younger than one year. Seven fungal species are to be considered rare, as their colonization frequency value is the lowest one, throughout the data set (CF% = 0.60; Figure 2).

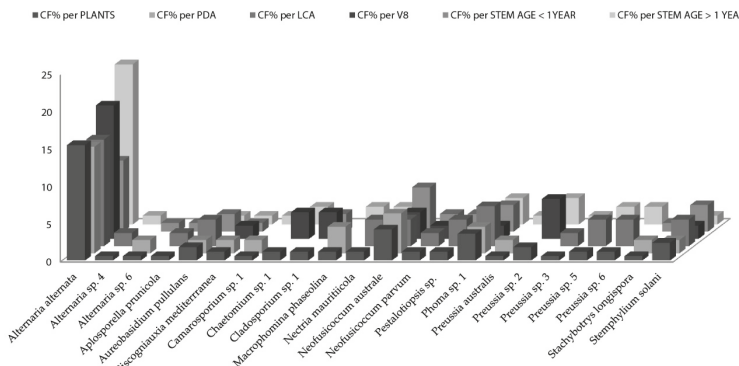


Figure 2. *Artemisia thuscula* colonization frequencies of endophytic fungi species per total number of plant individuals, per media (PDA, lignocellulose agar (LCA) and V8) and per stem age (>1 year and <1year). The y-axis data correspond to colonization frequency percentage.

Further, Sorensen–Dice similarity coefficient reveals proximate values among the similarities of the endophytic communities isolated on the three tested media (Table 10). Yet, none of them overpassed 52% similarity (i.e., LCA versus V8). As for the stem ages, the index showed a value of 43% similarity. Evidence for tissue specificity was previously demonstrated for phloem and xylem tissue, where the value of endophytic similarity reached 36% in roots of *Sophora tonkinensis* [67]. This suggests the necessity to broad both culture media and diversity of tissues to obtain a higher richness of endophytic fungal taxa.

Table 10. *A. thuscula* fungal endophytes isolated on different nutrient media and stem age: Sorensen–Dice coefficient of similarity.

	LCA	V8	≤1 Year	>1 Year
PDA	0.42	0.48	0.64	0.71
LCA		0.52	0.67	0.67
V8			0.67	0.59
≤1 year				0.43

Among the singleton species that occur only in the *Artemisia thuscula* plant individuals selected for this case study (i.e., limited to Tenerife) we have: *Aplosporella prunicola*, *Camarosporium* sp. 1, *Macrophomina phaseolina*, *Chaetomium* sp. 1, *Nectria mauritiicola*, *Neofusicoccum australe*, *Pestalotiopsis* sp., *Phoma* sp. 1 and *Stachybotrys longispora*. Except *Phoma* sp., all endophytic fungal species previously mentioned were isolated from only one nutrient medium. In addition, except *Camarosporium* sp. (isolated also from *Artemisia thuscula* in Palma Island), all endophytic fungal species previously named were isolated only from *Artemisia thuscula* in Tenerife Island.

3.2. Phylogenetic Relations

54 endophytic fungal ITS sequences and the associated GenBank sequences were used for the phylogenetic analysis (Table 3; sequences of strains HLP16, HLP22, HLP28, HLP33, HLP48A, HLP4,

HTF29, HTF33, HTF43, and HTF61 are not listed and are only available at request). The dataset consists of 603 characters after alignment, 43 characters are conserved, and 447 characters are parsimony informative, while 557 are variable characters. Bayesian Posterior Probabilities (BPPs) given below each node are shown on the upper branches.

Ten orders (Diaporthales, Dothideales, Botryosphaeriales, Hypocreales, Trichosphaeriales, Amphisphaeriales, Xylariales, Capnodiales, Pleosporales and Eurotiales) are recognized (Figure 3). The phylogenetic tree divides the taxa in five main clades, leaving *Diaporthe* sequences unclustered. Clade 1 consists of Dothideales and Botryosphaeriales (BPP = 0.98), Clade 2 groups Hypocreales, Trichosphaeriales, Amphisphaeriales and Xylariales (BPP = 0.88), Clade 3 and Clade 4 contain Capnodiales (BPP = 0.79) and Pleosporales (BPP = 0.63), respectively while Clade 5 accommodates Eurotiales (BPP = 0.62).

Interestingly, *Diaporthe* sequences are not clustered but several show different branch lengths. Yet, taxa *D. novem* and *D. phaseolorum* do not differentiate. Endophytic fungi were basically identified using morphology; therefore, HLP15 and HLP23 were considered *D. phaseolorum* and *D. novem*, respectively while structures of HLP37 did not allow an accurate species level identification. Apparently, the ITS region in *Diaporthe* is evolving at higher rates than TEF1 or MAT genes [68], therefore presenting a wider variation than advisable for species boundaries. Thus, a slowly evolving gene region should be used in order to establish species limits [69]. Nevertheless, ITS sequence data can be used for reliable identification of phylogenetic relationships as long as they are interpreted with care [69]. Several arrangements of genera draw the attention, like *Aureobasidium* (Dothideales) and *Aplopsorella* (Botryosphaeriales) which are shown with an immediate common ancestor (BPP = 0.97). *Aplopsorella* has over 300 species and appears to be heterogenous; therefore not all species are likely to belong in Botryosphaeriaceae [70]. The ascomycete genus *Aureobasidium* is a member of the family Aureobasidiaceae within the class of the Dothideomycetes [71]. Dothideomycetidae subclass was emended by Schoch et al., 2006 [72] and a new subclass was proposed, Pleosporomycetidae, with an additional order, the Botryosphaerales.

Penicillium and *Aspergillus* sequences form two sister clades as expected (BPP = 0.60). Three species of *Neofusicoccum* are clustered with relevant support (BPP = 0.89) while *N. parvum* is drawn outside. Hypocreales taxa are split in two sister clusters along with *Stachybotrys*, *Grandibotrys*, *Melanopsamma* and *Sirastachys* in one sister clade although with no relevant support (BPP = 0.55) and *Nectria*, *Sarocladium* and *Corallomyctela* as another sister clade (BPP = 0.87). Also, internal clustering is revealed between several taxa of the mentioned genera. Trichosphaeriales and Amphisphaeriales are shown having a common recent ancestor (BPP = 0.98). Hypocreales is recognized as monophyletic [73]. The order Hypocreales incorporates Nectriaceae and Stachybotriaceae beyond other six families [74]. Maharachchikumbura et al., 2014 [75] found using a combined LSU, SSU, TEF and RPB2 sequences data that *Stachybotrys* and related taxa (Stachybotriaceae) form a sister cluster of *Nectria* and related taxa (Nectriaceae). The results obtained with the ITS region are in accordance with the combined inference obtained by Maharachchikumbura et al., [75]. The Nectriaceae group (BPP = 0.87) comprises *Nectria* (Nectriaceae), *Sarocladium* (Hypocreomycetidae) and *Corallomyctella*—shown to comprise two distinct clades in Nectriaceae [76]. The second cluster joints *Stachybotrys*, *Grandibotrys*, *Sirastachys*, *Stachybotrys* (Stachybotriaceae, Hypocreomycetidae) and *Melanopsamma* (Chaetosphaeriaceae, Sordariomycetidae). *Melanopsamma pomiformis* was recently excluded from the genus [77] and it was linked to the asexual morph *Stachybotrys albipes* [78]. Strains of Sordariomycetes clustered into six subclasses among which Diaporthomycetidae, Xylariomycetidae and Hypocreomycetidae [75]. Our Bayesian analysis resulted in a monophyletic clade (Clade 2) which accommodates Hypocreales (Hypocreomycetidae), Trichosphaeriales (Diaporthomycetidae), Amphisphaeriales (Xylariomycetidae) and Xylariales (Xylariomycetidae). Yet, Diaporthales taxa (*Diaporthe* spp.) were left outside this clade. A resulting parsimonious tree of multi-locus based (LSU, ITS, and TEF1) sequences shows that the genus *Diaporthe* has paraphyletic origins [79]. Xylariales and Amphisphaeriales were found as sister clusters in Xylariomycetidae sharing a common ancestor [80].

Yet, the clade which accommodates Xylariomycetidae is a sister clade of Diaporthomycetidae (Diaporthales) and Hypocreomycetidae (Hypocreales).

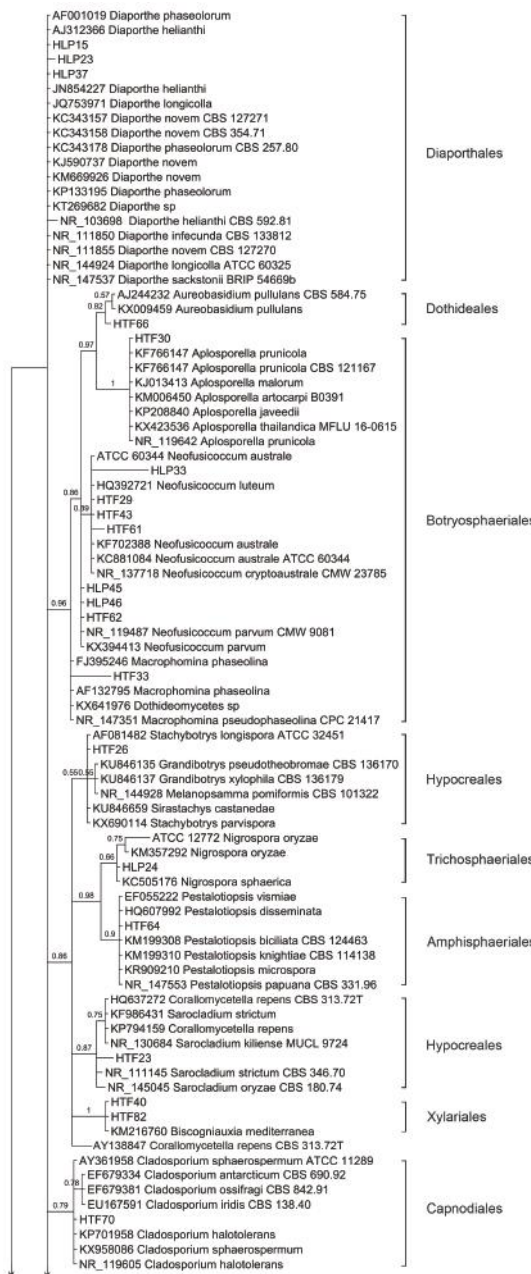


Figure 3. Cont.



Figure 3. Bayesian phylogenetic tree based on ITS rDNA sequence variants of the endophytic fungi isolated from *A. thyscula* and their associated external GenBank hits. The tree was rooted with *Caloscypha fulgens* sequence as outgroup. The Bayesian clade-credibility values (posterior probabilities) are indicated at internodes (BPP). The scale bar represents the expected changes per site. Sequences coded with HLP/HTF were obtained from endophytic fungi, sequences coded with taxa names are associated external sequences and the ones coded with CBS/L/ATCC/NRRL/B/MFLU/CMW/MUCL/BRIP were obtained from type strains.

Cladosporium sequences are clustered (BPP = 0.79) and different branch lengths between species are revealed, grouping *C. ossifragi*, *C. antarcticum* and *C. iridis* (BPP = 0.78). Conversely, *Aplosporella* sequences do not differentiate in between, showing all species with same branch lengths.

Stemphylium sequences are grouped but support does not avail this grouping (BPP = 0.53). *Phoma*-like sequences are clustered as expected (BPP = 0.93) showing higher differences between *Phoma*, *Didymella*, *Dothiorella* and *Notophoma* on one side (BPP = 0.93) and *Paraphoma chrysanthemicola* on the other side (BPP = 0.88). It is curious that several sequences of endophytes are grouped in a sister clade of *Alternaria* clade, *Phoma*-like clade and *Stemphylium* clade with high probability (BPP = 1), indicating different branch lengths. *Alternaria* sequences are not grouped in a single cluster but different branch lengths are drawn among the species. Similarly, *Preussia* and *Sporormiella* taxa are spread. *Coniothyrium*-like sequences are clustered, but support has an average value, BPP = 0.78. *Coniothyriaceae* and *Camarosporiaceae* grouping as well as the *Coniothyrium*-like sequences cluster and its sister cluster of *Pleosporaceae* is supported by the findings of Wijayawardene et al., 2014 [81]. Mainly the sequences obtained from the endophytic strains are grouped with the external sequences as expected (i.e., morphological identification) but several are left unclustered. For instance, inside the group of *Pleosporales* three endophytic sequences (HLP16, HLP22 and HLP48A) appear as more related, forming a strong-supported cluster (BPP = 1). This apparently new lineage should be confirmed with another phylogenetic study based on large subunit and small subunit nuclear rDNA regions, where only *Pleosporales* taxa would be included. In the present study none of the methods used like the morphology (absence of the sporulating structures), BLAST alignment (values of similarity with GenBank provided sequences did not exceed 86%, 88% and 84% for HLP16, HLP22, and HLP48A, respectively) and the ITS inference, could provide their proper identification or genetic stronger alliances inside *Pleosporales*.

4. Conclusions

The present study suggests culturable endophytic species have specificity for a plant host and “preference” for nutrient medium. Therefore, this study indicates the apparent necessity of using different culture media so as to obtain a higher diversity of species.

Acknowledgments: We thank Kanika Chowdhary (Indian Institute of Technology) and Mariano Hernandez (Dept. Bioquímica, Microbiología, Biología Celular y Genética, Universidad de La Laguna) for their valuable contributions. Two anonymous reviewers are thanked for their comments and suggestions on the draft version of this script. The authors are grateful to Cristian Predescu and Alin Gayraud for the graphical elements - phylogenetic tree and abstract, respectively. Andreea Cosoveanu was supported by the pre-doc contract as research personnel in formation (sp. “Contratos predoctorales para la formación de doctores”), Call 2015 by Universidad de La Laguna and La Caixa.

Author Contributions: Andreea Cosoveanu and Raimundo Cabrera conceived and designed the experiments; Andreea Cosoveanu and Samuel Rodriguez Sabina performed the experiments; Andreea Cosoveanu and Raimundo Cabrera analyzed the data; Raimundo Cabrera contributed reagents/materials/analysis tools; Andreea Cosoveanu and Raimundo Cabrera wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest. The funding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

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Review

Mangrove-Associated Fungi: A Novel Source of Potential Anticancer Compounds

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Received: 16 July 2018; Accepted: 21 August 2018; Published: 24 August 2018

Abstract: Cancer is the second leading cause of death worldwide, and the number of cases is increasing alarmingly every year. Current research focuses on the development of novel chemotherapeutic drugs derived from natural as well as synthetic sources. The abundance and diversity in natural resources offer tremendous potential for the discovery of novel molecules with unique mechanisms for cancer therapy. Mangrove-derived fungi are rich source of novel metabolites, comprising novel structure classes with diverse biological activities. Across the globe, coastal areas are primarily dominated by mangrove forests, which offer an intensely complex environment and species that mostly remain unexplored. In recent years, many structurally diverse compounds with unique skeletons have been identified from mangrove fungi and evaluated for their antiproliferative properties. These compounds may serve as lead molecules for the development of new anticancer drugs. Mangrove endophytes can be modulated using epigenetic means or culture optimization methods to improve the yield or to produce various similar analogs. The present review provides an insight into the bioactive metabolites from mangrove endophytes reported during the period from 2012 to 2018 (up to April, 2018) along with their cytotoxic properties, focusing on their chemical structures and mode of action, as indicated in the literature.

Keywords: endophytic fungi; anticancer compounds; mangroves; co-culture; epigenetic modification

1. Introduction

Mangroves are salt-tolerant forest ecosystems, representing a lively ecosystem with an amalgam of land-dwelling and marine habitats with high biodiversity and socio-economic importance [1]. Marine fungi are among the most prominent species existing in mangrove forests, and support nutrient replenishment [2]. As per reports on marine fungi, in this ecological niche mangrove fungi make up the second largest group [3]. These fungi may occur as saprophytes, symbiotically or as a parasites in the mangrove ecosystem. In addition, these fungi belong to both the lower class, such as oomycetes and thraustochytrids, and the upper class, such as ascomycetes and basidiomycetes. Fungal secondary metabolites are structurally quite diverse, and their functions mostly depend on self-defense against other microorganisms [4]. Most often mangrove fungi flourish in challenging habitats, making them a rich source of bioactive metabolites. Endophytes are one of the various groups of mangrove fungi that have resulted in the identification of a large number of new bioactive metabolites of nutraceutical and pharmaceutical importance. These include antibiotic, anticancer, antidiabetic, antioxidant, antiviral, anti-inflammatory and immunosuppressive drugs, along with other pharmaceutical agents [5].

Cancer affects different organs, and is identified by the unchecked proliferation of abnormal cells that invade other healthy tissue. The treatment is primarily confined to chemotherapy. Besides being expensive, chemotherapy is known to cause severe side effects, making treatment problematic. The non-effectiveness of many existing drugs along with multi-drug resistance further aggravates the problem, making cancer treatment difficult. For medicinal chemists, the primary goal remains the discovery and identification of chemotherapeutic agents derived from natural products. Secondary metabolites have opened new avenues for the development of novel therapeutic agents [6,7]. Endophytic fungi, which are a less-explored area of the microbial community, have a tremendous potential to produce new metabolites that can be used for pharmaceutical applications. Since the initial report of the identification of paclitaxel, derived from an endophyte associated with Northwest Pacific yew by Stierle et al. [8], scientists have identified many other crucial anticancer molecules from fungal endophytes [6]. Many researchers were attracted to marine mangrove fungi because of their diversity, which may lead to the discovery of several novel natural products. With the remarkable advancements in spectroscopic techniques, separation methods and microplate-based sensitive *in vitro* assays, the natural product exploration of mangrove fungi has attracted special attention regarding novel and unexplored chemical scaffolds [9]. Of the various existing groups of mangrove fungi, endophytes have been identified as producers of new bioactive metabolites with pharmaceutical and nutraceutical importance.

Most of the endophytes have the potential to produce novel bioactive metabolites, which will undoubtedly boost novel drug discovery. However, higher similarity among microbes leads to the frequent identification of the same compound in the endophytes. During axenic cultivation, a specific portion of the biosynthetic genes are expressed while growing *in vitro*, and various genes stay masked or silent and do not express in laboratory conditions. For this reason, the routine method of fermentation yields metabolites without chemical diversity. Co-cultivation could help to overcome this problem and is preferred, with two or more microbes allowed to grow together. This approach offers a better competitive environment, allowing the increased production of constitutive as well as cryptic compounds that are not traced out in axenic cultures [10]. Several co-cultivation strategies such as different combinations of fungi, the co-cultivation of fungi with bacteria and the co-cultivation of different bacteria have been reported for the enhancement of the chemical diversity of marine-derived microorganisms [10].

The development of methodologies to induce the expression of biosynthetic transcription as well as the suppression of these genes plays a vital role in the search for new secondary metabolites. The regulation of the enzymes that control metabolite production can be achieved by changing epigenetic mechanisms such as DNA methylation and histone modifications (acetylation and phosphorylation) by using epigenetic modifiers [11]. As an example of the importance of epigenetic modulation in producing unknown natural products, when *Aspergillus niger* is cultivated over a two-week period in vermiculite-based semi-solid medium treated with suberoylanilide hydroxamic acid (SAHA), it leads to the isolation of a new fungal metabolite nygerone [12]. Hence the epigenetic approach can be a game changer in the production/enhancement of secondary metabolites.

The present review provides a comprehensive overview of the bioactive metabolites identified from mangrove endophytes during the period from 2012 to 2018 (up to April, 2018) including eighty novel compounds of the total 181 reported. The total number of compounds as well as novel compounds isolated from mangrove fungi during this period is presented in Figure 1. The origin, chemical structure of the biological targets and efficacies of these compounds are also discussed where available. The anticancer properties of many of these compounds are presented in Table 1. They are arranged based on the broader category of the taxonomic class of the cytotoxic compounds producing fungi. An attempt has also been made to review recent developments such as co-cultivation and epigenetic modifications in endophytic fungi to enhance the secondary metabolite production.

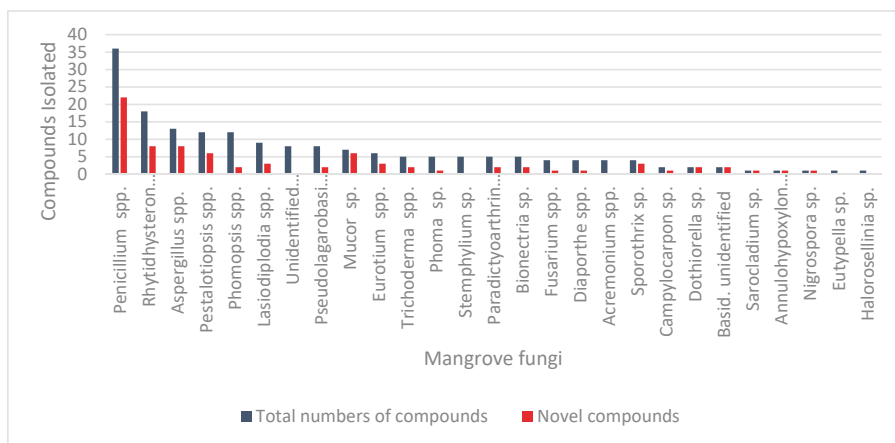


Figure 1. Novel anticancer bioactive compounds reported from mangrove fungi.

2. Bioactive Compounds in Mangrove Plants

2.1. Compounds Produced by Coelomycetes

Pestalotiopsis is the most noteworthy coelomycetous fungi and the species of *Pestalotiopsis* are known to produce the diverse array of novel compounds. Strobel and Long [13] described *Pestalotiopsis* as the “*E. coli* of the temperate and tropical rainforest systems”. The species *Pestalotiopsis* is widely recognized to be a prolific producer of a diverse array of metabolites that include alkaloids, chromones, coumarins, isocoumarin derivatives, lactones, peptides, phenols, phenolic acids, quinones, semiquinones, xanthones, terpenoids and xanthone derivatives along with an array of antimicrobial, antifungal, antitumor, antiviral, antineoplastic, and antioxidant compounds [14,15]. Some of the cytotoxic compounds reported from this genus such as demethylcisterol A3 (1), ergosta-5,7,22-trien-3-ol (2), stigmastan-3-one (3), stigmast-4-en-3-one (4), stigmast-4-en-6-ol-3-one (5), and flufuran (6) (Figure 2), were discovered from *Pestalotiopsis* spp., associated with Chinese mangrove *Rhizophora mucronata*. Compounds 2–6 showed cytotoxicity against human cancer cell lines HeLa, A549, and HepG, with IC₅₀ values in the range of 11.44–102.11 μM. Compound 1 had the most potential, with IC₅₀ values reaching the nM activity level from 0.17 to 14.16 nM. Flow cytometric investigation demonstrated that compound 1 inhibited the cell cycle at the G0/G1 phase in a dose-dependent manner with a significant induction of apoptosis on the three tested cell lines. The involvement of the mitochondria in compound-1-induced apoptosis was demonstrated using MMP [16].

The compounds 7-O-methylnigrosporolide (7), pestalotioprolides D–F (8, 9, 10) (Figure 2), were extracted from the *Pestalotiopsis microspora*, endophytic fungus obtained from the fruits of *Drepanocarpus lunatus* collected from Douala, Cameroon. An approximately ten-fold increase in the yield of compounds 9 and 10 compared to axenic fungal control was observed when *P. microspora* was co-cultured with *Streptomyces lividans*. Compounds 7–10 exhibited cytotoxicity against the L5178Y cell line with IC₅₀ values of 0.7, 5.6, 3.4, and 3.9 μM, respectively, and compound 9 also showed potent activity against the A2780 cell line displaying an IC₅₀ value of 1.2 μM [17].

Another study by Hemphill et al. reported a new compound pestalpolyol I (11) (Figure 2) of the polyketide group from *Pestalotiopsis clavispora*, the endophytic fungus obtained from petioles of the *Rhizophora harrisonii*, growing in Port Harcourt (Nigeria). Compound 11 showed cytotoxicity against the L5178Y cell line with an IC₅₀ value of 4.10 μM [18].

A new aromatic amine, pestalamine A (12) (Figure 2), was isolated from *P. vaccinia* from a branch of *Kandelia candel*, a viviparous mangrove species widely distributed in coastal and estuarine areas

of southern China. The structure of pestalamine A **12** was determined by spectroscopic methods, especially 2D NMR analyses. Compound **12** showed moderate cytotoxicity against MCF-7, HeLa, and HepG2 human cancer cell lines with IC₅₀ values of 40.3, 22.0, and 32.8 μ M, respectively [19].

Phomazines B (**13**), epicorazine A (**14**), epicorazine B (**15**), epicorazine C (**16**), exserohilone A (**17**) (Figure 2), were isolated from an endophytic fungus, *Phoma* sp. OUCMDZ-1847 associated with the fruit of *Kandelia candel* collected in Wenchang, Hainan Province, China. Compounds **13–17** showed cytotoxicity against the HL-60, HCT-116, K562, MGC-803, and A549 cell lines with IC₅₀ values in the range of 0.05 to 8.5 μ M [20].

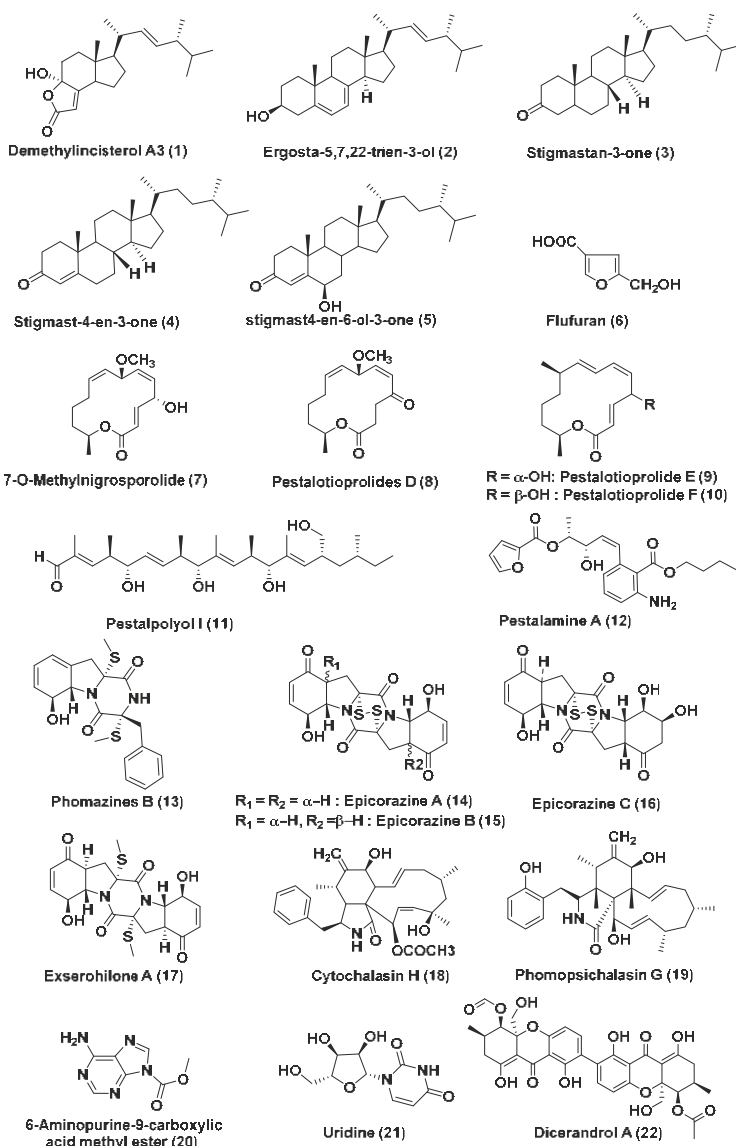


Figure 2. Structures of metabolites isolated from Coelomycetes (1–22).

Cytochalasin H (**18**) (Figure 2), was identified from *Phomopsis* sp., an endophytic fungus of mangrove origin in Zhanjiang, China. Compound **18** was shown to inhibit the cell cycle of A549 cells at the G2/M phase. Additionally, DNA fragmentation along with a decrease in the transmembrane potential of mitochondria was observed in A549 cells. It was also shown to regulate the expression level of Bax, P53, Bcl-xL, and Bcl-2. On treatment with Cytochalasin H, the migration capability was impaired in a dose-dependent manner [21].

A new cytochalasin, phomopsichalasin G (**19**) (Figure 2), was isolated from *Phomopsis* sp. xy21, and xy22, associated with *Xylocarpus granatum*, collected in Trang Province, Thailand. Compound **19** exhibited inhibitory activities against HCT-8, HCT-8/T, A549, MDA-MB-231, and A2780 cancer cell lines with IC₅₀ values of 7.5, 8.6, 6.4, 3.4, and 7.1 μM, respectively [22].

6-Aminopurine-9-carboxylic acid methyl ester (**20**) and uridine (**21**) (Figure 2), were isolated from *Phomopsis longicolla* HL-2232, an endophyte of *Bruguiera sexangula* var. *rhynchopetala*. The compounds **20** and **21** exhibited cytotoxicity against MCF-7 and A549 cell lines with IC₅₀ values of 14.9 and 8.6 μM, respectively [23].

Phomopsis sp. HNY29-2B, an endophyte isolated from the branch of *Acanthus ilicifolius* collected from the South China Sea in Hainan province, China, is reported as a source of the known phomoxanthones, including dicerandrol A (**22**) (Figure 2), dicerandrol B (**23**), dicerandrol C (**24**), diacetylphomoxanthone B (**25**) and penexanthone A (**26**) (Figure 3). Compound **22** exhibited broad-spectrum cytotoxic activity against MDA-MB-435, HCT-116, Calu-3 and Huh7 cell lines with IC₅₀ values of 3.03, 2.64, 1.76 and 4.19 μM, respectively. Compound **23** and **26** showed potent cytotoxic activities against MDA-MB-435, HCT-116, and Calu-3 (IC₅₀ < 10 μM), and displayed poor cytotoxic activity effects against the MCF-10A cell line (IC₅₀ > 50 μM). Compound **24** exhibited cytotoxic activity against the MDA-MB-435, HCT-116, Calu-3, MCF-10A cell lines with IC₅₀ values of 44.10, 42.63, 36.52, and 33.05 μM, respectively. Compound **25** exhibited cytotoxicity against the MDA-MB-435, HCT-116, Calu-3, Huh7 cell lines with IC₅₀ values of 14.40, 7.12, 4.14 and 29.20 μM and showed no cytotoxic effect on the MCF-10A cell line [24].

A new xanthone O-glycoside, 3-O-(6-O-α-L-arabinopyranosyl)-β-D-glucopyranosyl-1,4-dimethoxyxanthone (**27**) (Figure 3), was isolated from an endophytic fungus, *Phomopsis* sp. (ZH7) obtained from the stem of *Excoecaria agallocha* of a mangrove from Dong Sai on the South China Sea coast. Compound **27** had an inhibitory effect on the growth of HEp-2 and HepG2 cells displaying IC₅₀ values of 9 and 16 μM, respectively [25].

A highly oxygenated chloroazaphilone derivative, isochromophilone D (**28**), and a known analogue epi-Isochromophilone II (**29**) (Figure 3), were isolated from *Diaporthe* sp. SCSIO 41011, the endophytic fungus associated with *Rhizophora stylosa*, collected in Sanya city, Hainan Province, China. Compound **29** exhibited cytotoxic activity against the ACHN, OS-RC-2, and 786-O cell lines with IC₅₀ values ranging from 3.0 to 4.4 μM. Sorafenib, a positive control, exhibited cytotoxic activity against the tested cell lines with IC₅₀ in the range of 3.4 to 7.0 μM. Compound **28** showed activity against 786-O cells with an IC₅₀ of 8.9 μM. In a dose- and time-dependent manner, apoptosis was induced in 786-O cells by compound **28** [26].

New chromeno[3,2-c] pyridine, 5-deoxybostrycoïdin (**30**), and fusaristatin A (**31**) (Figure 3), were isolated from an endophytic fungus *Diaporthe phaseolorum* SKS019 isolated from the branches of the mangrove plant *Acanthus ilicifolius*, collected from Shankou in Guangxi province, China. Compound **30** exhibited cytotoxicity against the MDA-MB-435, and NCI-H460 human cancer cell lines with IC₅₀ values of 5.32 and 6.57 μM, respectively, and compound **31** showed growth-inhibitory activity against the MDA-MB-435 human cancer cell line with an IC₅₀ value of 8.15 μM [27].

Mycoepoxydiene (**32**) and deacetylmycoepoxydiene (**33**) (Figure 3) were isolated from a mangrove endophytic fungus *Phomopsis* sp. A818 isolated from the foliage of *Kandelia candel*, collected from the Mangrove Nature Conservation Area of Fugong, Fujian Province, China. Compounds **32** and **33** had IC₅₀ values of 7.85 and 14.61 μM, respectively, against MDA-MB-435 [28]. Mycoepoxydiene (**32**) (Figure 3), significantly suppressed antigen-stimulated degranulation and cytokine production in

mast cells and IgE-mediated passive cutaneous anaphylaxis in mice. Compound 32 suppressed antigen-induced activation of Syk, and subsequently inhibited the phosphorylation of PLC γ 1, Akt, and MAPKs such as extracellular signal-regulated kinase, c-jun N-terminal kinase, and p38 in mast cells. Mycoepoxydiene can inhibit the activation of mast cells and protect mice from a mast-cell-mediated allergic response through inhibiting the activation of Syk [29].

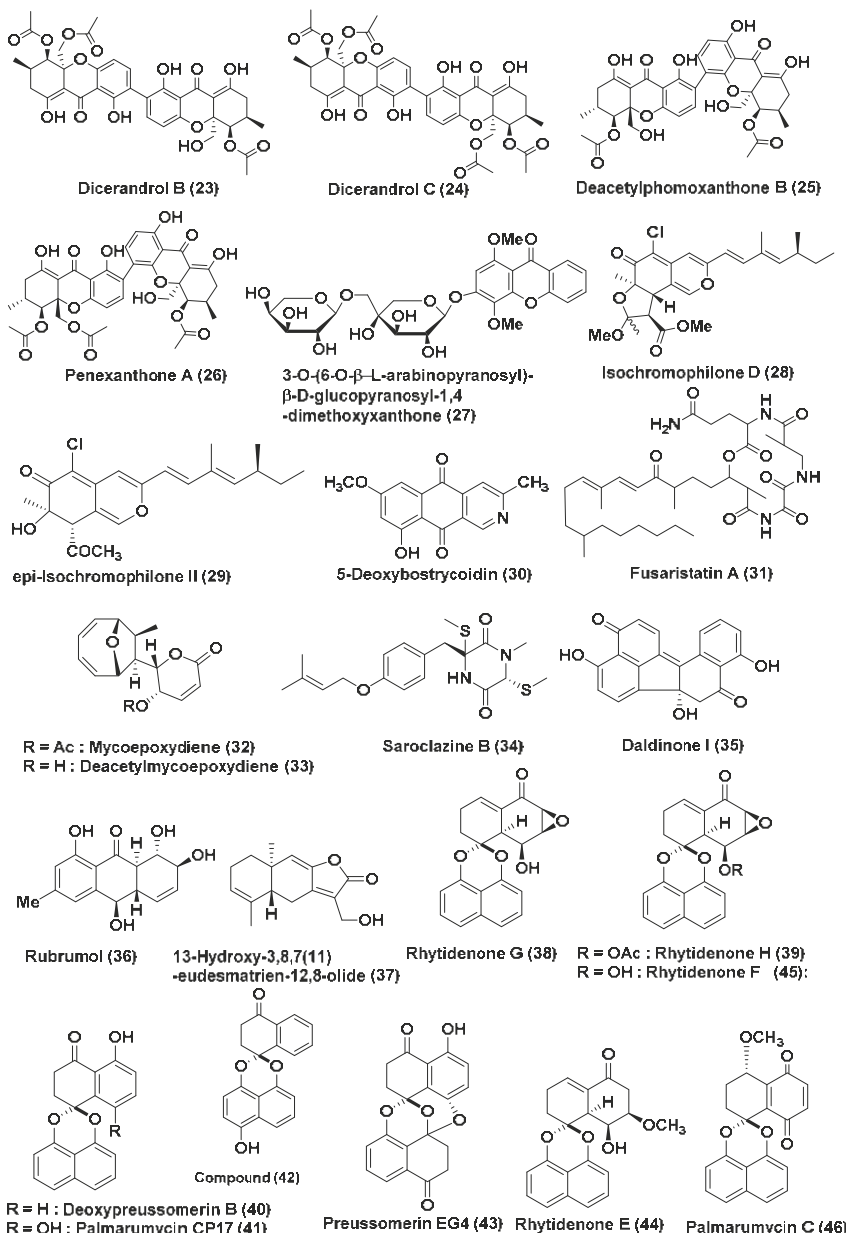


Figure 3. Structures of metabolites isolated from Coelomycetes (23–33) and Ascomycetes (34–46).

2.2. Compounds Produced by Ascomycetes

A new diketopiperazine derivative, saroclazine B (34) (Figure 3), was isolated from the mangrove-derived fungus *Sarocladium kilicense* HDN11-84 isolated from the rhizosphere soil of the mangrove plant *Thespesia populnea*, collected in Guangxi Province, China. Compound 34 showed cytotoxicity against HeLa cell lines with an IC₅₀ value of 4.2 μM [30].

Benzofluoranthene metabolites and daldinone I (35) (Figure 3) were extracted from *Annulohypoxylon* sp., an endophytic fungus associated with *Rhizophora racemosa*, collected in Cameroon. Compound 35 exhibited average to potent cytotoxicity with IC₅₀ values of 14.1 and 6.6 μM, against Jurkat J16 and Ramos cell lines, respectively. It was reported that compound 35 induces apoptotic cell death caused by the induction of intrinsic apoptosis [31].

A new anthraquinone rubrumol (36) (Figure 3) with poly-hydroxyl groups was isolated from a halo-tolerant endophytic fungus *Eurotium rubrum*, isolated from the salt-tolerant wild plant *Suaeda salsa* L. collected from the “BoHai” seaside, China. The biological effect of compound 36 on Topo I to relax supercoiled pBR322 DNA was investigated in the cleavable complex assay. The results indicated that compound 36 displayed biological activity compared to the positive control camptothecin. The relaxation activity of rubrumol (36) was stronger than that of camptothecin at the concentration of 100 μM. The band backward shifting and trailing of rubrumol (36) was observed at 100, 50, 10, 5 and 1 μM. Compound 36 also exhibited cytotoxic activities against A549, MDA-MB-231, PANC-1 and HepG2 human cancer cell lines, by MTT method. The inhibition rate for compound 36 against these four cancer cell line was less than 60% at 100 μg/mL, which implied that it displayed no significant cytotoxic activity [32].

The 13-Hydroxy-3,8,7(11)-eudesmatrien-12, 8-oxide (37) (Figure 3) was isolated from *Eutypella* sp. 1–15 isolated from the soil of the mangrove rhizosphere in Jimei, Fujian Province, China. Compound 37 exhibited potent anticancer activity against JEKO-1 and HepG2 with IC₅₀ values of 8.4 and 28.5 Mm, respectively [33].

Rhytidones G (38), H (39), deoxypreussomerin B (40), palmarumycin CP17 (41), 1-oxo-1,4-dihydronaphthalene-4-spiro-20-naphtho[400-hydroxy-100,800-de][10,30]-dioxine (42), preussomerin EG4 (43), rhytidonenone E (44), rhytidonenone F (45), palmarumycin C5 (46), (Figure 3), and 4,8-dihydroxy-3,4-dihydronaphthalen-1(2H)-one (47) (Figure 4), were isolated from *Rhytidhysteron rufulum* AS21B, an endophytic fungus associated with the leaves of *Azima sarmentosa*, collected from the mangrove forest in Samutsakhon province, Thailand. The culture in acidic medium enhanced the production of compounds 41 and 45, with a four-fold and eight-fold increase, respectively, which are present in minor quantities under normal culture condition. Compounds 38–47 exhibited cytotoxicity against Ramos cells with IC₅₀ values of 17.98, 0.018, 18.00, 33.1, 15, 82.9, 0.461, 0.048, 31.7 and 23.1 μM, respectively, while the control Ibrutinib exhibited cytotoxicity of 28.7 μM against the same cell line. Compounds 38–39, 44–45, and 47 exhibited cytotoxicity against the H1975 cell lines with an IC₅₀ value of 7.3, 0.252, 10.24, 1.17 and 50 μM, respectively, while control afatinib exhibited cytotoxicity of 1.97 μM against the same cell line [34].

Lasiodiplodia sp. 318#, an endophytic fungus associated with *Excoecaria agallocha*, collected from Guangdong Province, China, was the source of compound 2,4-Dihydroxy-6-nonylbenzoate (48) (Figure 4). Compound 48 exhibited cytotoxicity against the MMQ and GH3 cell lines with IC₅₀ values of 5.2 and 13.0 μM, respectively [35]. Previously, a new lasiodiplodin—ethyl-2,4-dihydroxy-6-(8'-hydroxynonyl)-benzoate (49)—was obtained from the same fungus. Compound 49 exhibited average cytotoxicity against the MDA-MB-435, HepG2, HCT-116, A549, and leukaemia THP1 cell lines with IC₅₀ values of 10.13, 12.50, 11.92, 13.31 and 39.74 μM, respectively [36].

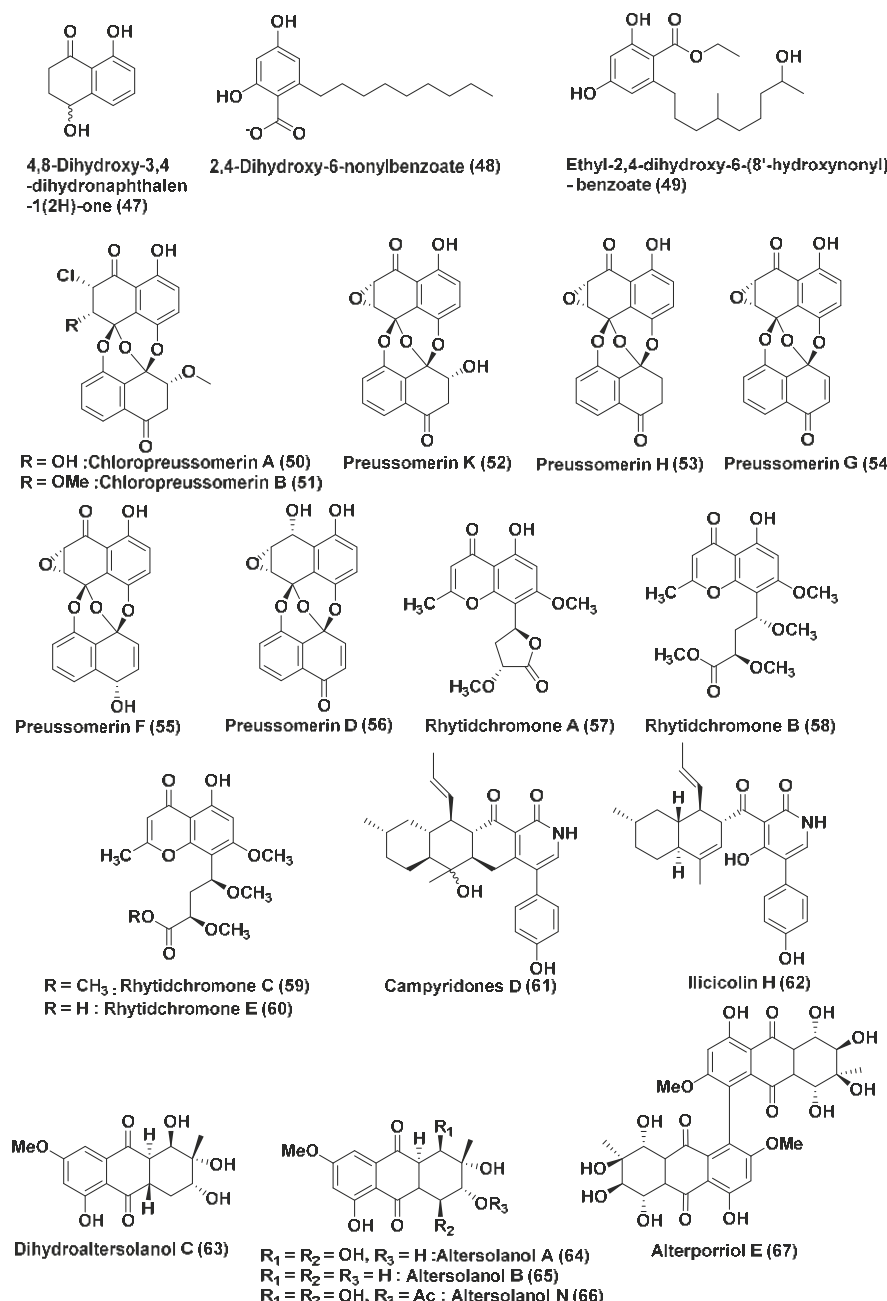


Figure 4. Structures of metabolites isolated from Ascomycetes (47–67).

Two new chlorinated preussomerins, chloropreussomerins A (50), and B (51), and a known preussomerin analog preussomerin K (52), preussomerin H (53), preussomerin G (54), preussomerin F (55), preussomerin D (56) (Figure 4), were obtained from the endophytic fungus *Lasiodiplodia theobromae*

ZJ-HQ1 isolated from *Excoecaria agallocha* collected from Guangdong Province, China. Compounds **50–51** and **56** were found to be active against the A549 and MCF-7 cell lines with IC₅₀ values ranging from 5.9–8.9 μM, and compounds **52–55** showed cytotoxicity against A549, HepG2, and MCF-7 human cancer cell lines with IC₅₀ values of 2.5–9.4 μM [37].

Four highly oxygenated chromones—rhytidchromone A (**57**), B (**58**), C (**59**) and E (**60**) (Figure 4)—were obtained from *Rhytidhysteron rufulum* BG2-Y, an endophyte associated with the leaves of *Bruguiera gymnorhiza* collected from Prachuab Kiri Khan Province, Thailand. Compounds **57–60** were found to be active against Kato-3 cell lines, with IC₅₀ values in the range of 16.0–23.3 μM, while the rhytidchromones A and C were active, with IC₅₀ values of 19.3 and 17.7 μM respectively, against MCF-7 cell lines [38].

Campyridone D (**61**) and ilicicolin H (**62**) (Figure 4) were extracted from *Campylocarpon* sp. HDN13-307, an endophyte associated with the root of the mangrove plant *Sonneratia caseolaris*. Compounds **61** and **62** showed cytotoxicity, with the IC₅₀ values of 8.8 μM and 4.7 μM, respectively, against HeLa cells [39].

Stemphylium globuliferum, an endophytic fungus associated with the Egyptian mangrove plant *Avicennia marina*, was the source of dihydroaltersolanol C (**63**), altersolanols A, B, N (**64**, **65**, **66**), and alterporriol E (**67**) (Figure 4) [40]. Compounds **63**, **64**, **65**, and **67** showed cytotoxicity with IC₅₀ values of 3.4, 2.5, 3.7 and 6.9 μM, respectively, towards L5178Y cells [41]. Compound **66** also showed good activity, with IC₅₀ values in the low micro-molar range towards L5178Y cells [42]. Mishra et al. [43] reported that compound **64** exhibited cytotoxicity against 34 human cancer cell lines in vitro, with mean IC₅₀ (IC₇₀) values of 0.005 μg/mL (0.024 μg/mL). It has also been reported that compound **64** is a kinase inhibitor and induces cell death by apoptosis through the caspase-dependent pathway, and that kinase inhibition might be the mechanism for the cytotoxic activity [44]. The pro-apoptotic and anti-invasive activity of compound **64** that occurred through the inhibition of the NF-κB transcriptional activity may be responsible for its antitumor potential [45].

Two new hydroanthraquinones, paradictyoarthrins A (**68**) and B (**69**) and the known compounds, preussomerin C (**70**), ymf 1029C (**71**) and altenusin (**72**) (Figure 5), were isolated from *Paradictyoarthrinium diffractum* BCC 8704, an endophyte associated with mangrove wood in Laem Son National Park, Ranong Province, Thailand. Compounds **68–72** were evaluated for cytotoxic activity against cancer cell-lines, KB, MCF-7, and NCI-H187, and noncancerous Vero cells. Compound **69** exhibited moderate cytotoxicity against KB, MCF-7 NCI-H187 and Vero cell lines with IC₅₀ of 3.1, 3.8, 9.5 and 5.6 μg/mL, respectively, whereas Compound **68** showed weaker activity with IC₅₀ values in the range of 23–31 μg/mL. Compounds **70–72** showed average to poor activity in tested cell lines. Compound ymf 1029C **71** showed relatively stronger cytotoxicity on NCI-H187 cells than other cell-lines (IC₅₀ 5.0 μg/mL) [46].

A marine anthraquinone derivative SZ-685C (**73**) (Figure 5) has been isolated from the mangrove endophytic fungus *Halorosellinia* sp. (No. 1403), which was found in the South China Sea. The IC₅₀s of SZ-685C in nonfunctioning pituitary adenoma (NFPA), MMQ, and RPC cells were 18.76, 14.51, and 56.09 μM, respectively. Hoechst 33342 dye/propidium iodide (PI) double staining and fluorescein isothiocyanate-conjugated Annexin V/PI (Annexin V-FITC/PI) apoptosis assays detected an enhanced the rate of apoptosis in cells treated with SZ-685C. Enhanced expression levels of caspase 3 and phosphate and tensin homologs were determined by Western blotting. The protein expression levels of Akt were decreased when the primary human NFPA cells were treated with SZ-685C. It has been observed that SZ-685C (**73**) induces the apoptosis of human NFPA cells through the inhibition of the Akt pathway in vitro. These findings suggest that SZ-685C may be a potentially promising Akt inhibitor and anti-cancer agent for the treatment of NFPA [47].

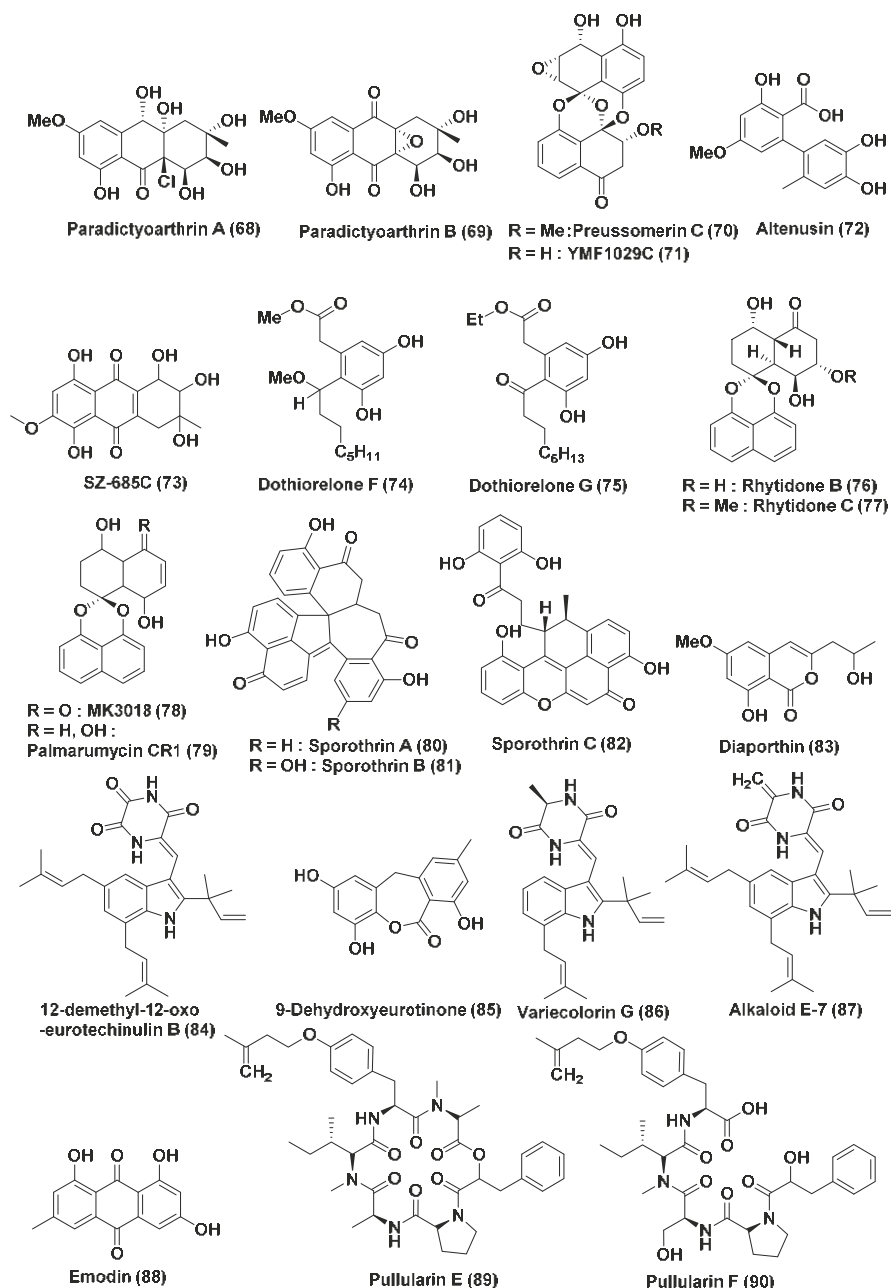


Figure 5. Structures of metabolites isolated from Ascomycetes (68–90).

SZ-685C (73), was previously reported to inhibit the proliferation of certain tumor cells. SZ-685C inhibited MMQ cell growth in a dose-dependent manner but showed little toxicity toward rat pituitary cells. The IC₅₀ of SZ-685C in MMQ cells and RPCs were 13.2 and 49.1 μ M, respectively. Increasing

numbers of apoptotic cells were observed in response to escalating concentrations of SZ-685C, and the expression level of prolactin was inhibited. Nevertheless, the level of prolactin mRNA was unchanged. Additionally, miR-200c was upregulated in MMQ cells compared with RPCs, and downregulation of miR-200c was observed in SZ-685C-treated MMQ cells. Furthermore, the overexpression of miR-200c weakened the effect of the SZ-685C-induced apoptosis of MMQ cells. It has been suggested that SZ-685C induces MMQ cell apoptosis in a miR-200c-dependent manner [48].

Two new polyketides, named dothiorelons F (74) and G (75) (Figure 5), were isolated from *Dothiorella* sp., an endophytic fungus associated with the bark of the mangrove tree *Aegiceras corniculatum* at the estuary of Jiulong River, Fujian Province, China. Compounds 74 and 75 showed significant cytotoxicity against the Raji cancer cell line, with an IC₅₀ value of 2 µg/mL [49].

New spironaphthalenes, rhytidones B–C (76, 77) and known MK3018 (78), palmarumycin CR1 (79) (Figure 5), were extracted from *Rhytidhysteron* sp. an endophytic fungus associated with the leaves of *Azima sarmentosa*, collected from the mangrove forest in Samutsakhon province, Thailand. Compound 76 was found to be poorly active against CaSki cells, with an IC₅₀ value of 22.81 µM, while compounds 77–79 showed average activity against the MCF-7 and CaSkicell lines with IC₅₀ values in the range of 14.47 and 25.59 µM [50].

Sporothrix sp. isolated from the bark of an inshore mangrove sample *Kandelia candel* in the South China Sea was the source of Sporothrin A (80), sporothrin B (81), sporothrin C (82), diaporthin (83) (Figure 5). Compounds 81–83 were found to show weak cytotoxic activity with IC₅₀ values of 20, 23, and 23 g/mL, respectively. Compound 80 exhibited strong inhibition of AChE in vitro (IC₅₀ was 1.05 µM) [51].

A new dioxopiperazine alkaloid, 12-demethyl-12-oxo-eurotechnulin B (84), and one new anthraquinone derivative 9-dehydroxyeurotinone (85), and known compounds, variecolorin G (86), alkaloid E-7 (87), and emodin (88) (Figure 5), were isolated from *Eurotium rubrum*, an endophytic fungus associated with the inner tissue of *Hibiscus tiliaceus*, collected from Hainan Island, China. Compounds 84–88 displayed cytotoxic activity against one or two of the MCF-7, SW1990, HepG2, NCI-H460, SMMC7721, Hela, and Du145 cell lines in the range of 15–30 mg/mL [52].

Two new peptides—pullularins E (89) and F (90) (Figure 5)—and three known compounds—pullularins A (91) and C (92) and verticillin D (93) (Figure 6)—were extracted from *Bionectria ochroleuca*, an endophytic fungus associated with the inner leaf tissues of *Somneratia caseolaris* from Hainan island, China. Compound 93 showed potent to moderate activity against L5178Y cell lines with an EC₅₀ value of <0.1 µg/mL. Compounds 89–92 also exhibited potent to average activity against the L5178Y cell lines, with EC₅₀ values ranging between 0.1 and 6.7 µg/mL [53].

2.3. Compounds Produced by Hyphomycetes

New isocoumarin derivatives, aspergisorcoumrins A (94) and B (95) (Figure 6) were obtained from the culture of the endophytic fungus *Aspergillus* sp. HN15-5D derived from the fresh leaves of the mangrove plant *Acanthus ilicifolius* collected from Dongzhaigang Mangrove National Nature Reserve on Hainan Island, China. Compounds 94 and 95 exhibited cytotoxicity against MDA-MB-435, with IC₅₀ values of 5.08 and 4.98 µM, respectively [54].

Two new 6,8(14),22-hexadehydro-5α,9α-epidioxy-3,15-dihydroxy sterols, nigerasterols A (96) and B (97) (Figure 6), were isolated from *Aspergillus niger* MA-132, an endophytic fungus associated with *Avicennia marina*. Compounds 96–97 displayed cytotoxicity against the HL60 cell line with an IC₅₀ value of 0.30 and 1.50 µM, respectively. Compounds 96–97 also exhibited potent activity against the A549 cell line with IC₅₀ values of 1.82 and 5.41 µM, respectively [55].

Four new quinazolinone alkaloids, namely, aniquinazolines A–D (98–101) (Figure 6), were identified from *Aspergillus nidulans* MA-143, an endophytic fungus associated with the leaves of the marine mangrove plant *Rhizophora stylosa*. Compounds 98–101 showed potent lethality against brine shrimp with LD₅₀ values of 1.27, 2.11, 4.95 and 3.42 µM, respectively, which were stronger than that of the positive control colchicine (with the LD₅₀ value of 88.4 µM) [56].

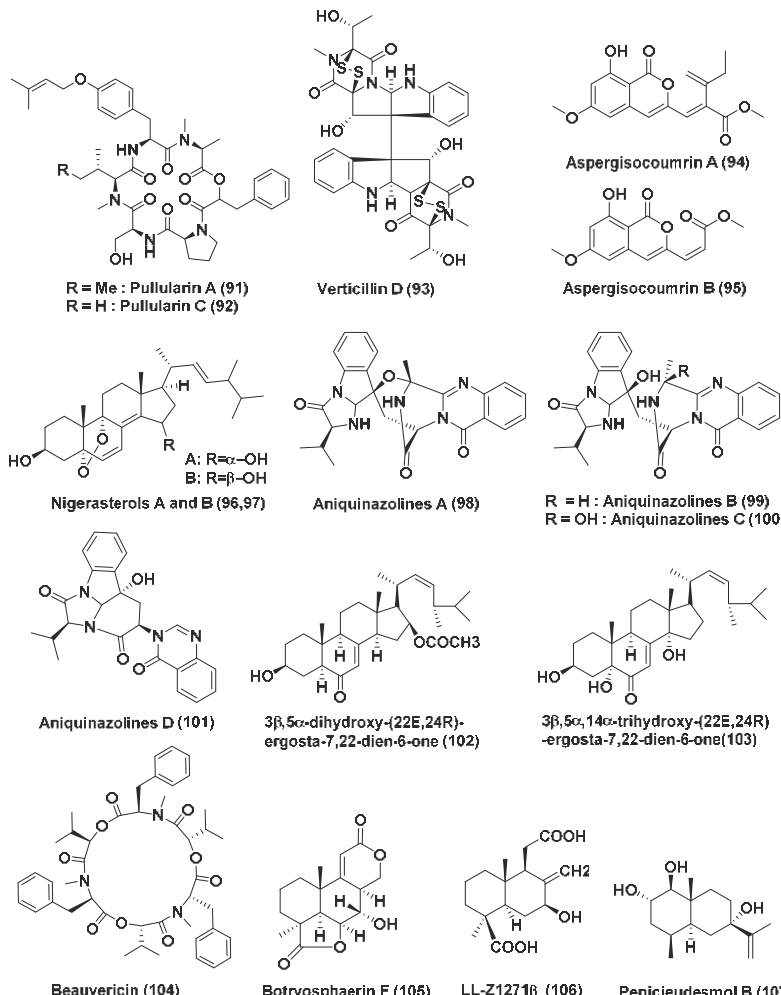


Figure 6. Structures of metabolites isolated from Ascomycetes (91–93) and Hyphomycetes (94–107).

The compounds 3 β ,5 α -Dihydroxy-(22E,24R)-ergosta-7,22-dien-6-one (**102**), 3 β ,5 α ,14 α -trihydroxy-(22E,24R)-ergosta-7,22-dien-6-one (**103**), and beauvericin (**104**) (Figure 6), were extracted from *Aspergillus terreus* (No. GX7-3B), a mangrove endophytic fungus isolated from the leaves of *Rhizophora stylosa*. Compounds **102** and **104** exhibited good or moderate cytotoxic activity against MCF-7, A549, HeLa and KB cell lines with IC₅₀ values of 4.98, 1.95, 0.68, 1.50 and 2.02, 0.82, 1.14, 1.10 μ M, respectively; compound **103** exhibited poor activity against the cell lines tested, namely MCF-7 (25.4 μ M), A549 (27.1 μ M), HeLa (24.4 μ M) and KB (19.4 μ M) [57].

A new compound, botryosphaerin F (**105**), and a known compound, LL-Z1271 β (**106**) (Figure 6), were obtained from the mangrove fungus *Aspergillus terreus* (No. GX7-3B) isolated from the branch of *Bruguiera gymnorhiza*, growing in a coastal salt marsh of the South China Sea in Guangxi Province, China. Compound **105** showed potent inhibiting activity towards the MCF-7 and HL-60 cancer cell lines with IC₅₀ values of 4.49 and 3.43 μ M, respectively, and compound **106** exhibited promising activity against the HL-60 cell line with an IC₅₀ value of 0.6 μ M [58].

A new eudesmane-type sesquiterpenoid, penicieudesmol B (**107**) (Figure 6), was isolated from the mangrove-derived endophytic fungus *Penicillium* sp. J-54 associated with the leaves of *Ceriops tagal*, which were collected in Dong Zhai Gang Mangrove Reserve in Hainan province, China. Compound **107** exhibited weak cytotoxicity against K-562 with an IC₅₀ value of 90.1 μM, with paclitaxel as the positive control (IC₅₀ = 9.5 μM) [59].

A new compound Penibenzophenone B (**108**) (Figure 7), was obtained from the endophytic fungus *Penicillium citrinum* HL-5126 isolated from the mangrove *Bruguiera sexangula* var. *rhynchopetala* collected in the South China Sea. The new compound **108** displayed cytotoxic activity against human A549 cell lines with an IC₅₀ value of 15.7 μg/mL [60].

Five new derivatives of macrolide antibiotic Brefeldin A (**109**), along with Brefeldin A 7-O-acetate (**110**) (Figure 7), were produced by an endophytic fungus, *Penicillium* sp., which was isolated from the healthy root of *Panax notoginseng*. Compounds **109–110** exhibited cytotoxic activity against the 293, HepG2, Huh7 and KB cell line with an ID₅₀ values from 0.024 to 0.62 μM. Further, studies of the cellular mechanism of compounds **109–110** showed that they arrested HepG2 cells at the S phase [61].

A new chaetoglobosin, penochalasin K (**111**) (Figure 7), was extracted from the mangrove endophytic fungus *Penicillium chrysogenum* V11. Its structure was elucidated by 1D, 2D NMR spectroscopic analysis and high resolution mass spectroscopic data. Compound **111** showed strong cytotoxicity against the MDA-MB-435, SGC-7901 and A549 cell lines with an IC₅₀ less than 10 μM [62].

The new epipolythiodioxopiperazine alkaloids, penicisulfuranols A–C (**112–114**) (Figure 7), were obtained from *Penicillium janthinellum* HDN13-309, the mangrove endophytic fungus associated with *Sonneratia caseolaris* collected from Hainan Province, China. Compounds **112–114** showed cytotoxicity against HeLa and HL-60 cell lines, with IC₅₀ values ranging from 0.1 to 3.9 μM [63].

Penicillium chrysogenum V11, a mangrove endophytic fungus was the source of a novel chaetoglobosin named penochalasin I (**115**), along with chaetoglobosins A (**116**), and cytoglobosin C (**117**) (Figure 7). Compound **115** exhibited marked cytotoxicity against MDA-MB-435 and SGC-7901 cells (IC₅₀ < 10 μM), and compounds **116** and **117** showed potent cytotoxicity against SGC-7901 and A549 cells (IC₅₀ < 10 μM) [64].

Using the one strain many compounds (OSMAC) approach, new diketopiperazines, spirobrocazine C (**118**) and brocazine G (**119**) (Figure 7) were characterized from *Penicillium brocae* MA-231, an endophytic fungus associated with *Avicennia marina* collected at Hainan Island, China. Compound **119** exhibited potent cytotoxic activity against the A2780 and A2780 CisR cell lines, with IC₅₀ values of 664 and 661 nM, respectively. This activity is higher than the cisplatin where the IC₅₀ values were reported as 1.67 and 12.63 μM, respectively. Compound **118** showed moderate activity against A2780 cells with an IC₅₀ value of 59 μM [65]. Previously reported disulfide-bridged diketopiperazine derivatives, brocazines A (**120**), B (**121**), E (**122**), F (**123**) (Figure 7) were isolated from the same fungus. Compounds **120–123** displayed cytotoxic activity against the Du145, HeLa, HepG2, MCF-7, NCI-H460, SGC-7901, SW1990, SW480, and U251 cell lines with IC₅₀ values ranging from 0.89 to 9.0 μM. Compounds **121** and **122** exhibited strong activity against the SW480 cell line with IC₅₀ values of 2.0 and 1.2 μM, respectively, while compound **123** showed potent activity against the DU145 and NCI-H460 cell lines, with IC₅₀ values of 1.7 and 0.89 μM, respectively [66].

The fungal metabolites TMC-264 (**124**), and PR-toxin (**125**) (Figure 7) were obtained from *Penicillium chermesinum* strain HLit-ROR2, an endophytic fungus associated with the root of *Heritiera littoralis*, collected from Samut Sakhon province, Thailand. Compound **124** showed cytotoxicity against T47D and MDA-MB231 cell lines with the IC₅₀ value of 1.08 and 2.81 μM, respectively, while a positive control, doxorubicin, showed activity with IC₅₀ of 1.55 and 2.24 μM respectively. Compound **124** also showed cytotoxic activity against the HepG2 cell line, with an IC₅₀ value of 3.27 μM, which was 11 times more potent than that of etoposide, an anticancer drug (IC₅₀ 35.66 μM). Besides, compound **124** selectively exhibited cytotoxic activity toward MOLT-3 and T47D cancer cells with the IC₅₀ values of 1.36 and 1.08 μM and with selectivity index (SI) values of 9 and 11, respectively (IC₅₀ 12.64 μM for the normal cell line, MRC-5). Compound **125** exhibited cytotoxicity against the HuCCA-1, HeLa, T47D,

and MDA-MB231 cell lines with IC₅₀ values in the range of 0.81–2.19 µM, which was comparable to that of doxorubicin (IC₅₀ 0.26–2.24 µM). Interestingly, the compound **125** exhibited cytotoxicity against the HL-60 cell line with an IC₅₀ value of 0.06 µM, which is superior to doxorubicin (IC₅₀ 1.21 µM). Compound **125** selectively showed cytotoxic activity against MOLT-3 and HL-60 cancer cell lines with IC₅₀ of 0.09 and 0.06 µM, respectively, with respective SI values of 40 and 61 (IC₅₀ 3.66 µM for the MRC-5 cell line) [67].

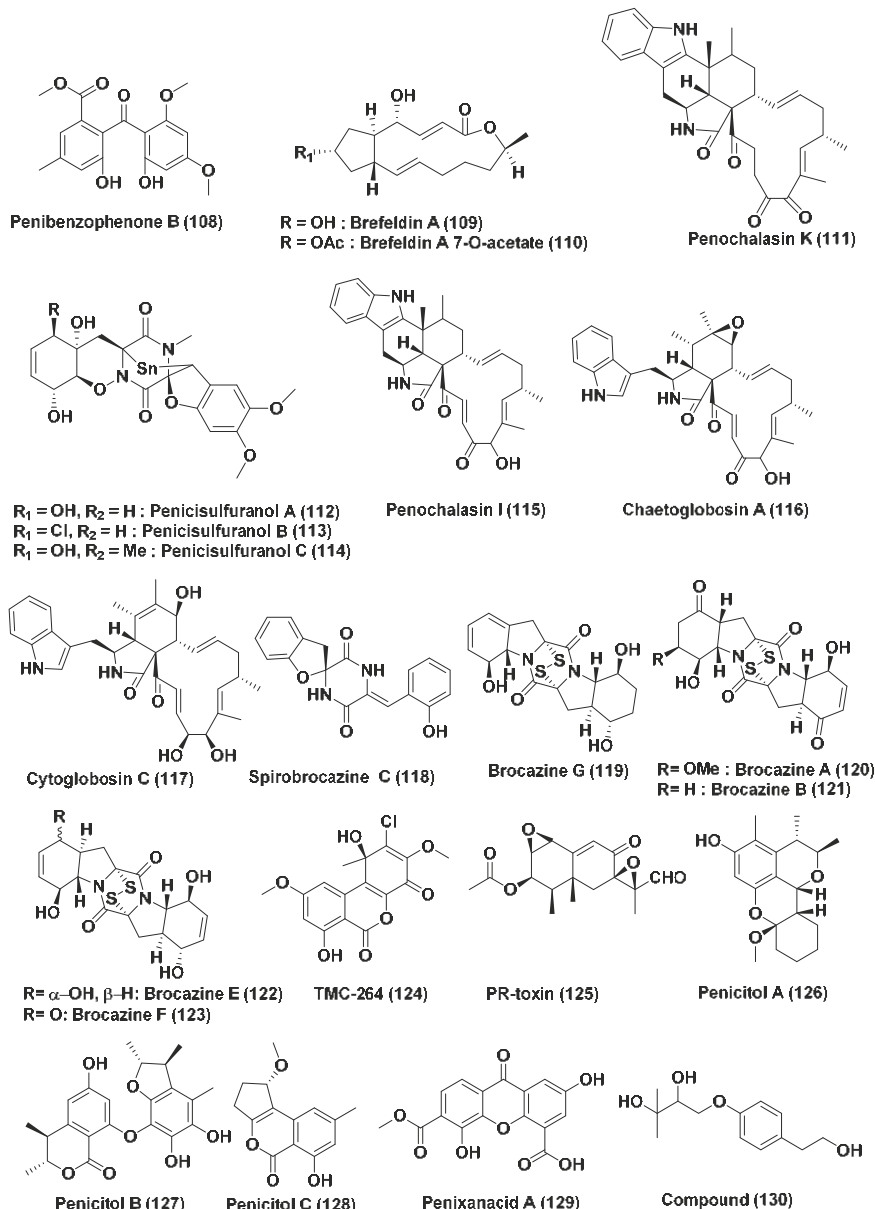


Figure 7. Structures of metabolites isolated from Hyphomycetes (108–130).

Three new citrinin analogs, penicitols A–C (**126–128**), and one new xanthone derivative, penixanacid A (**129**) (Figure 7), were isolated from *Penicillium chrysogenum* HND11–24 isolated from the rhizosphere soil of *Acanthus ilicifolius*. Compounds **126–127** showed good activity against the HeLa, BEL-7402, HEK-293, HCT-116, and A549 cell lines with IC₅₀ values of 4.6–10.5 and 3.4–9.6 µM, respectively. Compounds **128–129** exhibited cytotoxicity against the HeLa, BEL-7402, HEK-293, HCT-116, and A549 cell lines with IC₅₀ values of 10–40.5 µM [68].

Two new metabolites, compounds **130** (Figure 7) and **131** (Figure 8), were obtained from *Penicillium* sp. FJ-1, associated with *Avicennia marina* collected in Fujian, China. Compound **131** exhibited anti-proliferative activity against Tca8113, MG-63 and the normal liver cell line WRL-68 with an IC₅₀ value of 10 µM, 55 nM and 58 µM, respectively. Compound **131** has also shown a significant inhibition of tumor growth of human osteosarcoma when tested against nude mice. Compound **130** exhibited weak activity against Tca8113 and MG-63 cells with the IC₅₀ values of 26 µM and 35 µM, respectively, in the anti-proliferative assay. Taxol, the positive control, showed activity against Tca8113 and MG-63 cell lines with IC₅₀ values of 46 nM and 10 nM, respectively [69].

Meleagrin (**132**) (Figure 8) was isolated from the endophytic fungus *Penicillium* sp. GD6, associated with the Chinese mangrove *Bruguiera gymnorhiza* collected off the coasts of Zhanjiang, China. Compound **132** showed potent cytotoxic activity against two tumor cell lines, HL60 and A549, with IC₅₀ values of 9.7 and 8.3 µM, respectively [70].

Mangrove endophytic fungus, *Penicillium* 303# obtained from the sea water in Zhanjiang Mangrove National Nature Reserve in Guangdong Province, China, was the source of compounds **133**, **134**, **135** and **136** (Figure 8). Compounds **133–135** exhibited cytotoxicity against MDA-MB-435, HepG2, HCT-116, and A549 with IC₅₀ value of 11.9–37.82 µg/mL. Compound **136** showed strong cytotoxic activity against the MDA-MB-435 cell lines with IC₅₀ values of 7.13 µM and average to low activity against the HepG2 and HCT-116 cell lines with IC₅₀ values of 39.64 and 27.80 µM, respectively [71].

A new isobenzofuranone, 4-(methoxymethyl)-7-methoxy-6-methyl-1(3H)-isobenzofuranone (**137**) (Figure 8), was isolated from the mangrove endophytic fungus, *Penicillium* sp. ZH58, which was associated with the leaves of mangrove tree *Avicennia* from Dong Sai, Hainan, on the South China Sea coast. Compound **137** showed cytotoxic activity against the KB and KB_V200 cell lines with IC₅₀ values of 6 and 10 µg/mL, respectively [72].

The new and rare sulfur-containing curvularin derivatives sumalarins A, B, (**138**, **139**) C (**140**) (Figure 8), and dehydrocurvularin (**141**) (Figure 8) were obtained from *Penicillium sumatrense* MA-92, isolated from the rhizosphere of the mangrove *Lumnitzera racemose* collected at Wen Chang on Hainan Island, China. Compounds **138–141** displayed cytotoxic activities against the Du145, HeLa, Huh 7, MCF-7, NCI-H460, SGC-7901, and SW1990 cell lines, with IC₅₀ values in the range of 3.8 to 10 µM [73].

Penicillium sp. ZH16, a mangrove endophytic fungus, obtained from the South China Sea, was the source of the furanocoumarin 5-methyl-8-(3-methylbut-2-enyl) furanocoumarin (**142**) (Figure 8). Compound **142** exhibited cytotoxicity against the KB and KB_V200 cell lines with IC₅₀ values of 5 and 10 µg/mL, respectively [74].

Zhang et al. [75] reported the production of the compounds (3S)-6-oxo-de-O-methylasiodiplodin (**143**), (3R)-de-O-methylasiodiplodin (**144**), (3R)-nordinone (**145**) (Figure 8), by the co-culturing of mangrove endophytic fungus *Trichoderma* sp. 307 isolated from the stem bark of *Clerodendrum inerme*, and the aquatic pathogenic bacterium *Acinetobacter johnsonii* B2 isolated from the stem bark of *Clerodendrum inerme*, collected in Zhanjiang Mangrove National Nature Reserve in Guangdong Province, China. Compound **144** exhibited good cytotoxic activity against the GH3 and MMQ cell lines with IC₅₀ values of 6.44 and 6.58 µM, respectively, and against rat normal pituitary cells (RPC) with an IC₅₀ value of 6.94 µM. Compound **145** displayed average cytotoxic activity against the GH3 and MMQ cell lines with IC₅₀ values of 12.33 and 10.13 µM, respectively, and against RPC cell lines with an IC₅₀ value of 100.03 µM. Compound **143** was less active against the GH3 and MMQ cell lines with IC₅₀ values of 21.42 and 13.59 µM, respectively, and against the RPC cell lines with an IC₅₀ value of 142.8 µM as a positive control [75].

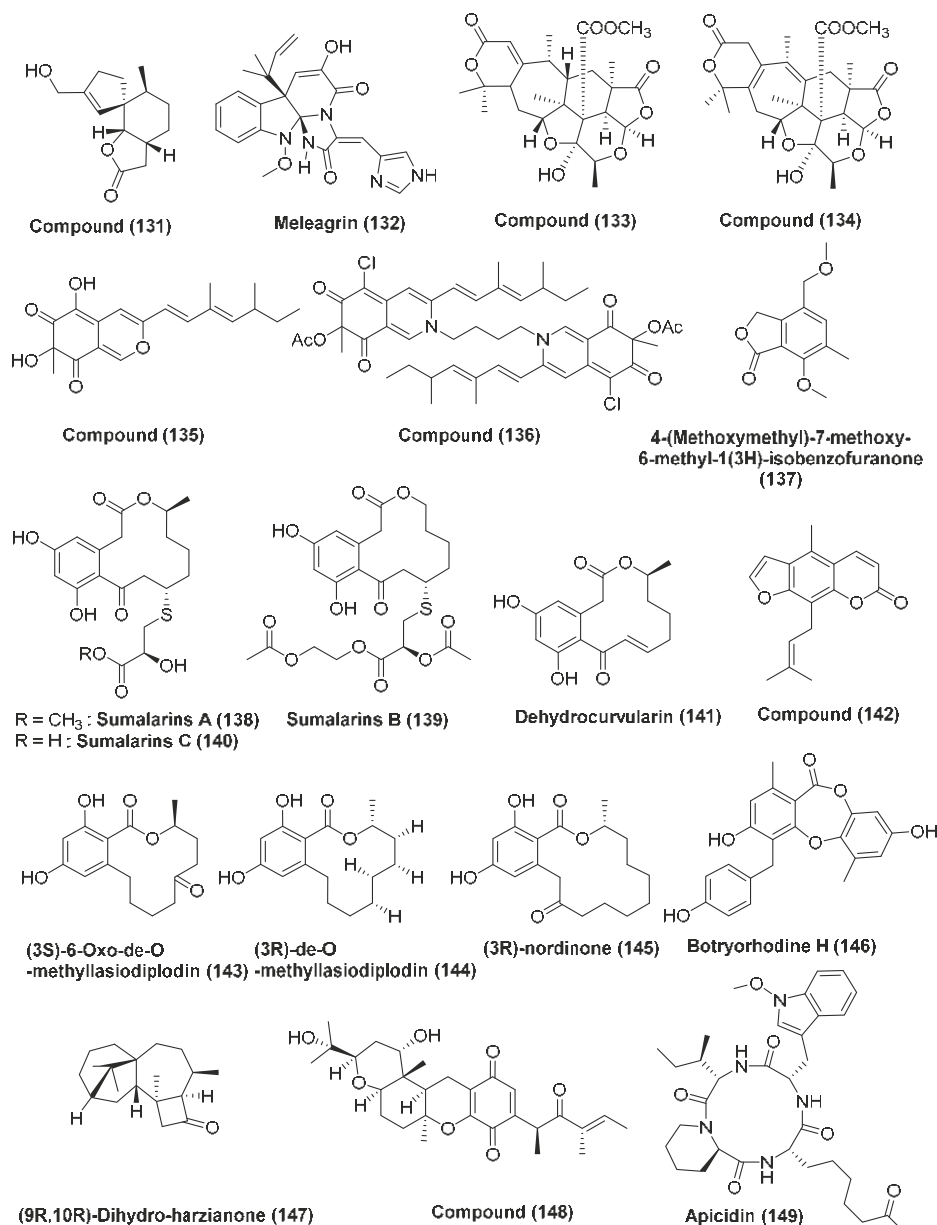


Figure 8. Structures of metabolites isolated from Hyphomycetes (131–149).

Zhang et al. [76] reported the production of a new depsidone, botryorhodine H (146) (Figure 8), by co-culturing mangrove endophytic fungus *Trichoderma* sp. 307 and *Acinetobacter johnsonii* B2. Compound 146 exhibited good cytotoxic activity against the MMQ and GH3 cell lines with IC₅₀ values of 3.09 and 3.64 μM, respectively.

A new harziane diterpenoid, named (9R,10R)-dihydro-harzianone (**147**) (Figure 8), was isolated from *Trichoderma* sp. Xy24, an endophytic fungus residing in the leaves, stems, and peels of the mangrove plant *Xylocarpus granatum*, collected in the Sanya district of Hainan province, China. Compound **147** was active against the HeLa and MCF-7 cell lines with IC₅₀ values of 30.1 µM and 30.7 µM, respectively [77].

The endophytic fungus *Nigrospora* sp. MA75 associated with *Pongamia pinnata*, which led to the isolation of a new compound 2,3-didehydro-19a-hydroxy-14-epicochlioquinone B (**148**) (Figure 8) when grown in medium containing 3.5% NaI. Compound **148** showed potent cytotoxicity against the MCF-7, SW1990, and SMMC7721 cell lines, with IC₅₀ values of 4, 5, and 7 µg/mL, respectively [78].

A known cyclic peptide, beauvericin (**104**) (Figure 6), was obtained from *Fusarium* sp. (No. DZ27) an endophytic fungus residing inside the bark of *Kandelia candel* from Dongzhai mangrove, Hainan, China, in the South China Sea. Compound (**104**) showed cytotoxic activity against the KB and KBv200 cell lines with IC₅₀ values of 5.76 and 5.34 µM, respectively. It induces apoptosis through the mitochondrial pathway, including the decrease of relative oxygen species generation, the loss of mitochondrial membrane potential, the release of cytochrome c, the activation of Caspase-9 and -3, and the cleavage of PARP. Additionally, the regulation of Bcl-2 or Bax was not involved in the apoptosis induced by beauvericin in KB and KBv200 cells [79].

An inhibitor of histone deacetylase, Apicidin (**149**) (Figure 8), was isolated from *Fusarium* sp., an endophytic fungus associated with the leaf of mangrove *Kandelia candel* planted at Dongzhai Harbor on Hainan Island, China. Apicidin showed good cytotoxic activity against GLC-82 cells with the IC₅₀ value of 6.94 ± 0.27 µM. Apicidin suppressed proliferation and invasion, and induced apoptosis via the mitochondrial pathway in GLC-82 cells, including the loss of ΔΨm, the release of cytochrome c from mitochondria, the activation of caspase-9 and -3, and the cleavage of poly-ADP-ribose polymerase [80]. Apicidin **149**, was previously isolated from the mangrove endophytic fungus ZZF42 from the South China Sea and exhibited selective in vitro cytotoxicity towards KB and KBv200 with IC₅₀ values of less than 0.78 µg/mL [81].

An unusual alkaloid 2-acetyl-1,2,3,4-tetrahydro-β-carboline (**150**), fusamine (**151**), and 3-(1-aminoethylidene)-6-methyl-2H-pyran-2,4(3H)-dione (**152**) (Figure 9), were isolated from the culture broth of *Fusarium incarnatum* (HKI0504), an endophytic fungus of the mangrove plant *Aegiceras corniculatum*. Compound **150** exhibits weak antiproliferative activity against HUVEC and the K-562 cell line with GI₅₀ values of 41.1 and 33.3, respectively. Compounds **151** and **152** exhibit anti-proliferative activity against HUVEC and the GI₅₀ K-562 cell line with GI₅₀ values of 37.3, 37.6, and 41.1, 33.3, respectively. The compounds **150**, **151** and **152** displayed cytotoxic activity against HeLa cells with a CC₅₀ value of 23.8, 23.3, and 23.8 µM, respectively. The standard imatinib exhibited anti-proliferative activity against HUVEC and the GI₅₀ K-562 cell line with a GI₅₀ value of 18.5 and 0.17, respectively, and cytotoxic activity against HeLa cells with a CC₅₀ value of 65.8 µM [82].

Acremonium sp., an endophytic fungus residing inside the leaves of *Sonneratia caseolaris* collected from Hainan, China, was the source of a dimeric anthracene derivative torrubieillin B (**153**) (Figure 9). Compound **153** exhibited potent cytotoxic activity against the cisplatin-sensitive cell lines Cal27, Kyse510, HCC38, A2780, MDA-MB-231 with IC₅₀ values ranging from 0.3 to 1.5 µM, and against the cisplatin-resistant cell lines Cal27, Kyse510, HCC38, A2780, MDA-MB-231 with IC₅₀ values ranging from 0.2 to 2.6 µM. The positive control cisplatin exhibited cytotoxic activity against the tested cell lines with IC₅₀ values in the range of 1.5–38.1 µM [83].

Waol A (**154**), pestalotiopene A (**155**), and cytosporone E (**156**) (Figure 9) were obtained from the endophytic fungus *Acremonium strictum*, isolated from the mangrove tree *Rhizophora apiculata*, collected on the island of CatBa, Vietnam. Compounds **154**–**156** showed moderate cytotoxic activity against human cisplatin-sensitive (IC₅₀ values 27.1, 76.2, and 8.3 µM, respectively) and resistant A2780 cell lines (IC₅₀ values 12.6, 30.1, and 19.0 µM, respectively) [84].

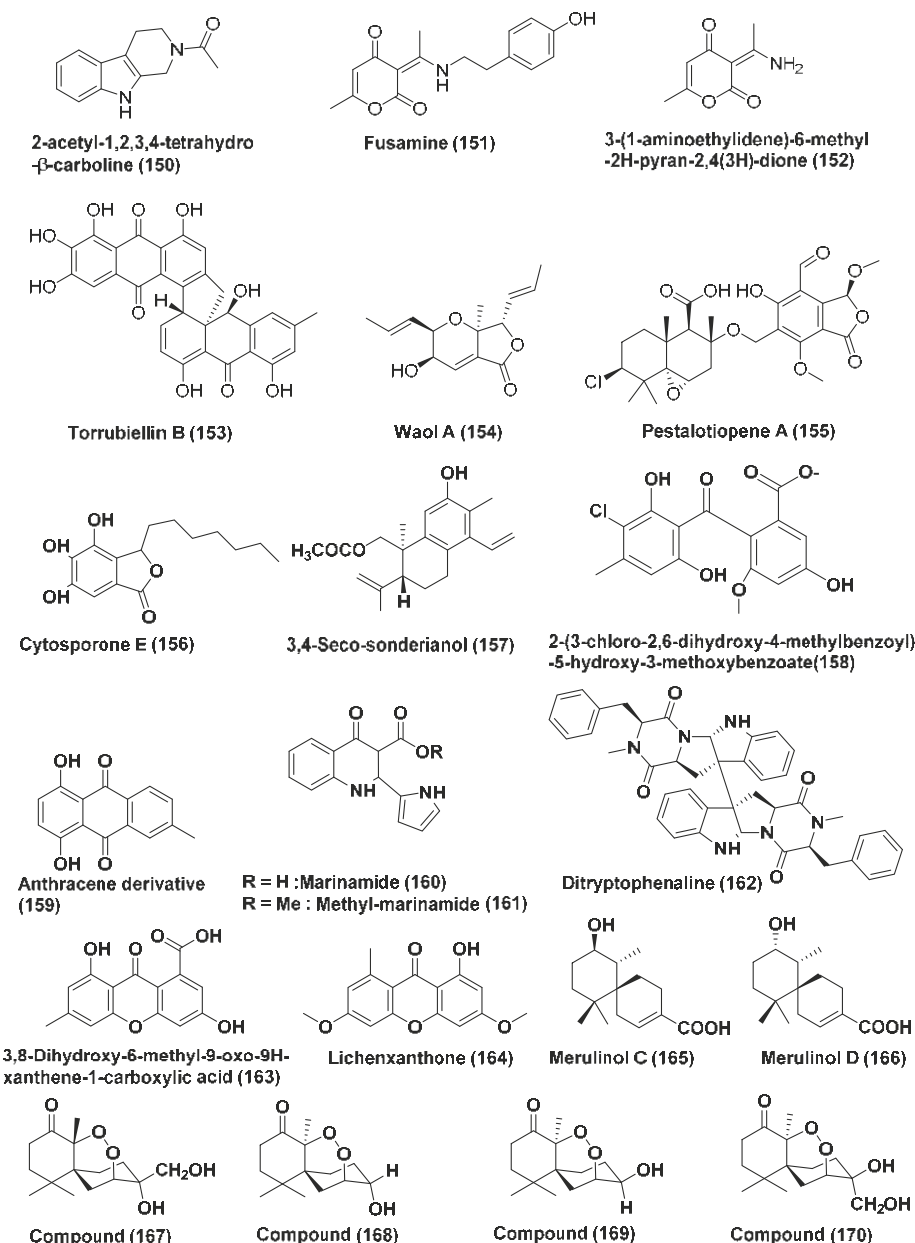


Figure 9. Structures of metabolites isolated from Hyphomycetes (150–170).

The compound 3,4-seco-sonderianol (157) (Figure 9), a known diterpenoid, was obtained from the endophytic fungus J3 of *Ceriops tagal*, collected in Hainan province, China. Compound 157 showed cytotoxic activity against the K562, SGC-7901, and BEL-7402 cell lines with IC₅₀ values of 9.2, 15.7, and 25.4 µg/mL, respectively. The positive control paclitaxel displayed activity against the K562, SGC-7901 and BEL-7402 cell lines with IC₅₀ values of 5.1, 1.6 and 6.3 µg/mL, respectively [85].

The compound 2-(3-chloro-2,6-dihydroxy-4-methylbenzoyl)-5-hydroxy-3-methoxybenzoate (**158**) (Figure 9), was extracted from the endophytic fungus No. ZH-3 from the South China Sea. Compound **158** showed cytotoxic activity against the hepG2 cell line ($IC_{50} = 25 \mu\text{g}/\text{mL}$) [86].

Mangrove endophytic fungus No.5094, which was collected in the South China Sea was the source of anthracene derivative (**159**) (Figure 9). Compound **159** exhibited potent activity towards the KB and KBv200 cell lines with LD_{50} values of 5.5 and $10.2 \mu\text{M}$, respectively [87].

Marinamide (**160**) and methyl marinamide (**161**) (Figure 9) were obtained by co-cultures of two marine-derived mangrove endophytic fungi (strains Nos. 1924 and 3893) from the South China Sea coast. Their structures were elucidated using comprehensive spectra methods. Compound **160** was found to be cytotoxic with IC_{50} values of 7.0, 0.4, 91 nM and $0.529 \mu\text{M}$, respectively against HepG2, 95-D, MGC823 and HeLa cells. For the similar cell lines, compound **161** exhibited cytotoxicity with IC_{50} values of 2.52, 1.54, 13.0 and $0.110 \mu\text{M}$, respectively [88].

Mangrove endophytic fungus No. Gx-3a in the South China Sea was the source of ditryptophenaline (**162**) (Figure 9). Compound **162** exhibited potent cytotoxic activity against KB and KBv200 cells, with LD_{50} values of $8.0 \mu\text{M}$ and $12.0 \mu\text{M}$ [89].

The 3, 8-Dihydroxy-6-methyl-9-oxo-9H-xanthene-1-carboxylate (**163**) and lichenxanthone (**164**) (Figure 9), were isolated from the mangrove endophytic fungus No.SK7RN3G₁ from the South China Sea. Compounds **163–164** exhibited cytotoxicity against the HepG2 cell line with IC_{50} values of 20 and $25 \mu\text{g}/\text{mL}$, respectively [90].

2.4. Compounds Produced by Basidiomycetes

Two new chamigrane sesquiterpenes, merulinols C and D (**165**, **166**) (Figure 9), were isolated from the culture of the basidiomycetous fungus XG8D, isolated from the healthy leaves of *Xylocarpus granatum*, collected in Samutsakorn province, Thailand. Compounds **165** and **166** selectively displayed cytotoxicity against KATO-3 cells with IC_{50} values of 35.0 and $25.3 \mu\text{M}$, respectively [91].

Compounds **167**, **168**, **169** and **170** (Figure 9) were extracted from *Pseudolagarobasidium acaciicola*, associated with *Bruguiera gymnorhiza* from Samut Sakhon province, Thailand. These compounds were evaluated for their cytotoxicity against the HuCCA-1, A549, MOLT-3, HepG2, HL-60, MDA-MB231, T47D, and HeLa cancer cell lines and normal human embryonic lung cell lines (MRC-5). Compound **167** displayed cytotoxicity against the HuCCA-1, A549, MOLT-3, HepG2, HL-60, MDA-MB231, and T47D cancer cell lines with IC_{50} values in the range of 0.28–37.46 μM and against the MRC-5 normal cell line with an IC_{50} value of $17.92 \mu\text{M}$. Compound **167** selectively showed activity against the HL-60 cell line with an IC_{50} value of $0.28 \mu\text{M}$ and a selectivity index (SI) value of 64.0. Compound **168** showed activity against the A549, MOLT-3, HepG2, HL-60, MDA-MB231, T47D, HeLa and MRC-5 cells IC_{50} values in the range of 12.09–170.08 μM . Compounds **169** and **170** were active against the cell lines tested, IC_{50} 15.20–76.97 μM for **169** and IC_{50} 18.31–154.51 μM for **170**. Compound **170** also selectively exhibited cytotoxic activity toward the HL-60 cell line (IC_{50} 18.31 μM) with an SI value of 4.4 [92].

A new nor-chamigrane endoperoxide, 3-*epi*-Stepperoxide A (**171**), along with the known sesquiterpenes steperoxide A (**172**), merulin B (**173**), and merulin C (**174**) (Figure 10) were isolated from *Pseudolagarobasidium acaciicola*, an endophyte residing inside the mangrove plant *Bruguiera gymnorhiza*. Compounds **171** and **172** showed strong cytotoxic activity with IC_{50} ranges of 0.68–3.71 and 0.67 – $5.25 \mu\text{g}/\text{mL}$, respectively, against MOLT-3, HuCCA-1, A549, HepG2, HL-60, MDA-MB-231, T47D, and HeLa cells. Compound **173** exhibited weak activity against MOLT-3, A549, HepG2, HL-60, MDA-MB-231 and T47D cells with an IC_{50} ranging from 11.94–49.08 $\mu\text{g}/\text{mL}$, but was inactive toward HuCCA-1 and HeLa cells at $50 \mu\text{g}/\text{mL}$. Compound **174** showed the most potent cytotoxic activity against HL60 cancer cells, with an IC_{50} value of $0.08 \mu\text{g}/\text{mL}$, whereas it displayed activity toward the MOLT-3, HuCCA-1, A549, HepG2, MDA-MB-231, T47D, and HeLa cell lines with an IC_{50} range of 0.19–3.75 $\mu\text{g}/\text{mL}$ [93].

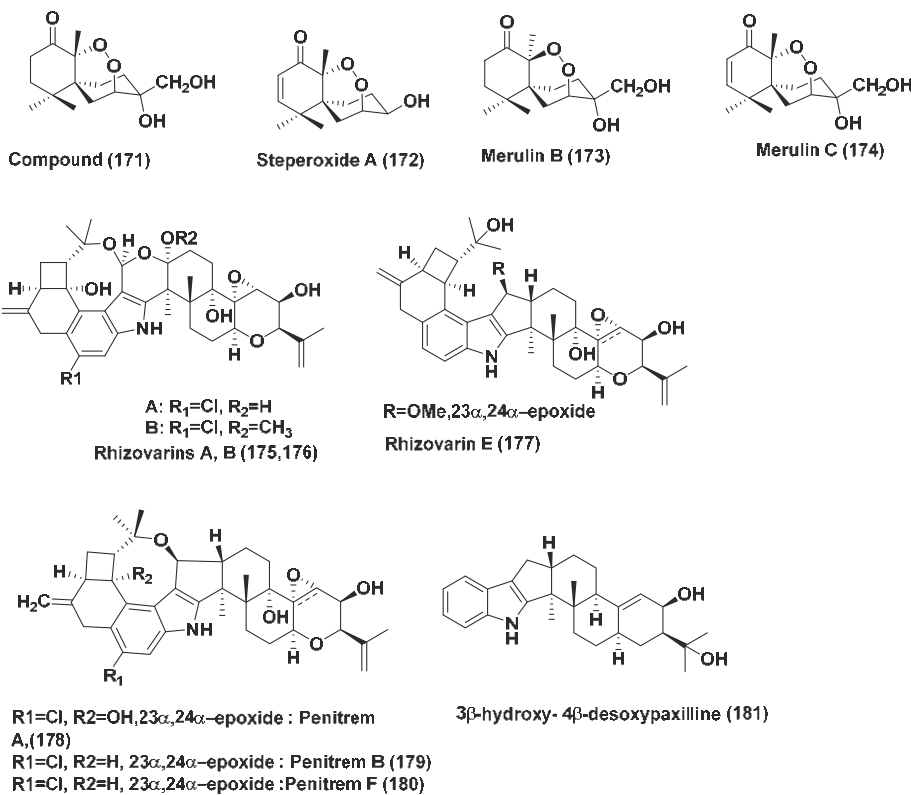


Figure 10. Structures of metabolites isolated from Basidiomycetes (171–174) and from Zygomycetes (175–181).

2.5. Compounds Produced by Zygomycetes

The rhizovarins A, B, E (175–177), penitrem A, C, F (178–180) and 3β-hydroxy-4β-desoxypaxilline (181) (Figure 10) were obtained from *Mucor irregularis* QEN-189, an endophytic fungus residing inside the inner tissue of *Rhizophora stylosa*, collected in Hainan Island, China. Compounds 175–181 showed cytotoxicity against the human A-549 cell lines with IC₅₀ values of 11.5, 6.3, 9.2, 8.4, 8.0, 8.2, and 4.6 μM, while compounds 175, 176, 178–181 were cytotoxic against the human HL-60 cell lines with IC₅₀ values of 9.6, 5.0, 7.0, 4.7, 3.3 and 2.6 μM, respectively. Adriamycin, a positive control, exhibited activity against A-549 and HL-60 cell lines with IC₅₀ values of 0.30 and 0.06 μM, respectively [94].

Table 1. Novel anticancer bioactive compounds reported from mangrove fungi.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by <i>Cochamyces</i>							
1	<i>Pestalotiopsis</i> sp.	<i>Rhizophora mucronata</i>	Not reported	Demethylincisterol A3 (1) Ergosta-5,7,22-trien-3-ol (2), stigmastan-3-one (3), stigmast-4-en-3-one (4), stigmast-4-en-6-ol-3-one (5), flufuran (6)	HeLa, A549 and HepG	In the range of 0.17 to 14.16 nM	[16]
2	<i>Pestalotiopsis microspore</i>	<i>Drepanoecarpus lunatus</i>	Cameroon	Compound (7) Pestalotioprolide D-F (8–10) Pestalotioprolide E (9)	L5178Y A2780	0.7, 5, 6, 34, and 3.9 μ M 1.2 μ M	[17]
3	<i>Pestalotiopsis clavigignora</i>	<i>Rhizophora harrisonii</i>	Port Harcourt, Nigeria	Pestalopolyol (11)	L5178Y	4.10 μ M	[18]
4	<i>Pestalotiopsis viciinii</i>	<i>Kantelia candel</i>	China	Pestalamine A (12)	MCF-7, HeLa, and HepG2	40.3, 22.0, and 32.8 μ M	[19]
5	<i>Phoma</i> sp. OUCMDZ-1847	Fruit sample of <i>Kandelia</i> <i>candel</i>	Wenchang, Hainan Province, China	Phomazines B (13), epicorazine A (14), epicorazine B (15), epicorazine C (16), exserohilone A (17)	HL-60, HCT-116, KS62, MGC-803, and A549 Cells	In the range 0.05 to 8.5 μ M	[20]
6	<i>Phomopsis</i> sp.		Zhanjiang, China	Cytochalasin H (18)	A549 cells	Arrested A549 cells at the G2/M phase, inhibited the migration ability of A549 cells in a dose-dependent manner	[21]
7	<i>Phomopsis</i> spp. xy21 and xy22	Leaves <i>Xylocarpus granatum</i>	Trang Province, Thailand	Phomopsichalasin G (19)	HCT-8, HCT-8/T, A549, MDA-MB-231, and A2780 cancer Cells	7.5, 8.6, 6.4, 3.4, and 7.1 μ M	[22]
8	<i>Phomopsis longicolla</i> HL-2232	<i>Bruguiera sexangula</i> var. <i>rhynchosperma</i>		6-aminopurine-9-carboxylic acid Me ester (20), uridine (21)	B16F10, A549, HL-60 and MCF-7 Cells	14.9 and 8.6 μ M	[23]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by <i>Cochamyces</i>							
9	<i>Phomopsis</i> sp. HNY29-2B	Branch of <i>Acanthus ilicifolius</i>	South China Sea, Hainan Province, China	Dicerandrol A (22), Dicerandrol B (23), Penexanthone A (26)	MDA-MB-435, HCT-116, Calu-3 and Huh7 Cells	3.03, 2.64, 1.76 and 4.19 μ M	[24]
10	<i>Phomopsis</i> sp. (ZH76)	<i>Excoecaria agallocha</i>	Dong Sai, South China Sea coast	Dicerandrol C (24) diacetyl phomoxanthone B (25)	MDA-MB-435, HCT-116, Calu-3, MCF-7, A Cells	<10 μ M, 44.10, 42.63, 36.52, and 33.05 μ M	
11	<i>Diaporthe</i> sp. SCSIO 41011	<i>Rhizophora stylosa</i>	Sanya city, Hainan Province, China	3-O-(6-O- α -L-arabinopyranosyl)- β -D-glucopyranosyl-1,4-dimethoxyxanthone (27)	HEp-2 and HepG2	9 and 16 μ M	[25]
12	<i>Diaporthe phascolorum</i> SKS019	Branches of <i>Acanthus ilicifolius</i>	Shankou, Guangxi province, China	Isochromophilone D (28)	786-O cells	8.9 μ M	[26]
13	<i>Phomopsis</i> sp. A818	Foliage of <i>Kandelia candel</i>	Fujian Province, China	epi-Isochromophilone II (29)	ACHN, OS-RC-2, and 786-O cells,	In the range of 3.0 to 4.4 μ M, Sorafenib (3.4 to 7.0 μ M)	[26]
14	<i>Phomopsis</i> sp. A818	Foliage of <i>Kandelia candel</i>	Fujian Province, China	5-deoxybstrocydien (30)	MDA-MB-435 and NCI-H460 cancer cells	5.32 and 6.57 μ M	[27]
15	<i>Sarcodidium kilense</i> HDN11-84	rhizosphere soil of <i>Thespesia populnea</i> ,	Guangxi Province, China	Fusaristatin A (31)	MDA-MB-435 cancer cells	8.15 μ M	
14	<i>Phomopsis</i> sp. A818	Foliage of <i>Kandelia candel</i>	Fujian Province, China	Mycooxydine (32), deacetylmycooxydine (33)	MDA-MB-435	7.85 and 14.61 μ M	[28]
15	<i>Sarcodidium kilense</i> HDN11-84	rhizosphere soil of <i>Thespesia populnea</i> ,	Guangxi Province, China	Suppress antigen-stimulated degranulation and cytokine production in mast cells and IgE-mediated passive cutaneous anaphylaxis in mice			[29]
15	<i>Sarcodidium kilense</i> HDN11-84	rhizosphere soil of <i>Thespesia populnea</i> ,	Guangxi Province, China	Saroclazine B (34)	HeLa Cells	4.2 μ M	[30]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
16	<i>Annulohypoxylon</i> sp.	<i>Rhizophthora racemosa</i>	Cameroun	Daldinone I (35)	Ramos and Jurkat J16	6.6 and 14.1 μ M, Potently blocks autophagy, a potential pro-survival pathway for cancer cells	[31]
Relaxation activity/The band backward shifting and trailing off rubrunol (36) was observed at 100, 50, 10, 5 and 1 μ M							
17	<i>Eurotium rubrum</i>	<i>Suaeda salsa</i>	"BoHai" seaside, China	Rubrunol (36)	Topo I	A549, MDA-MB-231, PANC-1 and HepG2	[32]
18	<i>Eutypella</i> sp. 1–15	Soil of mangrove rhizosphere in limei, Fujian Province, China	Not reported	13-Hydroxy-3,8(11)- endesmatrien-12,8-olide (37)	JERCO-1 and HepG2	8.4 and 28.5 μ M	[33]
19	<i>Rhytidhysteron</i> <i>nigulum</i> AS21B	Leaves of <i>Azima</i> <i>sambentosa</i>	Samutsakhon province, Thailand	Rhytidones G (38), H (39), deoxypseudohermin B (40), palmarumycin CP17 (41), 1-oxo-1,4-dihydronaphthalene- 4-spiro-20-naphthol[400]-hydroxy- 10,800-de[10,30]-dioxine (42), pseudohermin EG4 (43), rhytidone E (44), rhytidone F (45), palmarumycin C5 (46), and 4,8-dihydroxy-3,4- dihydronaphthalen-1(2 <i>H</i>)-one (47)	Ramos lymphoma Ramos lymphoma	17.98, 0.018, 18.00, 33.1, 15.82, 9.0461, 0.048, 31.7 and 23.1 μ M, (ibrutinib 28.7 μ M)	[34]
Compounds (38), (39), (44), (45), and (47)				H1975 Cell	7.3, 0.252, 10.24, 1.17 and 50 μ M (afatinib 1.97 μ M)		

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
20	<i>Lasiotiplodia</i> sp. 318#	<i>Excoecaria agallocha</i>	Guangdong Province, China	2,4-Dihydroxy-6-nonylbenzoate (48)	MMQ and GH3 Cells	5.2 and 13.0 μ M	[35]
21	<i>Lasiotiplodia</i> sp. 318#	<i>Excoecaria agallocha</i>	Guangdong Province, China	Ethyl-2,4-dihydroxy-6-(80-hydroxynonyl)benzoate (49)	MDA-MB-435, HepG2, HCT-116, A549 and leukaemia THP1 Cells	0.13, 12.50, 11.92, 13.31 and 39.74 μ M	[36]
22	<i>Lasiotiplodia theobromae</i> ZJ-HQ1	<i>Acanthus ilicifolius</i>	Guangdong Province, China	Chloropreussomerins A and B (50, 51), Preussomerin D (56)	A549 and MCF-7	In the range of 5.9–8.9 μ M	[37]
23	<i>Rhytidhysteron rufifolium</i> BG2-Y	Leaves of <i>Bruguiera gymnorhiza</i>	Pak Nam Pran, Prachub Kiri Khan Province, Thailand	Rhytidhormone A (57), B (58), C (59), and E (60)	Kato-3 Cells	16.0–23.3 μ M	[38]
24	<i>Campylocarpon</i> sp. HDN13-307	Root of <i>Sommeria casularis</i>	China	Campyridone D (61), and illicolin H (62)	MCF-7 cells	19.3–17.7 μ M	[39]
25	<i>Stenphytium globaliforme</i>	<i>Apicennia marina</i>	Hurghada, Egypt	Dihydroaltersolanol C (63), Altersolanol A (64), Altersolanol B (65), Alterporriol E (67)	HeLa	8.8 and 4.7 μ M	[39]
26	<i>Paradicthyothrinium diffactum</i> BCC 8704	Associated with mangrove wood	Laem Son National Park, Ranong Province, Thailand	Altersolanol N (66)	L5178Y	3.4, 2.53, 3.78 and 6.9 μ M	[40–42]
Low micromolar range (% growth-1.4)							
				Paradicthyothrin A (68)	KB, MCF-7, NCI-H187, Vero Cells	In the range of 23–31 μ g/mL	
				Paradicthyothrin B (69)	KB, MCF-7, NCI-H187, Vero Cells	3.1, 3.8, 9.5, and 5.6 μ g/mL	[46]
				Preussomerin C (70), ymf 1029C (71) and allenusin (72)	KB, MCF-7, NCI-H187, Vero Cells	Moderate to poor activity	
				ymf 1029C (71)	NCI-H187 cells	5.0 μ g/mL	

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
27	<i>Haloscellinia</i> sp. (No. 1403)	—	South China Sea	SZ-685C (73)	NFPA, MMQ and RfC cells	18.76, 14.51, and 56.09 μ M	[47]
28	<i>Dothiorella</i> sp.	<i>Aegiceras corniculatum</i>	Fujian Province, China	Dothiorelone F (74), Dothiorelone G (75)	Raji cancer	2 μ g/mL	[49]
29	<i>Rhytidites</i> sp.	Leaves of <i>Azima sambentosa</i>	Samut Sakhon province, Thailand	Rhytidones B-C (76, 77), MK3018 (78), Palmarumycin CRI (79)	MCF-7 and CaSki Cells	In the range of 14.47 and 25.59 μ M	[50]
30	<i>Sporothrix</i> sp.	Bark, <i>Kandelia candel</i>	South China Sea	Rhytidones B (76)	CaSki	22.81 μ M	[51]
				Sporothrin A (80)	Inhibition of AChE in vitro	1.05 μ M	[51]
				sporothrin B (81), sporothrin C (82), diaporthin (83)	HepG2	20, 23, and 23 μ g/mL	[51]
31	<i>Eurotium rubrum</i>	Semi-mangrove plant <i>Hibiscus tiliaceus</i>	Hainan Island, China	12-dimethyl-12-oxo-eurotechnidulin B (84), 9-dehydroxyeurotechnone (85), variecolorin G (86), alkaloid E-7 (87), and endodin (88)	HepG2, MCF-7, SW1990, HepG2, NCI-H460, SMMC7721, HeLa, and DU145	In the range of 15–30 μ g/mL	[52]
32	<i>Bionectria ochroleuca</i>	Inner leaf tissues of the plant <i>Sommereria casuarinae</i>	Hainan island, China	Pullularins E (89), F (90), pullularins A (91), and C (92)	L5178Y	EC_{50} values in the range of 0.1 and 6.7 μ g/mL	[53]
				verticilllin D (93)	L5178Y	<0.1 μ g/mL	[53]
33	<i>Aspergillus</i> sp. HN15-5D	Leaves, <i>Acanthus ilicifolius</i>	Hainan Island, China	Pullularins E (89), F (90), pullularins A (91), and C (92)	MDA-MB-435	5.08 and 4.98 μ M	[54]
34	<i>Aspergillus niger</i> MA-132	<i>Avicennia marina</i>	Hainan, China	Nigerasterol A (96) Nigerasterol B (97)	HL60	0.30 μ M, 1.50 μ M	[55]
35	<i>Aspergillus nidulans</i> MA-143	Leaves, <i>Rhizophora stylosa</i>		Nigerasterol A (96) Nigerasterol B (97)	A549	1.82 and 5.41 μ M	
				Aniquinazolines A–D (98–101)	Brine shrimp	LD ₅₀ 1.27, 2.11, 4.95 and 3.42 μ M, Colchicine LD ₅₀ 88.4 μ M	[56]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
36	<i>Aspergillus terreus</i> (No. GX7-3B)	Branch of <i>Bruguiera gymnorhiza</i> (Linn.)	South China Sea	$3\beta,5\alpha$ -dihydroxy- γ (22E,24R)-ergosta-7(22)-dien-6-one (102), Beauvericin (104)	MCF-7, A549, HeLa and KB	4.98 and 2.02, 1.95 and 0.82, 0.68 and 1.14, 1.50 and 1.10 μ M	[57]
37	<i>Aspergillus terreus</i> (No. GX7-3B)	Branch of <i>Bruguiera gymnorhiza</i> (Linn.)	South China Sea	$3\beta,5\alpha,14\alpha$ -trihydroxy- γ (22E,24R)-ergosta-7(22)-dien-6-one (103)	MCF-7, A549, HeLa and KB	25.4, 27.1, 24.4, 19.4 μ M	
38	<i>Penicillium</i> sp. J-54	Leaves, <i>Ceriops tagal</i>	Hainan province, China	Botryosphaerin F (105) and LL-Z1271 β (106)	MCF-7 and HL-60	4.49 and 3.43 μ M	[58]
39	<i>Penicillium citrinum</i> HL-5126	<i>Bruguiera sexangula</i> var. <i>rhynchosperata</i>	South China Sea	Penibenzophenone B (108)	K-562	0.6 μ M (paclitaxel, 9.5 μ M)	[59]
40	<i>Penicillium</i> sp.	<i>Panax notoginseng</i>	Wenshan, Yunnan province, China	Brefeldin A (109), Brefeldin A 7-O-acetate (110)	A549 Cells	90.1 μ M, (paclitaxel, 9.5 μ M)	[60]
41	<i>Penicillium chrysogenum</i> V11	Vein of <i>Myoporum boninoides</i>	Leizhou Peninsula, China	Penochalasin K (111)	MDA-MB-435, SGC-7901 and A549 cells	LD_{50} value from 0.024 to 0.62 μ M. Both the compounds arrested HepG2 cells at the S phase	[61]
42	<i>Penicillium janthinellum</i> HDN13-309	<i>Somneria caseolaris</i>	Hainan Province, China	Penicisulfuronol A-C (112–114)	HeLa and HL-60 Cells	<10 μ M	[62]
43	<i>Penicillium chrysogenum</i> V11	Not reported	Not reported	Penochalasin I (115), chaetoglobosins A (116), and cytoglobosin C (117) Compounds (116), and (117)	MDA-MB-435 and SGC-7901 cells	<10 μ M	[63]
44	<i>Penicillium brocae</i> MA-231	Mangrove plant <i>Avicennia marina</i>	Hainan Island, China	Spirobrocazine C (118)	SGC-7901 and A549 cells	<10 μ M	[64]
					A2780	59 μ M	
					A2780 and A2780 CisR	664 nM, 661 nM (cisplatin 1.67 and 12.63 μ M)	[65]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
45	<i>Penicillium brocae</i> MA-231	Mangrove plant <i>Avicennia marina</i>	Hainan Island, China	Procaaines A (120), B (121), E (122), F (123) Compounds (120) and (121) Compound (123)	Dul45, HeLa, HepG2, MCF-7, NCI-H460, SC-C-7901, SW1990, SW480, and U251 SW480 tumor cell line DU145 and NCI-H460 Cells, TMC-264 (124)	from 0.89 to 9.0 μ M 2.0 and 1.2 μ M 1.7 and 0.89 μ M 1.08 and 2.81 μ M (doxorubicin 1.55 and 2.24 μ M)	[66]
Compounds Produced by Basidiomycetes							
46	<i>Penicillium chermesinum</i> strain HLt-ROR2	<i>Heritiera littoralis</i> ,	Samut Sakhon province, Thailand	PR-toxin (125)	HuCCA-1, HeLa, T47D, and MDA-MB231	3.27 μ M (Etoposide, 35.66 μ M)	[67]
47	<i>Penicillium chrysogenum</i> HND11-24	The rhizosphere soil of the mangrove plant <i>Acanthus ilicifolius</i>	China	Penicitols A (126) Penicitols B (127) Penicitols C (128) and Penixanacid A (129)	MOLT-3 T47D HuCCA-1, HeLa, T47D, and MDA-MB231 HL-60 cell line MOLT-3 and HL-60	1.36 μ M 1.08 μ M 0.81–2.19 μ M (doxorubicin, 0.26–2.24 μ M) 0.06 μ M (doxorubicin, 1.21 μ M) 0.09 μ M, 0.06 μ M	[67]
48	<i>Penicillium</i> sp. Ff-1	<i>Avicennia marina</i>	Fujian, China	Compound (130) Compounds (131) Compounds (131)	Tca8113 and MG-63 cells Tca8113 and WRL-68 MG-63 cells	26 and 35 μ M (Taxol, 46 and 10 mM) 10 and 58 μ M 55 nM	[69]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
49	<i>Penicillium</i> sp. GD6,	<i>Bruguiera gymnorhiza</i>	Zhanjiang, China	Meleagrin (132)	HL60 and A549	9.7 and 8.3 μ M	[70]
50	<i>Penicillium</i> 303#	Sea water	Guangdong Province, China	55, 7R, 9S, 10S, 11R, 12S, 13R, 22R, and 23R, (133), 7R, 9S, 10S, 11R, 12S, 13R, 22R, and 23R (134)	MDA-MB-435, HepG2, HCT-116, and A549	In the range of 11.9–37.82 μ g/mL	[71]
51	<i>Penicillium</i> sp. ZH58	Leaves, <i>Avicennia</i> sp.	Dong Sai, Hainan of the South China Sea coast	Compounds (135)	MDA-MB-435	7.13	
				Compound (136)	HepG2 and HCT-116	39.64 and 27.80 μ M	
52	<i>Penicillium stumattense</i> MA-92	Rhizosphere, <i>Lumnitzera racemose</i>	Wen Chang in Hainan Island, China	4-(methoxymethyl)-7-methoxy-6-methyl-1(3H)-isobenzofuranone (137)	KB and KBv200 cells	6 and 10 μ g/mL	[72]
53	<i>Penicillium</i> sp. ZH16	<i>Avicennia</i> sp.	South China Sea	Sumalatins A–C (138, 139, 140), and dehydrocurvularin (141)	Dul45, HeLa, Huh7, MCF-7, NCI-H460, SGC-7901, and SW1990 Cells	In the range of 3.8 to 10 μ M	[73]
54	<i>Trichoderma</i> sp. 307			5-methyl-8-(3-methylbut-2-enyl)furanocoumarin (142)	KB and KBv200	5 and 10 μ g/mL	[74]
	Co cultured with <i>Ametobacter johnsonii</i>	Stem bark, <i>Clerodendrum inerme</i>	Guangdong Province, China	(3S)-6-oxo-de-O-methylasiodiplodin (143) (3R)-de-O-methylasiodiplodin (144)	GH3 and MMQ Cells RPC	21.42 and 13.59 μ M, 142.8 μ M, 6.44 and 6.58 μ M, 6.94 μ M	[75]
	B2			(3R)-nordinone (145)		12.33 and 10.13 μ M, 100.03 μ M,	
55	<i>Trichoderma</i> sp. 307 co-culturing with <i>Ametobacter johnsonii</i>	Stem bark of <i>Clerodendrum inerme</i>	Guangdong Province, China	Botryosphaeric acid H (146)	MMQ GH3 Cells	3.09 and 3.64 μ M	[76]
56	<i>Trichoderma</i> sp. XY24	Leaves, stems and peels of <i>Xylecarpus granatum</i>	Hainan province, China	(9R,10R)-dihydro-harzianone (147)	HeLa and MCF-7 Cells	30.1 μ M and 30.7 μ M	[77]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
57	<i>Nigrospora</i> sp. MA75	<i>Pongamia pinnata</i>	Guangxi Zhuang Autonomous Region of China	2,3-didehydro-19 α -hydroxy-14 β -epicichloquinone B (148)	MCF-7, SW1990, and SMMC721	4, 5, and 7 μ g/mL	[78]
58	<i>Fusarium</i> sp. (No. DZ27)	Bark of <i>Kandelia candel</i>	Dongzhai mangrove, Hainan, China	Beauvericin (104)	KB and KBv200 cells	5.76 and 5.34 μ M	[79]
59	<i>Fusarium</i> sp.	Leaf of mangrove <i>Kandelia candel</i>	Dongzhai Harbor of Hainan Island, China	Apicidin (149)	GLC-82 cells	6.94 μ M	[80]
60	Unidentified fungus ZZF42	South China Sea	Not reported	Apicidin (149)	KB and KBv200	0.78 μ g/mL	[81]
61	<i>Fusarium incarnatum</i> (HKU0504)	<i>Aegiceras corniculatum</i>	Not reported	2-acetyl-1,2,3,4-tetrahydro- β -carboline (150)	HUVEC and K-562 HeLa cell	IC_{50} 41.1 and 33.3 μ M	[82]
	<i>Fusarium incarnatum</i> (HKU0504)	<i>Aegiceras corniculatum</i>	Not reported	Fusamine (151)	HUVEC and K-562 HeLa cell	IC_{50} 37.3 and 37.6 μ M	[82]
	<i>Fusarium incarnatum</i> (HKU0504)	<i>Aegiceras corniculatum</i>	Not reported	3-(1-aminoethylidene)-6-methyl-2H-pyran-2,4(3H)-dione (152)	HUVEC and K-562 HeLa cell	IC_{50} 41.1 and 33.3 μ M	[82]
62	<i>Acremonium</i> sp.	leaves of <i>Sonneratia caseolaris</i> c	Dong Zhai Gang Mangrove Garden, Hainan, China	Tortriubellin B (153)	Cisplatin sensitive Cal27, Kysse510, HCC38, A2780, MDA-MB231 Cisplatin resistant, Cal27, Kysse510, HCC38, A2780, MDA-MB231	In the range of 0.3 to 1.5 μ M In the range of 0.2 to 2.6 μ M	[83]
63	<i>Acremonium strictum</i>	<i>Rhizophora apiculata</i>	Island of Cat Ba, Vietnam	Wao A (154), Pestalotiopene A (155) Cytopsporone E (156)	Cisplatin-sensitive, A2780 Cisplatin-Resistant A2780	27.1, 76.2, and 8.3 μ M 12.6, 30.1, and 19.0 μ M	[84]
64	Endophytic fungus J3	<i>Ceriops tagal</i>	Hainan Province, China	3,4-seco-sonderianol (157)	K562, SGC-7901, and BEI-7402 Cells	9.2, 15.7, and 25.4 μ g/ml	[85]
65	Endophytic fungus No. ZH-3	Not reported	South China Sea	2-(3-chloro-2,5-dihydroxy-4-methylbenzoyl)-3-methoxybenzoate (158)	HepG2 cell line	25 μ g/mL	[86]

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Ascomycetes							
66	Endophytic fungus No. 5094	Not reported	South China Sea	Anthracene derivative (159)	KB and KBv200	LD ₅₀ values of 5.5 and 10.2 μ M	[87]
67	Co-cultures of two mangrove endophytic fungi (strains Nos. T324 and 3893)	Not reported		Marinamide (160)	HepG2, 95-D, MGCC832 and HeLa Cells	7.0, 0.4, 91 nM and 0.529 μ M	[88]
68	Endophytic fungus No.Gx-3a	Not reported	South China sea	Methyl marinamide (161)	HepG2, 95-D, MGCC832 and HeLa Cells	2.52, 1.54, 13, 0.110 μ M	[88]
69	Mangrove endophytic fungus No-SK7RN3G1	Not reported	South China Sea	Dityryptophenoline (162)	KB, KBv200	8.0 and 12.0 μ M	[89]
70	Basidiomycetous fungus XG8D	leaves of <i>Xyllocarpus granatum</i>	Samut Sakhon province, Thailand	3,8-dihydroxy-6-n-methyl-9-oxo- β -xanthene-1-carboxylate (163), Lichenanthrone (164), Merulinols C and D (165, 166)	HepG2 cell line	20 and 25 μ g/mL	[90]
Compounds Produced by Basidiomycetes							
71	<i>Pseudolagrobasidium acacicola</i> , <i>Bruguiera gymnorhiza</i>		Samut Sakhon province, Thailand	Compound (167)	HuCCA-1, A549, MOLT-3, HepG2, MDA-MB231, T47D	0.28–37.46 μ M	[91]
				Compound (168)	MRC-5	IC ₅₀ 17.92 μ M	
					HL-60 cell line	IC ₅₀ 0.28 μ M	
				Compound (169)	A549, MOLT-3, HepG2, HL-60, MDA-MB231, T47D, HeLa cancer cell	12.09–170.08 μ M	[92]
				Compound (170)	HuCCA-1, A549, MOLT-3, HepG2, HL-60, MD-A-MB231, T47D, HeLa cancer cell	18.31–154.51 μ M	
					HL-60	18.31 μ M	

Table 1. Cont.

Sr. No.	Fungus	Host Plant(s)	Plant Part or Tissue Locality of Host Plants	Compounds Isolated	Cell Line	IC_{50}/EC_{50} /Inhibition	Refs.
Compounds Produced by Basidiomycetes							
72	<i>Pseudolagarbasidium acacicola</i>	<i>Brugiera gymnorhiza</i>	Not reported	Endoperoxide (171), Steperoxide A (172)	MOLT-3, HuCCA-1, A549, HepG2, HL-60, MDA-MB-231, T47D, and HeLa cancer Cells	In the range of 0.68–3.71 and 0.67–5.25 μ g/mL	[93]
Rhizovariins A, B, E (175, 176, 177), Penitrem A, C, F (178, 179, 180) and 3 β -hydroxy- α - β -desoxypaxilline (181)							
73	<i>Mucor irregularis</i> QEN-189	<i>Rhizophora stylosa</i>	Hainan Island, China	Penitrem A, C, F (178, 179, 180) and 3 β -hydroxy- α - β -desoxypaxilline (181)	MOLT-3, A549, HepG2, HL-60, MDA-MB-231 and T47D Cells	In the range of 11.94–49.08 μ g/mL	[94]
Rhizovariins A, B, E (175, 176, 177), Penitrem A, C, F (178, 179, 180) and 3 β -hydroxy- α - β -desoxypaxilline (181)							
					MOLT-3, HuCCA-1, A549, HepG2, MDA-MB-231, T47D, and HeLa Cells	In the range of 0.19–3.75 μ g/mL	

3. Methods Used for the Activation of Silent Biosynthetic Genes

Recent studies in the marine-based microorganisms have shown that these microorganisms are a rich source for novel bioactive compounds. Salinosporamide A (marizomib), a microbial compound isolated from marine *Salinispora* bacteria with proteasome inhibitory activity is expected to be a future anti-cancer drug, and is presently under clinical trials [95]. However, the reoccurrence of the same compound as discovered in terrestrial sources, in marine microorganisms often leads to serious issues. Advances in molecular biology have enhanced our understanding regarding how to exploit the genetic potential of bacteria and fungi to produce newer chemical entities apart from those that are currently known, which have yet to be explored [96,97]. It has been reported that under laboratory conditions, biosynthetic genes are not expressed as such, as only limited bioactive compounds are produced by these microbes. To overcome these limitations, different strategies have been proposed, including culturing promising strains in varying culture media and under a variety of culture conditions [98], mixing cultures of two or more microbe variants and epigenetic modifications that treat microbes with epigenetic modifiers such as histone deacetylase inhibitors or DNA methyl transferase to initiate the transcription of silent genes [99,100] to enhance the variation and diversity of the produced metabolites.

3.1. The Co-Culture Strategy

Microbes in natural ecosystem conditions always harbor and flourish in co-existence with a variety of microbes. Antagonism and competition for limited resources often lead to high competition among species, and microbes adopt various defense strategies, which favor the production of important bioactive secondary metabolites [101]. The co-culturing of two or more different microbes at the laboratory scale might mimic the ecological setting and induce the cascade of genes responsible for biosynthesis that are normally are masked under optimum culture parameters. Co-cultivation of two *Aspergillus* species derived from mangroves produced the new alkaloid asperginic acid and the previously recognized compounds neoaspergillic acid and ergosterol, with antibacterial activity [102]. Li et al. [103] co-cultured two mangrove epiphytes and identified a novel xanthone derivative compound that showed antifungal activity. Two new alkaloids, marinamide, and marinamide methylether, were reported from mangrove-derived endophytic fungi with a cytotoxic effect when grown in mixed fermentation [88]. Pestalone, a chlorinated prenylsecocanthraquinone, was produced by the marine-derived fungus *Pestalotia* sp. when grown in the presence of the marine-derived bacterium *Thalassopia* sp., which belongs to the Gram-negative group. [104]. When *Libertella* sp., a marine-based fungi, were cultured in the presence of the bacteria *Thalassopia* sp., it resulted in the production of diterpenoid libertellenones of fungal origin [105]. In another set of studies, when the bacterium *Sphingomonas* sp. was grown in the presence of *Aspergillus fumigatus*, a novel compound glionitrin A, a diketopiperazine disulfide, was identified and appeared to show strong cytotoxicity against HCT-116, A549, AGS and DU145 cells [106]. These studies suggest that co-cultivation has tremendous potential to generate novel chemical entities from microbes when cultured under laboratory conditions.

3.2. Epigenetic Modification

The addition of epigenetic modifiers to fungi would allow us to induce cryptic fungal gene clusters. This technique can be applied to any fungal strain and does not require strain-dependent genetic manipulation. Williams et al. [107] reported that epigenetic modifiers could be rationally employed to access silent natural product pathways. Histone deacetylase (HDAC) or DNA methyltransferase (DMAT) are often used as epigenetic agents to change the transcription rate of some genes [108]. Henrikson et al. [12] reported the identification of nygerone A from *A. niger* when grown with suberoylanilide hydroxamic acid (SAHA). Wang et al. [109] reported induced metabolite generation in *Penicillium citreonigrum* when grown in the presence of methyl transferase inhibitor, 5-azacytidine (5-AZA). When *Hypoxyylon* sp., an endophytic fungi, was treated with the epigenetic modifiers SAHA and AZA it enhanced the production of volatile organic compounds (VOCs) [110].

The marine endophytic fungus *Leucostoma persoonii* from *Rhizophora mangle* enhanced the production of cytosporones B, C, E and R in HDAC inhibited fermentation [111]. These studies provide evidence that the use of epigenetic modifiers modulate secondary metabolite production, resulting in different gene expressions.

4. Conclusions

Mangrove fungi are a ubiquitous source of novel bioactive metabolites with the potential to display anticancer properties. It is interesting to observe the chemical diversity in these metabolites, which include simple glycoside (27) and peptide molecules (pullularins E, 89; F, 90 and apicidin, 149) as well as complex stereospecific structures such as cytochalasin H (18), phomopsichalasin G (19), aniquinazolines A–D (98–101) and penitrem A, B and F (178–180). Chemical diversity plays an important role in the drug discovery pipeline, as this provides structurally diverse scaffolds that display similar activity via different modes and/or mechanisms of action. This phenomenon is also observed in mangrove fungal metabolites, as they show potent anticancer activity via different mechanisms of action such as apoptotic cell death (SZ-685C, 73; beauvericin, 104), the inhibition of kinase proteins involved in signal transduction pathways (Mycoepoxydiene, 32; Altersolanol A, 64; and the inhibition of topoisomerase I (36). Although many metabolites demonstrated moderate cytotoxic activities against cancer cell lines, only a few displayed superior activity than the standard anticancer drugs (98–101, 119, 124). It can be suggested that the rational derivatization of metabolites may provide molecules with better activity against a wide range of cancer cell lines. In addition, the identified metabolites with broad-spectrum anticancer activity need to be investigated to establish their mechanisms of action and to develop as novel anticancer therapeutics.

Author Contributions: The manuscript was critically evaluated by S.K.D., V.P., M.S.R. and M.K.G. The chemical structures were drawn by V.P. and M.K.G. and they assisted in the preparation of Table 1. The manuscript has been read and approved by all named authors.

Funding: This research received no external funding.

Acknowledgments: The authors are thankful to Alok Adholeya, Sustainable Agriculture Division, The Energy and Resources Institute (India) for continuous support.

Conflicts of Interest: The authors declare no conflict of interest.

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Review

Endophytic Fungi: A Source of Potential Antifungal Compounds

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Received: 20 May 2018; Accepted: 16 June 2018; Published: 25 June 2018

Abstract: The emerging and reemerging forms of fungal infections encountered in the course of allogeneic bone marrow transplants, cancer therapy, and organ transplants have necessitated the discovery of antifungal compounds with enhanced efficacy and better compatibility. A very limited number of antifungal compounds are in practice against the various forms of topical and systemic fungal infections. The trends of new antifungals being introduced into the market have remained insignificant while resistance towards the introduced drug has apparently increased, specifically in patients undergoing long-term treatment. Considering the immense potential of natural microbial products for the isolation and screening of novel antibiotics for different pharmaceutical applications as an alternative source has remained largely unexplored. Endophytes are one such microbial community that resides inside all plants without showing any symptoms with the promise of producing diverse bioactive molecules and novel metabolites which have application in medicine, agriculture, and industrial set ups. This review substantially covers the antifungal compounds, including volatile organic compounds, isolated from fungal endophytes of medicinal plants during 2013–2018. Some of the methods for the activation of silent biosynthetic genes are also covered. As such, the compounds described here possess diverse configurations which can be a step towards the development of new antifungal agents directly or precursor molecules after the required modification.

Keywords: endophytic fungi; antifungal compounds; medicinal plants; co-culture; epigenetic modification

1. Introduction

The undisputed potential of fungi to produce bioactive secondary metabolites has long been established. To date, merely 5% of the entire number of species has been elaborated of an estimated 1.5 million. Out of these (69,000 species), merely 16% (11,500) have been cultured [1]. Amongst the fungi, endophytes represent a wide source of unexplored and uncharacterized microorganisms capable of producing novel metabolites. Endophytes generally exist asymptotically, coexisting with their hosts and representing an underutilized group of microorganisms for the discovery of new compounds. Endophytes produce diverse metabolites and have the ability to synthesize compounds which are solely produced and isolated from higher plants [2,3]. Strobel and Daisy [4] commented that endophytes could be a goldmine of secondary metabolites. *Pestalotiopsis* sp. can be considered as “the *E. coli* of the rain forests” and *P. microspora*, a “microbial factory” of bioactive secondary metabolites. As per them, numerous chemical structures such as Ambuic acid, Cryptocandin, Taxol, Torreyanic acid, Subglutinol A

and B, and many others have been identified. An array of metabolites of different chemical classes profiles have been deciphered, such as alkaloids, cytochalasines, flavonoids, furandiones, phenylpropanoids, lignans, peptides, phenol, phenolic acids, steroids, terpenoids, quinones, aliphatic acid, and chlorinated compounds. Secondary metabolites derived from endophytes comprise classes of compounds such as steroids, xanthones, phenols, isocoumarins, perylene derivatives, quinines, furandiones, terpenoids, depsipeptides, and cytochalasins, which are identified to possess biological activities with antibiotic, antiviral, volatile antibiotic, anticancer, antioxidant, insecticidal, antidiabetic, and immunosuppressive properties [5–9]. Endophytes play a major role in the physiological activities of host plants, influencing the enhancement of stress, insects, nematodes, and disease resistance [10–13].

This review covers the antifungal fungal metabolites reported from endophytic fungi from medicinal plants during 2013–2018 and their potential as antifungal agents. The antifungal activity of these compounds against the selected fungal pathogens are described briefly and some details such as producing organisms, plant sources, place of collections, and the antifungal properties of many of these compounds are shown in Tables 1 and 2.

2. Medicinal Plants

2.1. Compounds Produced by Coelomycetes

Pestalotiopsis is an important genus of coelomycetes and different species of this genus have been identified for production of bioactive compounds for various biological properties which include antimicrobial, antifungal, antiviral, antineoplastic, and antioxidant activities [14]. Some of the antifungal reported from this genus includes a new monoterpane lactone, (3R,4R,6R,7S)-7-hydroxyl-3,7-dimethyl-oxabicyclo[3.3.1]nonan-2-one (1) (Figure 1), along with one related known compound, (3R,4R)-3-(7-methylcyclohexenyl)-propanoic acid (2) (Figure 1), were discovered from endophytic fungus *Pestalotiopsis foedan* obtained from the branch of *Bruguiera sexangula* in Hainan, China. Compounds 1–2 showed antifungal activity against *Botrytis cinerea* and *Phytophthora nicotianae* with MIC values of 3.1 and 6.3 µg/mL, respectively, while the known antifungal drug ketoconazole showed comparable activity (MIC 3.1 µg/mL each) Compound 2 also exhibited satisfactory activity against *Candida albicans* (MIC value of 50 µg/mL) while ketoconazole showed MIC of 6.3 µg/mL [15].

Plants of *Dendrobium officinale* were collected in Yandang Mountain, Zhejiang Province, China. *Pestalotiopsis* sp. DO14 was obtained from the shoots of *D. officinale* endophytic fungus which yielded two novel antifungal constituents, (4S,6S)-6-[(1S,2R)-1,2-dihydroxybutyl]-4-hydroxy-4-methoxytetrahydro-2H-pyran-2-one (3) and (6S,2E)-6-hydroxy-3-methoxy-5-oxodec-2-enoic acid (4), and two known compounds, LL-P880γ (5) and LL-P880α (6) (Figure 1) were isolated. Compounds 3–6 exhibited good anti-fungal activities (MIC ≤ 50 µg/mL) against *C. albicans*, *Cryptococcus neoformans*, *Trichophyton rubrum*, and *Aspergillus fumigatus*. Compounds 3 and 4 possess the strong activities with the MIC values ≤ 25 µg/mL against tested strains [16].

Endophytic fungus *Pestalotiopsis fici* obtained from the branches of *Camellia sinensis* collected from the suburb of Hangzhou, China was the source of a new a-pyrone derivative ficipyrone A (7) (Figure 1). Compound 7 showed antifungal activity against the plant pathogen *Gibberella zeae* with an IC₅₀ value of 15.9 µM (the positive control ketoconazole showed an IC₅₀ value of 6.02 µM) [17].

Endophytic fungus *Pestalotiopsis mangiferae* associated with *Mangifera indica* Linn collected from Maduravoyal, Tamil Nadu Province, India was the source of a new phenolic compound 4-(2,4,7-trioxa-bicyclo[4.1.0]heptan-3-yl) phenol (8) (Figure 1). Compound 8 exhibited strong antifungal activity against *C. albicans* with MIC value of 0.039 µg/mL, while nystatin showed MIC 10.0 µg/mL [18].

Phomopsis is an important genus that is a prolific producer of bioactive compounds including Cytochalasin H (9) (Figure 1), which was isolated from the endophytic fungus *Phomopsis* sp. of *Senna spectabilis* (Fabaceae) collected from São Paulo, Brazil. Compound 9 exhibited activity against *Cladosporium cladosporioides* and *C. sphaerospermum* with MIC values of 10.0 and 25.0 µg, respectively, while the MIC of nystatin the reference compound was 1.0 µg [19].

Endophytic fungus *Phomopsis* sp. isolated from *Aconitum carmichaeli* collected in Huize County, Yunnan Province, China yielded (14 β ,22E)-9,14-dihydroxyergosta-4,7,22-triene-3,6-dione (10) and (5 α ,6 β ,15 β ,22E)-6-ethoxy-5,15-dihydroxyergosta-7,22-dien-3-one (11), calvasterol A (12), and ganodermaside D (13) (Figure 1). All compounds were evaluated for their antifungal activities against *Candida albicans*, *Aspergillus niger*, *Pyricularia oryzae*, *Fusarium avenaceum*, *Hormodendrum compactum*, and *Trichophyton gypseum*. Compound 10 exhibited average antifungal activities against *C. albicans*, *H. compactum*, and *A. niger*, with MIC values of 64, 64, and 128 μ g/mL, respectively. Compound 11 showed poor inhibitory activity against *C. albicans* and *F. avenaceum* with MIC values of 128 μ g/mL. Compounds 12 and 13 showed average inhibitory activities against *F. avenaceum* (MIC 64 μ g/mL for both compounds). Only compound 12 exhibited weak antifungal activities against *P. oryzae* and *T. gypseum* (MIC of 128 and 256 μ g/mL, respectively) [20].

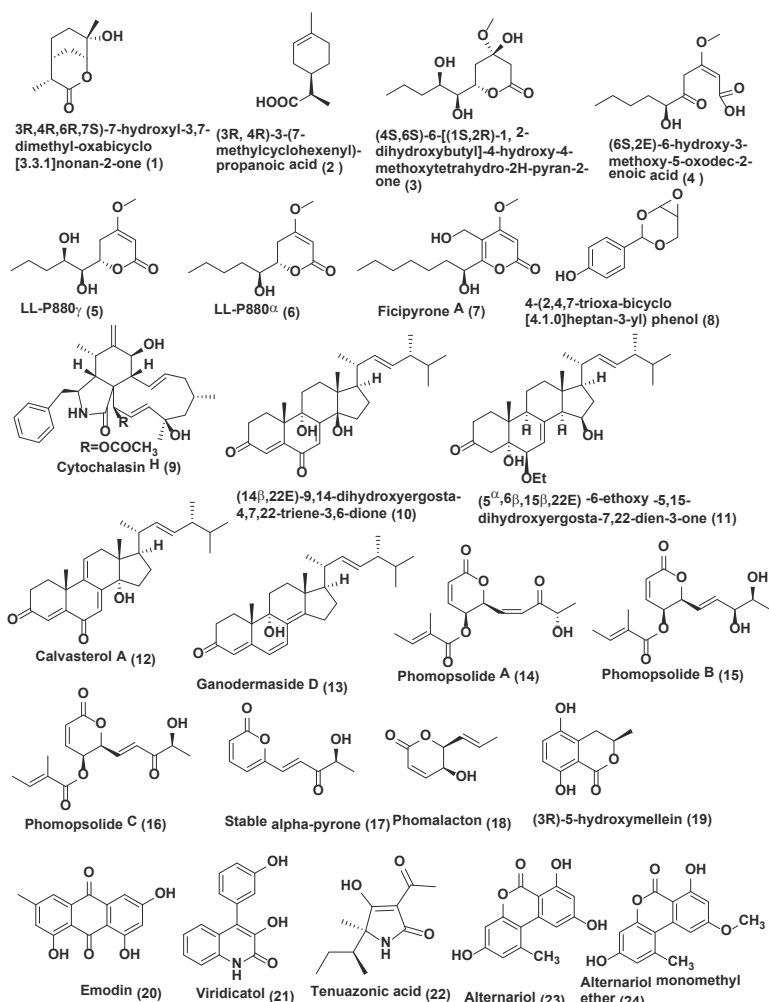


Figure 1. Structures of metabolites isolated from Coelomycetes (1–24).

Diaporthe maritima an endophytic fungus obtained from needles of *Picea* sp. in the Acadian forest of Eastern Canada yielded three dihydropyrone, phomopsolides A (14), B (15), and C (16), and a stable alpha-pyrone (17) (Figure 1). Compound 14 demonstrated growth inhibition at 25 μM against *Microbotryum violaceum* and *Saccharomyces cerevisiae* whereas Compounds 15–17 were active at 250 μM [21].

Another coelomycete *Phoma* is known to produce diverse compounds [22]. From *Phoma* sp. an endophytic fungus of the plant *Fucus serratus* yielded phomalacon (18), (3R)-5-hydroxymellein (19) and emodin (20) (Figure 1). Phomalacone (18), (3R)-5-hydroxymellein (19) and emodin (20) Compounds 18–20 exhibited antifungal activity against *Microbotryum violaceum* with 5, 6 and 5 mm zone of inhibition (0.05 mg was pipetted onto 9 mm sterile filter disk) [23].

Viridicatol (21) (Figure 1), tenuazonic acid (22), alternariol (23), and alternariol monomethyl ether (24) (Figure 1) were isolated from endophytic fungi *Phoma* sp. WF4 of *Eleusine coracana* grown under semi-hydroponic conditions Arkell Field Station, Arkell, ON, Canada. Compounds 21–24 caused reasonable breakage of *Fusarium graminearum* hyphae in vitro [24].

Endophytic fungus *Rhizopycnis vagum* Nitaf 22 obtained from the healthy root of *Nicotiana tabacum* grown at China Agricultural University Beijing, China was the source of Rhizopycnin D (25) and TMC-264 (26) (Figure 2). Compounds 25 and 26 showed strong inhibition of the spore germination of *Magnaporthe oryzae* with IC₅₀ values of 9.9 and 12.0 $\mu\text{g}/\text{mL}$, respectively [25].

A new polychlorinated triphenyl diether named microsphaerol (27) (Figure 2) has been isolated from the endophytic fungus *Microsphaeropsis* sp. (internal strain No. 8883), and phytochemical investigation of the endophytic fungus *Seimatosporium* sp. (internal strain No. 8883) associated with *Salsola oppositifolia* from Playa del Ingles (Gomera, Spain) led to the isolation of a new naphthalene derivative named seimatorone (28) (Figure 2). In antifungal assay, compounds 27 and 28 showed activity against *Microbotryum violaceum* with 9 and 5 mm zone of inhibition (0.05 mg was pipetted onto 9 mm sterile filter paper disk). In addition, there was some growth within the zone of inhibition [26].

Endophytic fungus *Colletotrichum gloeosporioides* associated with *Michelia champaca* isolated from São Paulo State University (UNESP), Araraquara, São Paulo, Brazil was found to be source of a new compound, 2-phenylethyl 1H-indol-3-yl-acetate (29) (Figure 2). Compound 29 displayed good activity against *Cladosporium cladosporioides* and *C. sphaerospermum* which was analogous to nystatin, the positive control [27].

Colletonoic acid (30) (Figure 2) was isolated from *Colletotrichum* sp. from Gomera (Spain). Colletonoic acid exhibit antifungal activity against *Microbotryum violaceum* with 7 mm zone of inhibition (0.05 mg was pipetted onto 9 mm a sterile filter paper disk) [28].

Coniothyrium sp., an endophytic fungus associated with *Salsola oppositifolia* from Gomera in the Canary Islands, was the source of known hydroxy anthraquinones 1,7-dihydroxy3-methyl-9,10-antraquinone (31), 1,6-dihydroxy-3-methyl-9,10-antraquinone (phomarin) (32), and 1-hydroxy-3-hydroxymethyl-9,10-antraquinone (33) (Figure 2) along with four new derivatives having a tetralone moiety, namely coniothyrinones A–C (34–36) (Figure 6) and D (37) (Figure 2). The absolute configurations of coniothyrinones A (34), B (35), and D (37) were determined by TDDFT calculations of CD spectra, allowing the determination of the absolute configuration of coniothyrinone C (36) as well. Coniothyrinones A (34), B (35), and D (37) could be used as ECD reference compounds in the determination of absolute configuration for related tetralone derivates. Compounds 31–37 showed inhibitory effects against the fungus *Microbotryum violaceum* with 7, 10, 8, 7.5, 6, 8 and 7.5 mm zone of inhibition (0.05 mg/9-mm sterile filter paper disk). Compounds 32 and 34 exhibited strong antifungal activity against *M. violaceum* (10 and 9 mm zone of inhibition) and *B. cinerea* (7.5 and 12.5 mm zone of inhibition) when tested under similar conditions [29].

2.2. Compounds Produced by Ascomycetes

Xylaria is very important ascomycetous genus and a good sources of novel bioactive compounds, and some of the compounds reported to have drug-able properties relevant for drug discovery [30,31]. Nine oxygenated guaiane-type sesquiterpenes (38–46) and three isopimarane diterpenes (47–49), (Figure 2)

were obtained from *Xylaria* sp. YM 311647, an endophytic fungus associated with *Azadirachta indica* collected from Yuanjiang County, Yunnan Province, China. All compounds were evaluated for their antifungal activities against *Candida albicans*, *Aspergillus niger*, *Pyricularia oryzae*, *Fusarium avenaceum*, and *Hormodendrum compactum*. Compounds 38–46 were moderately active against *C. albicans* and *H. compactum* (MIC values ranging from 32 to 256 µg/mL), while compound 47–49 were more active against all the tested strains (MIC values ranging from 16 to 256 µg/mL). Compound 49 exhibited the most promising activity against *C. albicans* and *P. oryzae* with MIC values of 16 µg/mL [32].

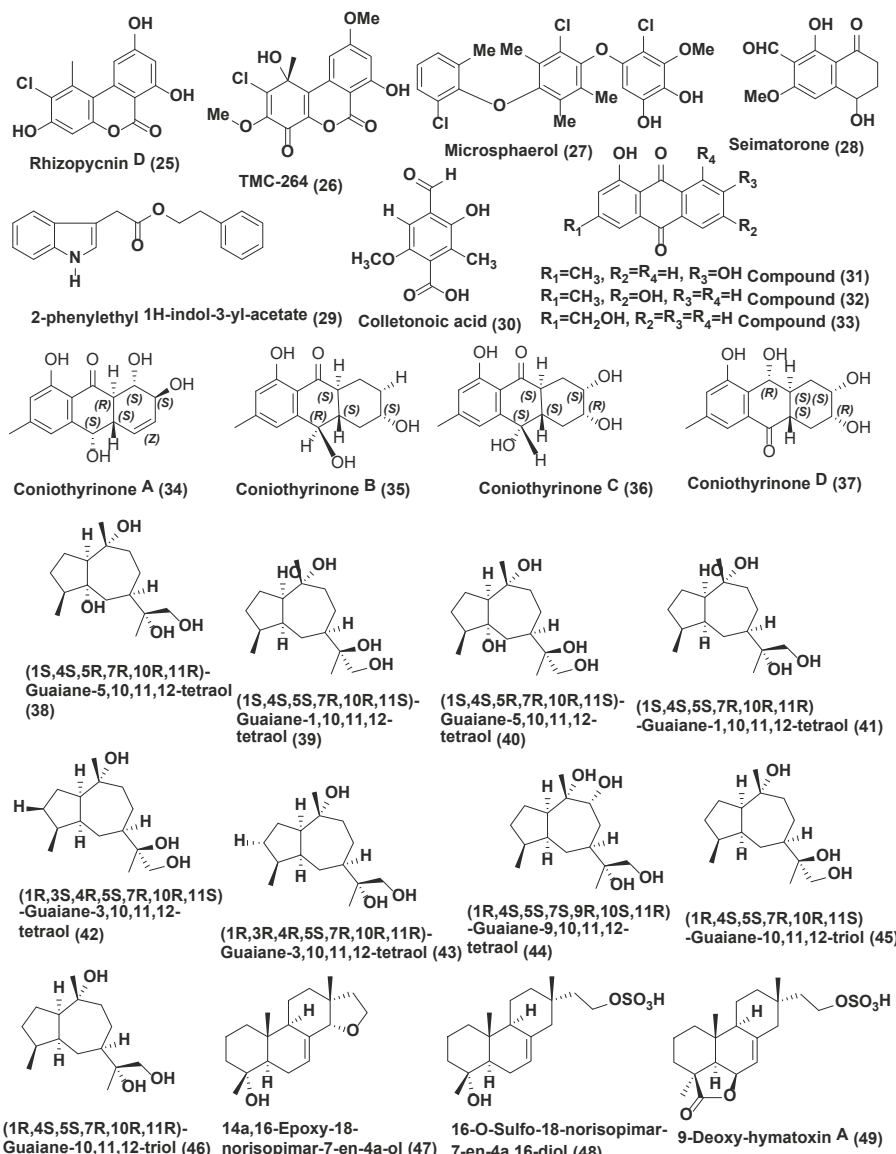


Figure 2. Structures of metabolites isolated from Coelomycetes (25–37) and Ascomycetes (38–49).

Endophytic fungus *Xylaria* sp. YM 311647 associated with *Azadirachta indica* from Yuanjiang County, China was also reported to produce five new guaiane sesquiterpenes, (50–54) (Figure 3). The antifungal activities of 50–54 were evaluated by means of the broth microdilution method against *C. albicans*, *A. niger*, *P. oryzae*, *F. avenaceum* and *H. compactum*. Compounds 50–54 exhibited average or poor antifungal activities against *P. oryzae* and *H. compactum* (MIC values in the range of 32–256 µg/mL). Among them, 53 exhibited the most promising inhibitory activity against *P. oryzae* with a MIC value of 32 µg/mL. Compounds 52 and 53 showed average antifungal activities against *H. compactum* with MIC values of 64 µg/mL. In addition, 53 and 54 exhibited the most promising antifungal activities against *C. albicans* with MIC values of 32 µg/mL. Compound 52 showed average inhibitory activities against *C. albicans*, *A. niger*, and *H. compactum* with MIC values of 64 µg/mL. All compounds showed no notable inhibitory activities against *Fusarium avenaceum* [33]. Amazonian endophytic fungus *X. fejeensis* residing in *Croton lechleri* yielded nonenolide, xyolide (55) (Figure 3). Compound 55 exhibited antifungal activity against oomycetes *Pythium ultimum* with a MIC value of 425 µM [34].

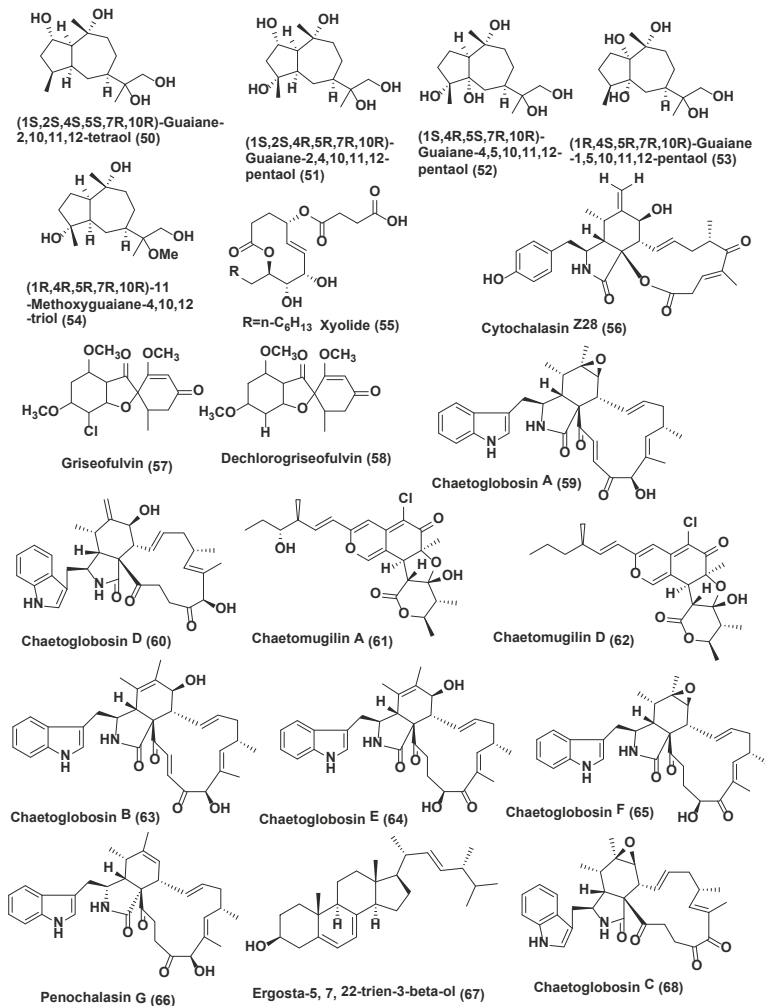


Figure 3. Structures of metabolites isolated from Ascomycetes (50–68).

Endophytic fungus *Xylaria* sp. XC-16 associated with *Toona sinensis* was isolated from Yangling, Shaanxi Province, China and was observed to produce a potent antifungal compound Cytochalasin Z28 (56) (Figure 3), displaying enhanced activity with an MIC of 12.5 μM as opposed to the antifungal activity possessed by hymexazol possessing an MIC value of 25 μM against the plant pathogen *Gibberella saubinetii* [35].

Various isolates of *Xylaria* produce griseofulvin (57) (Figure 3); *Xylaria* sp. PSU G12 associated with *Garcinia hombroniana* [36], and *X. cubiensis* residing in *Asimina triloba* [37], along with 13 strains of *Xylaria* sp. inhabiting *Pinus strobus* and six strains associated with *Vaccinium augustifolium* found in the Acadian forest of New Brunswick and Nova Scotia Canada [38] are known to produce griseofulvin and a few of them can also produce dechlorogriseofulvin (58) (Figure 3) [36,38].

Griseofulvin (57) is very potent against the phytopathogenic fungi, but not against oomycetes [38]. Griseofulvin has been validated to possess antifungal activity against *Alternaria mali*, *B. cinerea*, *C. gloeosporioides*, *Corticium sasaki*, *Fusarium oxysporum* and *Magnaporthe grisea* in vitro with IC₅₀ values of 18.0, 5.0, 1.7, 11.0, 30.0, and 1.7 $\mu\text{g}/\text{mL}$, respectively. Dechlorogriseofulvin (58) demonstrated poor activity, with an IC₅₀ value of 200 $\mu\text{g}/\text{mL}$ for each fungus. Griseofulvin (57) also hinders the growth of *M. grisea*, *C. sasaki*, *B. cinerea*, *Puccinia recondite* and *Blumeria graminis* f. sp. *hordei* in vivo, with a percentage of fungal control of 95, 100, 60, 90 and 90, respectively, at 150 $\mu\text{g}/\text{mL}$. Griseofulvin (57) is used to cure dermatophytic infections caused by fungi such as *Epidermophyton* and *Trichophyton* species [38].

Chaetomium is another genus of ascomycete and prolific producer of bioactive compounds [39,40]. Chaetoglobosin A (59) and D (60) (Figure 3) were isolated *Chaetomium globosum* CDW7, an endophyte from *Ginkgo biloba* located in Taixing and Nanjing in Jiangsu Province and Chengdu in Sichuan Province, China. Compounds 59 and 60 showed antifungal activity against *Sclerotinia sclerotiorum* with IC₅₀ values of 0.35 and 0.62 $\mu\text{g}/\text{mL}$, respectively, compared with carbendazim (0.17 $\mu\text{g}/\text{mL}$) [41].

Compounds Chaetomugilin A (61), Chaetomugilin D (62) Chaetoglobosin A (59), Chaetoglobosin B (63), Chaetoglobosin E (64), Chaetoglobosin F (65), and Penochalasin G (66) (Figure 3) were obtained from *C. globosum* endophyte obtained from seeds of *Panax notoginseng* collected at the Wenshan, Yunnan, China. Compounds 59 and 61–66 exhibited antifungal activity against *Phoma herbarum* (MIC in the range of 16–128 $\mu\text{g}/\text{mL}$) and *Epicoccum nigrum* (MIC in the range of <1–16 $\mu\text{g}/\text{mL}$). Both fungi are phytopathogenic fungi causing root rot of *Panax notoginseng* [42].

Ergosta-5,7,22-trien-3beta-ol (67) (Figure 3) was isolated from *Chaetomium cupreum* ZJWCF079 of *Macleaya cordata*. It exhibited antifungal activity against *Sclerotinia sclerotiorum* and *B. cinerea*, plant pathogenic fungi with EC₅₀ values of 125 $\mu\text{g}/\text{mL}$ and 190 $\mu\text{g}/\text{mL}$ respectively, but had no effects on *Pythium ultimum*, *Rhizoctonia solani* and *F. oxysporum* [43].

Chaetoglobosin A (59) (Figure 3), D (60), E (64), C (68), (Figure 3) G (69), and R (70) (Figure 4) were isolated from *Chaetomium globosum* No.04 obtained from barks of *Ginkgo biloba*, growing in Linyi, Shandong Province, China. Compounds 59, 60, 64, and 68–70 showed good growth inhibitory activity at a concentration of 20 $\mu\text{g}/\text{disk}$, against *Rhizopus stolonifer* and *Coniothyrium diplodiella* [44].

A new tetranorlabdane diterpenoids botryosphaerin H (71) and a known tetranorlabdane diterpenes 13,14,15,16-tetranorlabd-7-en-19,6beta:12,17-diolide (72) (Figure 4) were obtained from *Botryosphaeria* sp. P483, an endophyte of *Huperzia serrata* collected in Xichou County, Yunnan Province, China. When tested at 100 $\mu\text{g}/\text{disk}$, compound 71 showed zone of inhibition of 9, 7, 7, 8, and 8 mm, against *Gaeumannomyces graminis*, *Fusarium solani*, *Pyricularia oryzae*, *Fusarium moniliforme*, and *F. oxysporum* while compound 72 showed zone of inhibition of 12, 10, 10, 11, 13 mm against *G. graminis*, *F. solani*, *P. oryzae*, *F. moniliforme*, and *F. oxysporum*. The standard Carbendazim (50 $\mu\text{g}/\text{disk}$) exhibited activity against *G. graminis*, *F. solani*, *P. oryzae*, *F. moniliforme*, and *F. oxysporum*, with the zone of inhibition of 14, 18, 15, 17 and 15 mm respectively [45].

Endophytic fungus *Botryosphaeria dothidea* KJ-1 associated with the stems of *Melia azedarach* collected at Yangling, Shaanxi Province, China was the source of pycnophorin (73), stemphyperylene (74), chaetoglobosin C (68), djalonensone (75) (Figure 4), alternariol (76), β -sitosterol glucoside (77),

and 5-hydroxymethylfurfural (78) (Figure 4). Stemphyperylene (74) exhibited good antifungal activity against *Alternaria solani* the plant pathogen with the MIC value of 1.57 μM comparable to commonly used fungicide, the carbendazim. Compounds 68, 73, and 75–78 showed good to average antifungal activities against *A. solani* (MICs of 6.25–25 μM) [46].

Two eicosanoic acids, 2-amino-3,4-dihydroxy-2-25-(hydroxymethyl)-14-oxo-6,12-eicosenoic acid (79) and myriocin (80) (Figure 4), were isolated from *Mycosphaerella* sp. an endophytic fungus of *Eugenia bimarginata* DC. (Myrtaceae) collected in Brazil (Savannah). These compounds displayed antifungal activities against several isolates of *C. neoformans* and *C. gattii*, with MIC values for compound 79 ranging from 1.3 to 2.50 $\mu\text{g}/\text{mL}$ and for compound 80 was 0.5 $\mu\text{g}/\text{mL}$ [47]. Both compounds exhibited antifungal activities against several isolates of *C. neoformans* and *C. gattii*, with MIC values ranging from 0.49 to 7.82 μM for compound 79 and 0.48–1.95 μM for compound 80 in another study. When checked by the checkerboard microtiter assay, both compounds exhibited synergistic activity against *C. gattii* with amphotericin B. Ultrastructural analysis divulges various signs of damage in *C. gattii* and *C. neoformans* cells treated with compounds 79 and 80, including deformities in cell shape, depressions on the surface, and withered cells. Compounds 79 and 80 showed less loss of cellular material in cells of *C. gattii* compared to those treated with amphotericin B.

The difference in cellular material loss increased in a test compound concentration-dependent manner. Compound 80 also induced the formation of several pseudohyphae, suggesting that it could reduce virulence in *C. gattii* cells [48].

Endophytic fungus *Guignardia* sp., associated with *Euphorbia sieboldiana* collected from Nanjing, Jiangsu, China was the source of guignardone N (81) and guignardic acid (82) (Figure 4). Both compounds were evaluated for their inhibitory effects alone and with fluconazole on the growth and biofilms of *Candida albicans*. At 6.3 $\mu\text{g}/\text{mL}$ combined with 0.031 $\mu\text{g}/\text{mL}$ of fluconazole, compounds 81 and 82 were found to have prominent inhibition on the growth of *C. albicans* with fractional inhibitory concentration (FIC) index values of 0.23 and 0.19, respectively. Combined with fluconazole, both (40 $\mu\text{g}/\text{mL}$ for (81) and 20 $\mu\text{g}/\text{mL}$ for (82) could also inhibit *C. albicans* biofilms and reverse the tolerance of *C. albicans* biofilms to fluconazole [49].

Antifungal hyalodendriol C (83), rhizopycnol D (84), palmariol B (85), TMC-264 (86), penicilliumolide B (87) and alternariol 9-methyl ether (88) (Figure 4) were obtained from the endophytic fungus *Hyalodendriella* sp. Ponipodef 12 associated with the healthy stems of the “Neva” hybrid of *Populus deltoides* Marsh × *P. nigra* L. were collected from Longhua in Hebei Province of China. Compound 83 displayed antifungal effects against the spore germination of *M. oryzae* with potent inhibition with the IC₅₀ value of 11.6 $\mu\text{g}/\text{mL}$, which was comparable with the positive control, carbendazim (IC₅₀ 6.9 $\mu\text{g}/\text{mL}$) [50]. Previously, it found that rhizopycnol D (84), palmariol B (85), TMC-264 (86), penicilliumolide B (87), and alternariol 9-methyl ether (88) exhibits antifungal activity against the spore germination of *M. oryzae* [51,52].

Mellein (89) (Figure 4), was isolated from *Pezicula* sp. associated with the twigs of *Forsythia viridissima*, Zhejiang Province, Southeast China. Antifungal activity of this compound was tested against *B. cinerea*, *Pythium ultimum*, *Fusarium oxysporum* f. sp. *cucumerinum*, *Colletotrichum orbiculare*, *Verticillium dahliae*, *Pyricularia oryzae*, *Pestalotia diospyri*, *Sclerotinia sclerotiorum* and *Fulvia fulva*. Compound 89 displayed antifungal activity against 9 plant pathogenic fungi, esp. *B. cinerea* and *F. fulva* with EC₅₀ values below 50 $\mu\text{g}/\text{mL}$ [53].

Endophytic fungus *Nodulisporium* sp. A21 associated with the leaves of *Ginkgo biloba* collected from Nanjing in Jiangsu Province, China was a source of anti-phytopathogenic sporothriolide (90) (Figure 4). In mycelia growth inhibition method, sporothriolide (90) showed antifungal activity against *Rhizoctonia solani* with the EC₅₀ value of 3.04 $\mu\text{g}/\text{mL}$ (11.6 μM) while the EC₅₀ of positive control carbendazim was 1.84 $\mu\text{g}/\text{mL}$ (9.6 μM). Sporothriolide (90) at 200 $\mu\text{g}/\text{mL}$ had a protective efficacy of 71.7% against Rice Sheath Blight in comparison with the protective efficacy 90.1% of the positive control of validamycin A at 200 $\mu\text{g}/\text{mL}$. Conidia of *Magnaporthe oryzae* could not form the germ tube and appressorium germinate in the sporothriolide with solution at the concentration of 1.5 $\mu\text{g}/\text{mL}$ (5%

DMSO). In vivo, sporothriolide at 50 µg/mL, compared to tricyclazole of 2.5 µg/mL, could control the developing of Rice Blast [54]. Sporothriolide (90) also showed antifungal activity against *Sclerotinia sclerotiorum* with EC₅₀ of sporothriolide against was 2.78 µg/mL (10.7 µM) while the EC₅₀ of positive control carbendazim was 0.17 µg/mL (0.89 µM). When it comes to the protective activity on rape leaves, the positive control carbendazim of 250 µg/mL was 57.6% and sporothriolide of 250 µg/mL was 41.5% [54].

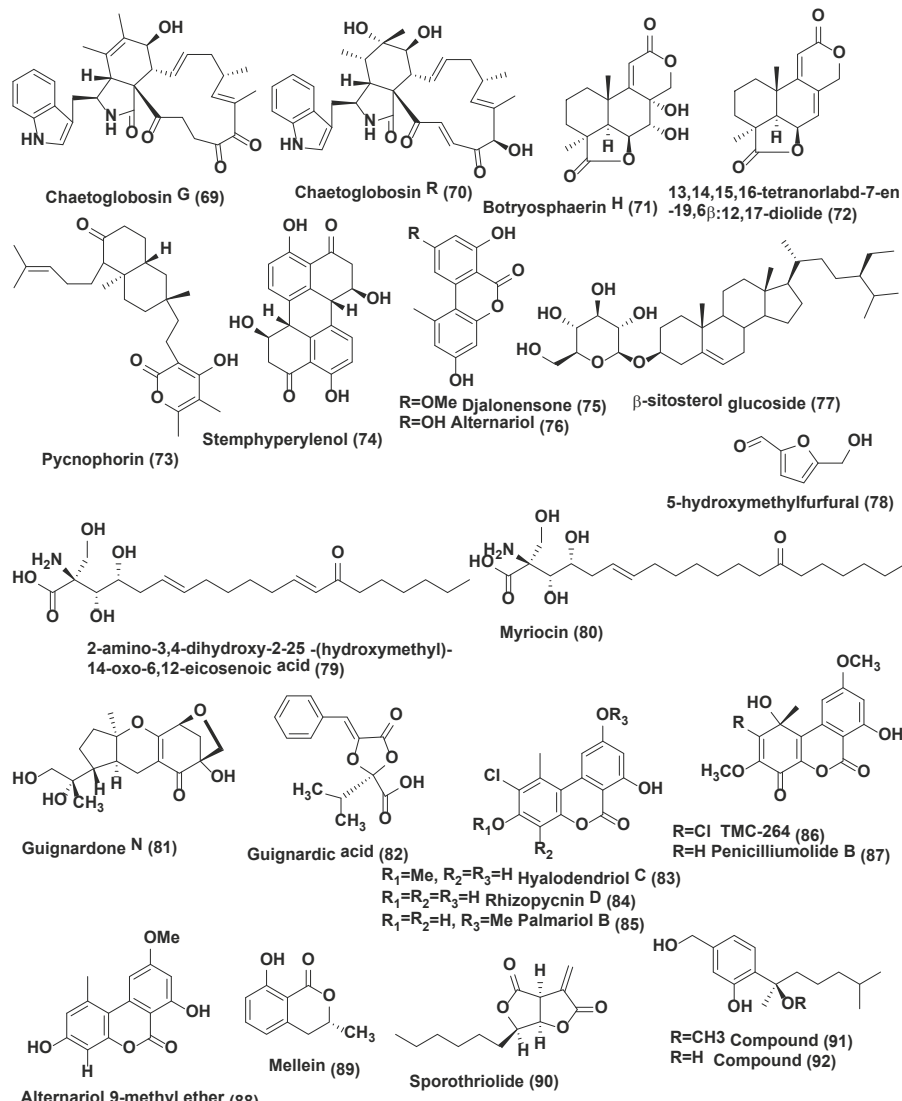


Figure 4. Structures of metabolites isolated from Ascomycetes (69–92).

Six phenolic bisabolane-type sesquiterpenoids (91, 92) (Figure 4) (93–96), along with a macrolide, pyrenophorin (97) (Figure 5) were isolated from *Lopherdermium nitens* DAOM 250027 endophyte of

Pinus strobus (eastern white pine) near Sussex, NB, Canada. These compounds were characterized based on interpretation of spectroscopic data (NMR, OR, UV) and HRMS. All compounds were tested for antifungal activity. Pyrenophorin (97) significantly reduced the growth of *Microbotryum violaceum* and *Saccharomyces cerevisiae* at 5 μ M, whereas sesquiterpenoids (91–96) were antifungal at 50 μ M to both species tested [55]. Isocoumarin derivative exserolide C (98) and (12R)-12-hydroxymonocerin (99) (Figure 5) were isolated from endophyte *Exserohilum* sp. associated with *Acer truncatum* collected from Beijing, China. Compounds 98 and 99 displayed antifungal activity against *Fusarium oxysporum*, with MIC value of 20 μ g/mL for both compounds, while Amphotericin B the positive control showed the MIC value of 0.63 μ g/mL [56].

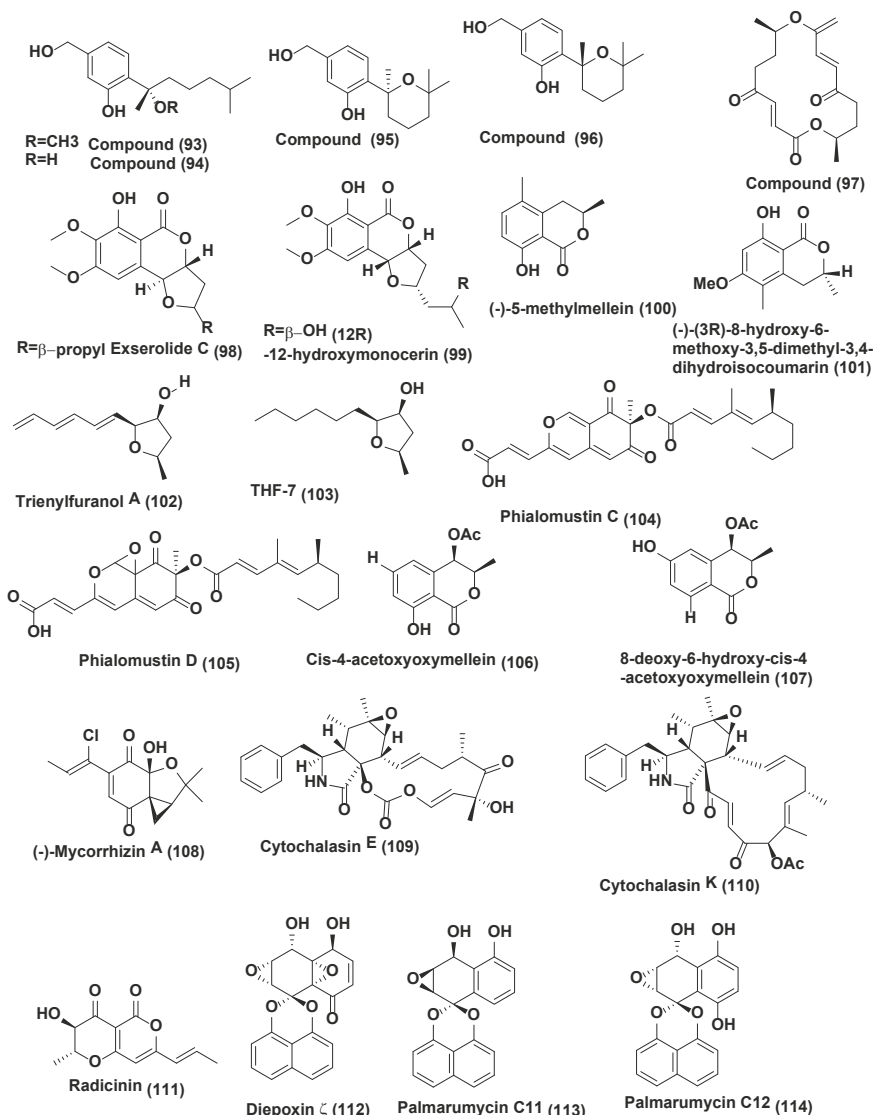


Figure 5. Structures of metabolites isolated from Ascomycetes (93–114).

Endophytic fungus of *Echinacea purpurea* associated with *Biscogniauxia mediterranea* EPU38CA from the wild in Missouri, USA was the source of (−)-5-methylmellein (**100**) and (−)-(3R)-8-hydroxy-6-methoxy-3,5-dimethyl-3,4-dihydroisocoumarin (**101**) (Figure 5). Compound **100** exhibited poor activity against *Phomopsis obscurans*, *P. viticola*, and *Fusarium oxysporum*, and stimulated the growth of *Colletotrichum fragariae*, *C. acutatum*, *C. gloeosporioides*, and *B. cinerea*. Compound **101** was acknowledged to be marginally more active in the microtiter method than 5-methylmellein [57]. Compound **101** was found to be slightly more active in the microtiter method than 5-methylmellein [57]. Trienylfuranol A (**102**) (Figure 5) was isolated from isolated from *Hypoxyylon submonticulosum* the endophyte of *Rubus idaeus* collected from Jordan Station, ON, Canada. It was identified based on high-resolution LC-MS and 1- and 2-D NMR spectroscopy. Absolute stereochemical configurations of the compounds were confirmed by NOE NMR experiments and by the preparation of Mosher esters. Complete hydrogenation of I yielded THF 7 (**103**) (Figure 5) that was used for stereochemical characterization and assessment of antifungal activity. Compound THF 7 (**103**) significantly inhibited the growth of *Saccharomyces cerevisiae* ($74 \pm 4\%$ inhibition) at a concentration of 250 µg/mL as compared with complete inhibition by nystatin at 10 µg/mL [58].

Endophytic fungus *Phialophora mustea* associated with *Crocus sativus* was the source of an unprecedented azaphilone derived skeleton, Phialomustin C (**104**) and D (**105**) (Figure 5). Compounds **104** and **105** showed potent activities against *Candida albicans*, with IC₅₀ values of 14.3 and 73.6 µM respectively [59].

An unidentified ascomycete, associated with *Melilotus dentatus* was the source of two new polyketide metabolites *cis*-4-acetoxymellein (**106**) and 8-deoxy-6-hydroxy-*cis*-4-acetoxymellein (**107**) (Figure 5). Compounds **106** and **107** displayed potent antifungal activities toward *Microbotryum violaceum* and *B. cinerea*, with 8 mm zone of inhibition for both fungi tested (0.05 mg was pipetted onto 9 mm sterile filter paper disk). In the case of *B. cinerea*, there was some growth within the zone of inhibition [60].

(−)-Mycorrhizin A (**108**) (Figure 6) was isolated from *Plectophomella* sp. while cytochalasins E (**109**) and K (**110**) (Figure 5) were isolated from *Physalospora* sp. Similarly, radicinin (**111**) (Figure 5) was purified from the endophytic fungus *Crataegus monogyna*. (−)-Mycorrhizin A showed good antifungal activity towards *Ustilago violacea* and *Eurotium repens*. Cytochalasins E (**109**) and K (**110**) showed potent activity against *E. repens* and *Mycotypha microspora*. Radicinin (**111**) (Figure 5) showed good activity against *E. repens* and *M. microspore* [61].

Diepoxin ζ (**112**), palmarumycin C11 (**113**), palmarumycin C12 (**114**) (Figure 5), cladospirone B (**115**), palmarumycin C6 (**116**), 1,4,7 β -trihydroxy-8-(spirodioxy-1',8'-naphthyl)-7,8-dihydronaphthalene (**117**), and palmarumycin C8 (**118**) (Figure 6) were obtained from *Berkleasmium* sp., an endophyte associated with *Dioscorea zingiberensis* from Hubei Province, China. Compounds **112–118** were evaluated for their antifungal activity against the spore germination of *M. oryzae*. Compounds **112–118** inhibited spore germination of *M. oryzae* with IC₅₀ values in the range 9.1–124.5 µg/mL. Palmarumycin C8 (**118**) showed the best inhibitory activity (IC₅₀ 9.1 µg/mL) among the compounds tested, although not as active as the positive control carbendazim (IC₅₀ 6.3 µg/mL) [62].

Bipolamide B (**119**) (Figure 6) was isolated from *Bipolaris* sp. MU34, the endophytic fungus associated with the leaves of *Gynura hispida* Thwaites collected from Mahidol University, Bangkok, Thailand. The compounds were characterized based on NMR and MS experiments. Bipolamide B (**119**) exhibited average antifungal activity with MIC values of 16, 32, 32, 64 and 64 µg/mL, against *Cladosporium cladosporioides*, *C. cucumerinum*, *Saccharomyces cerevisiae*, *Aspergillus niger* and *Rhizopus oryzae* respectively [63].

Altenusin (**120**) (Figure 6), a biphenyl derivative, was isolated from an endophytic fungus, *Alternaria alternata* Tche-153 of *Terminalia chebula*, collected from Bangkok, Thailand. Employing disk diffusion method and the microdilution checkerboard technique, altenusin (**120**) in amalgamation with each of three azole drugs, ketoconazole, fluconazole or itraconazole at their low sub-inhibitory concentrations displayed potent synergistic activity against *C. albicans* with the fractional inhibitory

concentration index range of 0.078 to 0.188 [64]. It is reported that *Schizosaccharomyces pombe* cells treated with altenusin were more rounded in shape than untreated cells which suggest that altenusin could act through the inhibition of cell wall synthesis or assembly in *S. pombe* [65].

Cladosporin (121) and isocladosporin (122) (Figure 6) were isolated from endophytic fungus *Cladosporium cladosporioides*. Compound 121 exhibited growth inhibition against *Colletotrichum acutatum*, *C. fragariae*, *C. gloeosporioides* and *Phomopsis viticola* at 30 μ M with 92.7%, 90.1%, 95.4%, and 79.9%, respectively. Similarly, compound 122 showed 50.4%, 60.2%, and 83.0% growth inhibition against *C. fragariae*, *C. gloeosporioides*, and *P. viticola*, respectively, at 30 μ M [66].

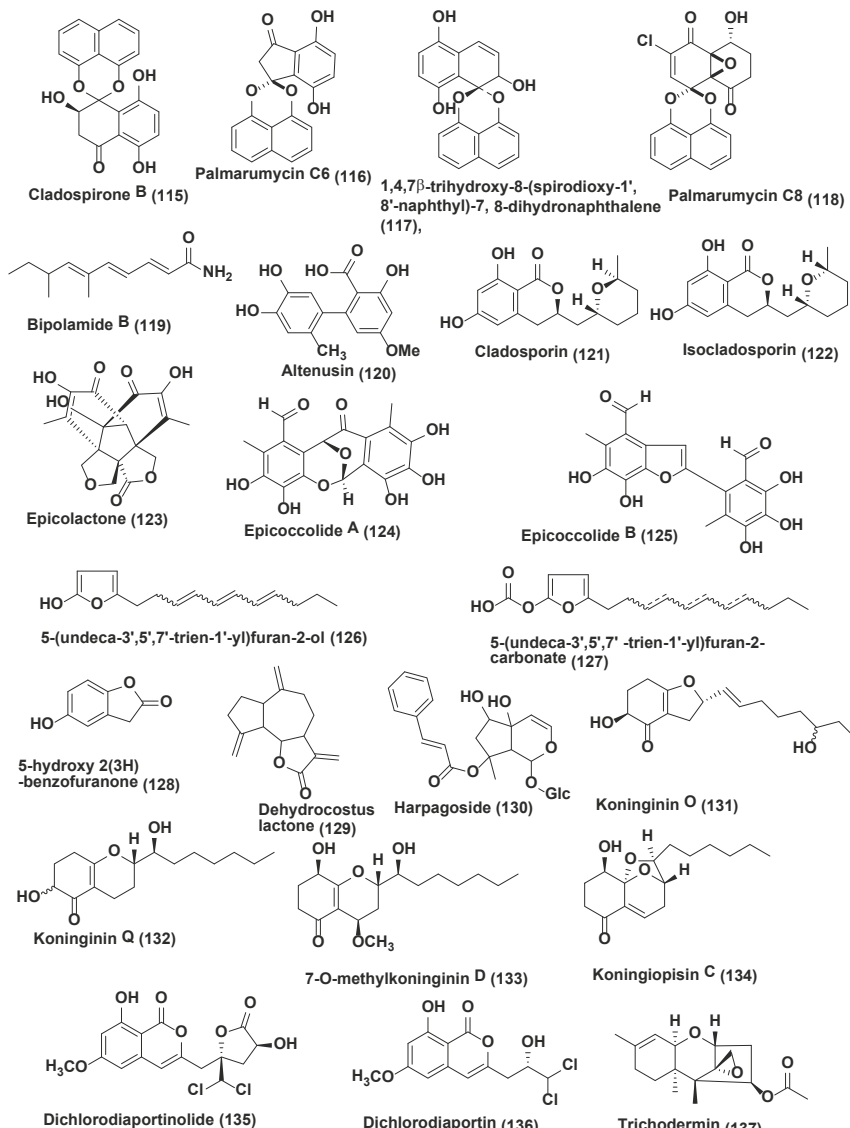


Figure 6. Structures of metabolites isolated from Ascomycetes (115–127) and Hyphomycetes (128–137).

Epicolactone (123) and epicoccolide A (124) and B (125) (Figure 6) polyoxygenated polyketides were obtained from an endophytic fungus, *Epicoccum* sp. CAFTBO, associated with stem bark and leaves of *Theobroma cacao* of Mount Kala, Republic of Cameroon. Compounds (123–125) showed good inhibitory effects on the mycelial growth of *Pythium ultimum* and *Aphanomyces cochlioides* and *Rhizoctonia solani* (MIC in the range of 20–80 µg per paper disc) [67].

5-methylmellein (100) (Figure 5) was isolated from endophytic fungus *Biscogniauxia mediterranea* Ohu 19B obtained from *Opuntia humifusa* (Cactaceae) of United States. Antifungal activity of compound 100 was evaluated using an in vitro microdilution broth assay against seven plant pathogens i.e., *Colletotrichum acutatum*, *C. fragariae*, *C. gloeosporioides*, *Fusarium oxysporum*, *B. cinerea*, *Phomopsis obscurans*, and *P. viticola*. *Phomopsis obscurans* was found to be being most susceptible (63.5% growth inhibition) at 150 µM at 120 h. The best growth inhibition (20.1%) to *F. oxysporum* was at 300 µM at 48 h. Lower doses (75 and 150 µM) of this compound caused stimulation of *B. cinera* and *C. fragariae*, while all doses caused stimulation of *C. acutatum* and *C. gloeosporioides* [68].

5-(undeca-3',5',7'-trien-1'-yl)furan-2-ol (126) and 5-(undeca-3',5',7'-trien-1'-yl)furan-2-carbonate (127) (Figure 6), two new alkylated furan derivatives, were recovered from the endophytic fungus *Emericella* sp. XL029 associated with the leaves of *Panax notoginseng* collected from Shijiazhuang, Hebei Province, China. Compound 126 displayed good antifungal activity against *Rhizoctorzia solani*, *Verticillium dahliae*, *Helminthosporium maydis*, *Fusarium oxysporum*, *Fusarium tricinctum*, *Botryosphaeria dothidea*, and *Alternaria fragariae* with MIC values ranging from 25 to 3.1 µg/mL, while compound 127 was found active against *V. dahliae*, *H. maydis*, *F. tricinctum*, *B. dothidea*, and *A. fragariae* with MIC values ranging from 50 to 12.5 µg/mL [69].

2.3. Compounds Produced by Hyphomycetes

5-hydroxy 2(3H)-benzofuranone (128), dehydrocostus lactone (129) and harpagoside (130) (Figure 6) were isolated from *Fusarium fujikuroi*, *Penicillium chrysogenum* and *Penicillium expunsum* endophytes of *Eleusine coracana* grown under semi-hydroponic conditions Arkell Field Station, Arkell, ON, Canada. Compounds 128–130 exhibited antifungal activity against *F. graminearum* with the MIC of 31.25, 250.00 and 31.25 µg/mL, respectively. An in vitro interaction between each compound and *Fusarium* was investigated using light microscopy and vitality staining where the results proposed a mixed fungicidal/fungistatic mode of action [70].

Endophytic fungus *Trichoderma koningiopsis* YIM PH30002 harbored in *Panax notoginseng* collected from Wenshan, Yunnan Province, China was the source of koninginin O (131), koninginin Q (132) and 7-O-methylkoninginin D (133) (Figure 6). The antifungal activities of these compounds were tested against phytopathogenic fungi, *Fusarium oxysporum*, *F. solani*, *F. flocciferum*, *Plectosphaerella cucumerina* and *Alternaria panax* which are causes of pathogens of root rot diseases of *P. notoginseng*. Koninginin O (131) and koninginin Q (132) exhibited poor activity against *F. oxysporum* and *P. cucumerina* (MIC of 128 µg/mL). 7-O-methylkoninginin D (133) also showed poor activity against *P. cucumerina* (MIC 128 µg/mL). Nystatin positive control showed antifungal activity with MICs at 32 µg/mL [71]. Koningiopisin C (134) (Figure 6) was also isolated from the same fungus. Koningiopisin C showed antimicrobial activities against *F. oxysporum*, *A. panax*, *F. solani* and *P. cucumerina* with MICs at 32, 64, 32, and 16 µg/mL, respectively [72].

Dichlorodiaportinolide (135) and dichlorodiaportin (136) (Figure 6) were isolated from endophytic fungus *Trichoderma* sp. 09 obtained from the root of *Myoporum bontioides* A. Dichlorodiaportinolide (135) and dichlorodiaportin (136) showed weak to high antifungal activities with MIC values ranging from 6.25 to 150 µg/mL against *Colletotrichum musae* and *Rhizoctonia solani* and were inactive to *Penicillium italic* and *Fusarium graminearum* (MIC values > 200 µg/mL) [73].

Trichodermin (137) (Figure 6) was isolated from endophytic fungus strain, *Trichoderma brevicompactum* 0248 obtained from *Allium sativum*. Trichodermin showed potent inhibitory activity against *Rhizoctonia solani*, with an EC₅₀ of 0.25 µg/mL and against *B. cinerea*, with an EC₅₀ of 2.02 µg/mL but relatively

poorly active against *Colletotrichum lindemuthianum* ($EC_{50} = 25.60 \mu\text{g}/\text{mL}$). Compound **137** exhibited good antifungal activity against the tested phytopathogens compared with the positive control Carbendazim [74].

Trichoderma koningiopsis YIM PH30002 collected at Wenshan, Yunnan Province of China was the source of two new metabolites koninginin R and S (138–139) (Figure 7). These isolated compounds showed certain antifungal activities against phytopathogens, *Fusarium flociferum* and *Fusarium oxysporum*. Compound **138** possess the weak activity against *F. oxysporum* and *F. flociferum* with the MICs at 128 $\mu\text{g}/\text{mL}$, while compound **139** displayed the poor activity against *F. oxysporum* with the MIC at 128 $\mu\text{g}/\text{mL}$ [75].

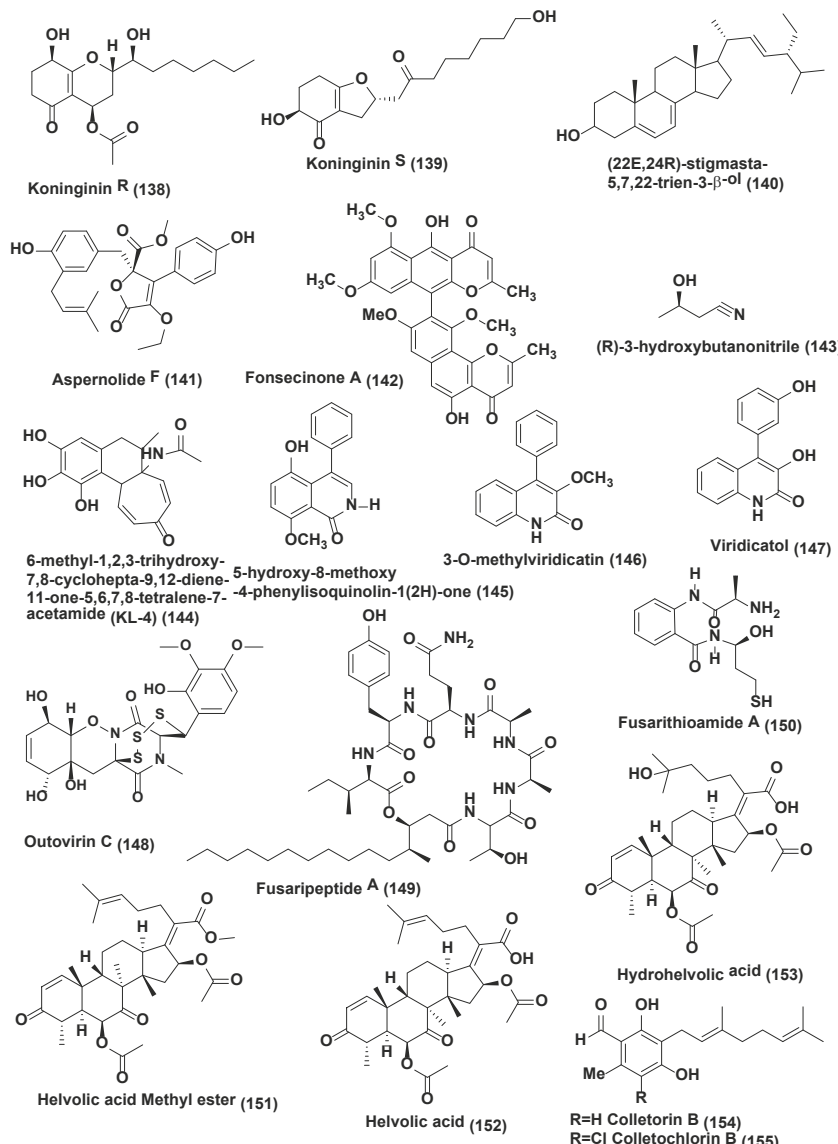


Figure 7. Structures of metabolites isolated from Hyphomycetes (138–155).

Stigmasterol derivative (22E,24R)-stigmasta-5,7,22-trien-3- β -ol (**140**) and a new butyrolactones, aspernolide F (**141**) (Figure 7) were obtained from the endophytic fungus *Aspergillus terreus* associated with the roots of *Carthamus lanatus* collected at Assiut, Egypt. Compounds **140–141** exhibited good activity against *C. neoformans* with IC₅₀ values of 4.38 and 5.19 μ g/mL respectively, compared to amphotericin B (IC₅₀ 0.34 μ g/mL) [76].

Fonsecinone A (**142**), and (R)-3-hydroxybutanonitrile (**143**) (Figure 7), were obtained from *Aspergillus* sp. KJ-9 an endophytic fungus associated with *Melia azedarach* which was collected at Yangling, Shaanxi Province, China, and identified by spectroscopic methods. Compounds **142** and **143** were active against *Gibberella saubinetti*, *Magnaporthe grisea*, *B. cinerea*, *C. gloeosporioides* and *A. solani* (MIC range of 6.25–50 μ M) [77].

6-methyl-1,2,3-trihydroxy-7,8-cyclohepta-9,12-diene-11-one-5,6,7,8-tetralene-7-acetamide (KL-4) (**144**) (Figure 7) was isolated from *Aspergillus* sp. obtained from the seeds of *Gloriosa superba* which were collected from Tirupati, India. KL-4 (**144**) exhibited good antifungal activity against *Saccharomyces cerevisiae*, *C. albicans* and *Cryptococcus gastricus* with MIC 25, 12.5, and 50 μ g/mL respectively [78].

Endophytic fungus *Penicillium* sp. R22 associated with *Nerium indicum* collected from Qinling Mountain, Shaanxi Province, China was the source of 5-hydroxy-8-methoxy-4-phenylisoquinolin-1(2H)-one (**145**) a new isoquinolone alkaloid along with 3-O-methylviridicatin (**146**) and viridicatol (**147**) (Figure 7) two known quinolinone alkaloids. Compound **145** exhibited good antifungal activity against *Alternaria brassicae*, *A. alternata* and *Valsa mali* with MIC value of 31.2 μ g/mL, compound **146** against *A. brassicae*, *B. cinerea* and *Valsa male* with MIC value of 31.2 μ g/mL, compound **147** against *A. brassicae*, *A. alternata* and *B. cinerea* with MIC value of 31.2 μ g/mL [79].

Trisulfide glovirin-like compound Outovirin C (**148**) (Figure 7), an epithiodiketopiperazine natural product, was identified from *Penicillium raciborskii*, an endophytic fungus associated with *Rhododendron tomentosum* were collected at the test site of University of Oulu, Finland. Outovirin C (**148**) showed antifungal activity when assayed by micro-spectrophotometry using a dose response growth inhibition assay. Outovirin C inhibited the growth of *Fusarium oxysporum*, *B. cinerea*, and *Verticillium dahliae* at a low concentration of 0.38 mM (207 μ g/mL) but a more significant growth inhibition was observed at the higher concentration of 0.76 mM (413 μ g/mL). Compound **148** was most active against *B. cinerea* (57% inhibition) and slightly less effective against *V. dahliae* (45% inhibition) [80].

Fusari peptide A (**149**) (Figure 7), a new cyclodepsipeptide, was isolated from the culture of the endophytic fungus *Fusarium* sp. associated with roots of *Mentha longifolia* growing in Saudi Arabia. Its structure was elucidated based on 1D and 2D NMR and HRESI and GC-MS experiments. The absolute configuration of the amino acid residues of **149** was assigned by chiral GC-MS and Marfey's analysis after acid hydrolysis. Compound **149** exhibited potent antifungal activity toward *C. albicans*, *C. glabrata*, *C. krusei*, and *A. fumigates* with IC₅₀ values of 0.11, 0.24, 0.19, and 0.14 μ M, respectively. Under similar condition control amphotericin B exhibited antifungal activity toward *C. albicans*, *C. glabrata*, *C. krusei*, and *A. fumigates* with IC₅₀ values of 0.3, 0.6, 0.5, 0.7 μ M, respectively [81].

Fusarithioamide A, a new benzamide derivative (**150**) (Figure 7) was isolated from *Fusarium chlamydosporium* associated with the leaves of *Anvillea garcinii* collected from Al-Azhar University, Saudi Arabia. Compound **150** exhibited good antifungal activity against *C. albicans* with inhibition zone diameters (IZD 16.2 mm and MIC 2.6 μ g/mL which is comparable to the positive control substance clotrimazole (IZD 18.5 mm and MIC 3.7 μ g/mL) [82].

A new helvolic acid derivative named helvolic acid methyl ester (**151**), together with two known helvolic acid compounds, helvolic acid (**152**) and hydrohelvolic acid (**153**) (Figure 7), were extracted from endophytic fungus *Fusarium* sp. associated with *Ficus carica* leaves collected from Qinling Mountain, Shaanxi Province, China. Compounds **151–153** exhibited good antifungal activity against *B. cinerea*, *C. gloeosporioides*, *F. oxysporum* f. sp. *niveum*, *Fusarium graminearum* and *Phytophthora capsici* with MIC value in the range of 12.5–25 μ g/mL while Carbendazim the standard showed MIC value in the range of 32.2–62.5 μ g/mL against the same fungi [83].

Collectorin B (154), colletochlorin B (155) (Figure 7), LL-Z1272 β (llicolin B) (156), and 4,5-dihydrodechloroaschlorin (157) (Figure 8) were extracted from endophytic fungus *Fusarium* sp. Collectorin B (154) and colletochlorin B (155) showed moderate antifungal activity towards *Ustilago violacea* and *F. oxysporum*. Compound 156 showed moderate antifungal activity towards *U. violacea* and *F. oxysporum*. Furthermore, 4,5-dihydrodechloroaschlorin (157) showed a very strong antifungal activity towards *Eurotium repens* [84].

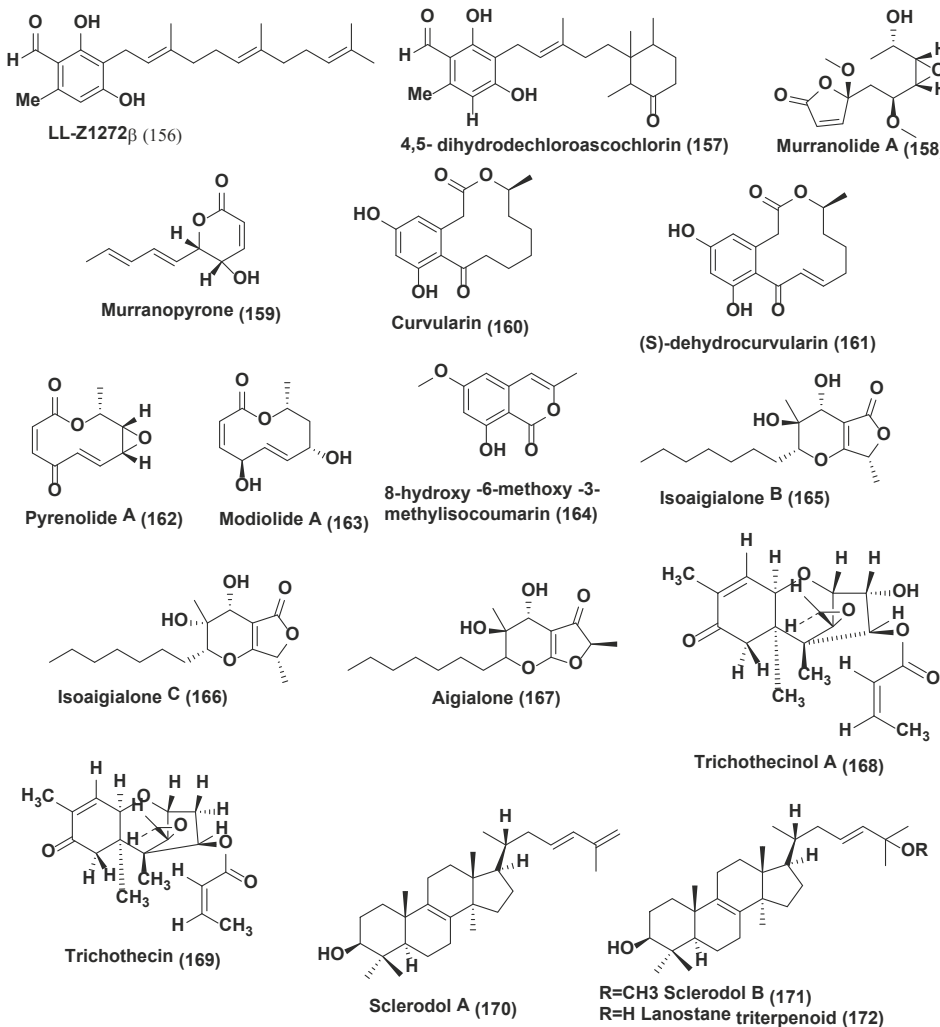


Figure 8. Structures of metabolites isolated from Hyphomycetes (156–169) and Basidiomycetes (170–172).

Murranolide A (158), murranopyrone (159), curvularin (160), (S)-dehydrocurvularin (161), pyrenolide A (162), modiolide A (163), and 8-hydroxy-6-methoxy-3-methylisocoumarin (164) (Figure 8) were extracted from the endophytic fungus *Curvularia* sp., strain M12, associated with the leaf of *Murraya koenigii* were collected from Rajshahi University, Bangladesh. Pyrenolide A (162) was observed to impair the mobility of *Phytophthora capsici* zoospores in a short time (30 min) at a low

concentration (100% at 0.5 µg/mL). Murranolide A (**158**), murranopyrone (**159**), curvularin (**160**), (S)-dehydrocurvularin (**161**), modiolide A (**163**), and 8-hydroxy-6-methoxy-3-methylisocoumarin (**164**) exhibited zoospore motility impairment activity at higher concentrations (IC_{50} : 50–100 µg/mL) [85].

Two new isoagialones, B (**165**) and C (**166**) (Figure 8), along with agialone (**167**) (Figure 8), were obtained from *Phaeoacremonium* sp., an endophytic fungus associated with the leaves of *Senna spectabilis* was collected in the Araraquara Cerrado area, in June 2001, Araraquara, São Paulo state, Brazil. These compounds were evaluated against *Cladosporium cladosporioides* and *C. sphaerospermum* using direct bioautography. Compounds **165** and **167** exhibited antifungal activity, with a detection limit of 5 µg, for both fungi, while compound **166** displayed weak activity (detection limit > 5 µg), with a detection limit of 25 µg. Nystatin was used as a positive control, showing a detection limit of 1 µg [86].

Trichothecinol A (**168**) (Figure 8) was obtained from *Trichothecium* sp. an endophytic fungus isolated from *Phyllanthus amarus* collected from Pune India. Compound **101** showed activity against *Cryptococcus albidus* (NCIM 3372) up to 20 µg/mL [87]. Trichothecin (**169**) (Figure 8), a sesquiterpene, was isolated from endophytic fungus *Trichothecium* sp. residing inside the leaves of *Phyllanthus* sp. collected from Pune India. Compound **169** exhibited anti-fungal activity against *Saccharomyces cerevisiae*, *Cryptococcus albidus* var *diffluens* NCIM 3371, *Cryptococcus albidus* var *diffluens* NCIM 3372, *Fusarium oxysporum*, *Penicillium expansum*, *Trichoderma viride*, *Paecilomyces varioti* and *Aspergillus niger* with MIC of 6, 20, 12, 10, 30, 40, 20 and 12 µg/mL, respectively [88].

2.4. Compounds Produced by Basidiomycetes

Two lanostane triterpenoids, sclerodols A (**170**) and B (**171**), and a known related lanostane triterpenoid (**172**) (Figure 8) were isolated from *Sclerotinia* UFSM Sc1 (Persoon) Fries an endophyte associated with *Eucalyptus grandis*. Both compounds were evaluated for their anti-candidal potential against *Candida albicans*, *C. tropicalis*, *C. crusei*, *C. parapsoriasis* for activities. Compound **171** showed good anticandidal activity against *C. albicans*, *C. tropicalis*, *C. crusei*, *C. parapsoriasis* with the MIC of 25.0, 25.0, 6.25 and 12.5 and MFC of 25.0, 25.0, 12.5 and 25.0 µg/mL respectively. Compounds **170** and **172** were less active against tested strain than compound **171** with the MIC in the range of 12.5–100 and MLC (minimal lethal concentration) of >100.0 µg/mL. Control nystatin exhibited showed anti-candidal activities against tested strains with the MIC in the range of 0.77–1.52 µg/mL and MLC in the range of 3.12–6.25 µg/mL [89].

Table 1. Antifungal compounds reported from endophytic fungi.

Sr. No.	Fungus	Plantsource	Compounds Isolated	Biological activity *	Ref.
1	<i>Pestalotopsis foetidum</i>	<i>Bursera sexangularis</i> Hainan, China	(3R,4R,6R,7S)-7-hydroxy-5,7-dimethyl-octahydro-3-(7-methylcyclohexenyl)-propanoic acid (2)	Compound 1 <i>B. cinerea</i> and <i>P. notitiae</i> (MIC 31 and 6.3 μ g/mL), ketocazole (MIC 3.1 μ g/mL each) Compound 2 <i>C. albanicus</i> MIC 50 μ g/mL [15]	
2	<i>Pestalotopsis</i> sp. DO14	<i>Dendrobium officinale</i> , Yandang Mountain, Zhejiang Province, China.	(4S,6S)-6-[(1S,2R)-1,2-dihydroxybutyl]-4-hydroxy-4- methoxytetrahydro-2H-pyran-2-one (3) and (6S,2E)-6-hydroxy-3-methoxy-5-oxo-dec-2-enoic acid (4), L-L-7880 Y (5), LL-7880 X (6)	Compounds 3–6 active against <i>C. albanicus</i> , <i>C. neoformans</i> , <i>T. rubrum</i> , and <i>A. fumigatus</i> ($\text{MIC} \leq 50$ μ g/mL) Compounds 3–4 active against <i>C. albanicus</i> , <i>C. neoformans</i> , <i>T. rubrum</i> , and <i>A. fumigatus</i> ($\text{MIC} \leq 25$ μ g/mL)	[16]
3	<i>Pestalotopsis fici</i>	<i>Camellia sinensis</i> Hangzhou, China.	Ficyprone A (7)	Compound 7 active against <i>G. zae</i> (IC_{50} 15.9 μ M), ketconazole (IC_{50} 6.02 μ M)	[17]
4	<i>Pestalotopsis mangiferae</i>	<i>Mangifera indica</i> Maduravoyal, Tamil Nadu Province, India.	4-(2,4,7-trioxa-bicyclo[4.1.0]heptan-2-yl) phenol (8)	Compound 8 active against <i>C. albanicus</i> (MIC 0.039 μ g/mL), Nystatin (MIC 10.6 μ g/mL)	[18]
5	<i>Phomopsis</i> sp.	<i>Sonja spectabilis</i> Sao Paulo, Brazil	Cytochalasin H (9)	Compound 9 active against <i>C. cladosporioides</i> and <i>C. sphaerospermum</i> (MIC 1.0 and 25.0 μ g, respectively), Nystatin (MIC = 1.0 μ g)	[19]
6	<i>Phomopsis</i> sp.	<i>Acutium corniculatum</i> , Huize County, Yunnan Province, China.	(14 β ,22 β)-9,14-dihydroxyergosta-4,7(22)-fene-3 β -olone (10), (5 α ,6 β ,15 β ,22 β)-6-ethoxy-5,15-dihydroxyergosta-7(22)-fene-3 β -one (11), calasterol A (12), and gomodermaside (13)	Compound 10 active against <i>C. albanicus</i> , <i>H. competitum</i> , and <i>A. niger</i> (MIC, 64, 64, and 128 μ g/mL, respectively). Compound 11 active against <i>C. albicans</i> and <i>E. aviculorum</i> (MIC = 128 μ g/mL). Compounds 12 and 13 active against <i>F. avenaceum</i> (MIC, 64 μ g/mL). Compound 12 active against <i>P. oryzae</i> and <i>T. sugii</i> (MIC 1.28 and 256 μ g/mL)	[20]
7	<i>Diplothele maritima</i>	<i>Picea</i> sp., Acadian forest of Eastern Canada.	Phenoposolidope A (14), B (15), and C (16), and a stable alpha-pharyrone (17)	Compound 14 active against <i>M. violaceum</i> and <i>S. cerevisiae</i> at 25 μ M, Compounds 15–17 demonstrated growth inhibition at 250 μ M	[21]
8	<i>Phoma</i> sp.	<i>Fucus serratus</i> .	Phomalacton (18), (3R)-5-hydroxymelikone (19) and erinodin (20)	Compounds 18–20 active against <i>M. violaceum</i> with 5, 6 and 5 mm zone of inhibition	[23]
9	<i>Phoma</i> sp. WF4	<i>Elatine coracina</i> Arkell Field Station, Arkell, ON, Canada	Viridicatol (21), tenuazonic acid (22), alternatriol (23), and alternatriol monomethyl ester (24)	Compounds 21–24 caused dramatic breakage of <i>E. graminifolium</i> hyphae in vitro	[24]
10	<i>Rhizopus nigricans</i> Nitaf 22	<i>Nicotiana tabacum</i> , China Agricultural University, Beijing 100193, China.	Rhizopycin D (25) and TM-C-264 (26)	Compounds 25–26 inhibited the spore germination of <i>M. oryzae</i> with IC_{50} values of 59 and 12.0 μ g/mL, respectively	[25]
11	<i>Micromphaleurosis</i> sp. <i>Sclerotiorum</i> sp.	<i>Salsola oppositifolia</i> , Playa del Ingles, Gomera, Spain	Microsphaerol (27) Sematoneone (28)	Compounds 27 and 28 active against <i>M. violaceum</i> with 9 and 5 mm zone of inhibition. In addition, there was some growth with in zone of inhibition comparable to that of the positive control nystatin	[26]
12	<i>Colletotrichum</i> <i>gloeosporioides</i>	<i>Michelia champaca</i> Sao Paulo State University, Araçatuba, São Paulo, Brazil.	2-phenylethyl 1H-indol-3-yl-acetate (29)	Compound 29 active against <i>C. cladosporioides</i> and <i>C. sphaerospermum</i>	[27]
13	<i>Colletotrichum</i> sp.	Gomera (Spain)	Colletomic acid (30)	Compound 30 active against <i>M. violaceum</i> with 7 mm zone of inhibition	[28]
14	<i>Coniothyrium</i> sp.	<i>Salsola oppositifolia</i> Gomera in the Canary Islands.	1,7-dihydroxy-3-methyl-9-(0-anthrquinone) (31), 1,6-dihydroxy-3- -methyl-9-(0-anthrquinone) (32), and 1-hydroxy-3-hydroxymethyl-9-(0-anthrquinone) (33) coniothyrones A-D (34–37)	Compounds 31–37 active against <i>M. violaceum</i> with 7, 10, 8, 7.5, 6, 8 and 7.5 mm zone of inhibition. Compounds 32–34 active against <i>M. violaceum</i> (10 and 9 mm zone of inhibition) and <i>B. cinerea</i> (7.5 and 12.5 mm zone of inhibition) when tested under similar conditions	[29,30]

Table 1. Cont.

Sr. No.	Fungus	Plantsource	Compounds Isolated	Biological activity *	Ref.s
Compounds Produced by Ascomycetes					
15	Xylaria sp. YM 311647	Azadina indica, Yunnanjiang County, Yunnan Province, China	(S,4S,5R,7R,10R,11R)-Guaine-5,10(11,12)-diol (38) (S,4S,5S,7R,10R,11S)-Guaine-1,10(11,12)-diol (39) (S,4S,5S,7R,10R,11S)-Guaine-5,10,11(12)-triol (40) (R,5S,5S,7R,10R,11R)-Guaine-1,10,11(12)-triol (41) (R,5S,4R,5S,7R,10R,11S)-Guaine-3,10,11,12-teranol (42) (R,4S,5S,7S,9R,10S,11R)-Guaine-9,10,11,12-teranol (43) (R,4S,5S,7S,9R,10S,11S)-Guaine-10,11,12-teranol (44) (R,4S,5S,7R,10R,11R)-Guaine-10,11,12-teriol (45) 14a,16-Epoxy-18-norcoprin-7-en-4a-h (47), 16-C-5-hydroxy-18-norcoprin-7-en-4a-h, 16-diol (48), and 9-Deoxy-5-hydroxy A (49)	Compounds 38–54 active against <i>C. allicans</i> and <i>H. compactum</i> (MIC in the range of 32 to 256 µg/mL), compounds 47–49 active against <i>C. allicans</i> , <i>A. niger</i> , <i>P. oryzae</i> , <i>F. avenaceum</i> , and <i>H. compactum</i> MIC in the range of 16 to 256 µg/mL. Compound 49 exhibited the potent inhibitory activity against <i>C. allicans</i> and <i>P. oryzae</i> with MIC values of 16 µg/mL	[32]
16	Xylaria sp. YM 31164	Azadina indica, Yunnanjiang County, Yunnan Province, China	(S,2S,4R,5R,7R,10R)-Guaine-2,4,10,11,12-pentaol (50), (S,2S,4R,5R,7R,10R)-Guaine-4,5,10,11,12-pentaol (52), (S,4R,5S,7R,10R)-Guaine-4,5,10,11,12-pentaol (53), (R,4S,5R,7R,10R)-Guaine-4,5,10,11,12-pentaol (54), Xyloide (55).	Compounds 50–54 active against <i>P. oryzae</i> and <i>H. compactum</i> (MIC in the range of 32–256 µg/mL). Compound 53 active against <i>P. oryzae</i> (MIC, 32 µg/mL). Compounds 52 and 53 active against <i>H. compactum</i> (MIC, 64 µg/mL). Compound 53 and 54 active against <i>C. allicans</i> (MIC, 32 µg/mL). Compound 52 active against <i>C. allicans</i> , <i>A. niger</i> , and <i>H. compactum</i> (MIC, 64 µg/mL).	[33]
17	X. kegerensis	Crotalaria lechleri.		Compound 55 active against <i>P. ultimum</i> (MIC, 425 µM)	[34]
18	Xylaria sp. XC-16	Tonka sinensis (Yangling, Shaanxi Province, China)	Cytchalasin Z28 (56)	Compound 56 active against <i>C. sativinelli</i> (MIC of 12.5 µM). Hymexazol (MIC = of 25 µM)	Zhang et al. [35]
19	Xylaria sp. strain FOD10, Xylaria sp. PSU-G12X, catensis, 13 strains of Xylaria sp.	Aliis holophylla; <i>Garcinia hanburyana</i> ; <i>Asimina triloba</i> ; <i>Pinus strobus</i> ; <i>Vaccinium angustifolium</i> , New Brunswick and Nova Scotia, Canada	Griseofulvin (57)	Griseofulvin (57) Inhibits <i>A. nadi</i> , <i>B. cinerea</i> , <i>Calotrichidium gleosporioides</i> , <i>Corticium sepiarium</i> , <i>F. oxysporum</i> and <i>M. grisea</i> in vitro (I_{C_50} values of 180, 5.0, 1.7, 1.0, 30.0, and 1.4 µg/mL, respectively). Compound 57 active against <i>M. grisea</i> , <i>C. sessile</i> , <i>B. cinerea</i> , <i>P. recutitae</i> and <i>B. graminis</i> f. sp. <i>hordei</i> in vivo, with % of fungal control at 95, 100, 60, 90 and 90, respectively, at 150 µg/mL.	[36–38]
20	Xylaria sp.		Dechlorogriseofulvin (58)	Compound 58 showed weak antifungal activity with an I_{C_50} value of 200 µg/mL against <i>M. grisea</i> , <i>C. sessile</i> , <i>B. cinerea</i> , <i>P. recutitae</i> and <i>B. graminis</i> f. sp. <i>hordei</i> in vivo.	[36,38]
21	Chaetomium globosum CDW7	Ginkgo biloba China	Chaetoglobosin A (59) and D (60)	Compounds 59–60 active against <i>S. sclerotiorum</i> with I_{C_50} values of 0.35 and 0.62 µg/mL, respectively, carbendazim (0.17 µg/mL)	[41]
22	Chaetomium globosum	Seeds of <i>Panax notoginseng</i> collected at the Wenshan, Yunnan, China	Chaetoglobosin A (59), Chaetomuglin D (62), Chaetoglobosin B (63), Chaetoglobosin E (64), Chaetoglobosin F (65) and Pencalathsin G (66)	Compounds 59 and 61–66 active against <i>P. helianthi</i> (MIC in the range of 16–28 µg/mL) and <i>E. nigrum</i> (MIC in the range of <1–16 µg/mL).	[42]
23	Chaetomium cupreum ZJWCRD79		Ergosterol-5,7,22-trien-3-beta-ol (67)	Compound 67 against <i>S. sclerotiorum</i> and <i>B. cinerea</i> with EC ₅₀ values of 125 µg/mL and 190 µg/mL, respectively.	[43]
24	Chaetomium globosum NC004	Barks of <i>Cinchona ledgeri</i> , Linyi, Shandong Province, China.	Chaetoglobosin A (59), D (60), E (64), C (68), C (69), R (70)	Compounds 59, 60, 64, and 68–70 active against <i>R. stolonifer</i> and <i>C. apiphilicola</i> at a concentration of 20 µg/disk.	[44]
25	Boryosphphaeria sp. T483	Huperzia serrata, Xichou County, Yunnan Province, China	Botryosphaerin H (71) 13,14,15,16-tetranorlabd-7-en-19,6β,12,17-diolide (72)	At 10 µg/disk, compound 71 showed zone of inhibition of 9, 7, 7, 8, and 8 mm, against <i>G. gummans</i> , <i>F. solani</i> , <i>P. oryzae</i> , <i>F. moniliforme</i> , and <i>F. avenaceum</i> ; compound 72 showed zone of inhibition of 12, 10, 10, 11, and 13 mm against <i>G. gummans</i> , <i>F. solani</i> , <i>P. oryzae</i> , <i>F. moniliforme</i> , and <i>F. avenaceum</i> ; carbendazim (50 µg/disk) showed the zone of inhibition of 14, 18, 15, 17, 15 mm against <i>G. gummans</i> , <i>F. solani</i> , <i>P. oryzae</i> , <i>F. moniliforme</i> , <i>F. avenaceum</i> , respectively.	[45]

Table 1. Cont.

Sr. No.	Fungus	Plant source	Compounds isolated	Biological activity *	Ref.
26	<i>Boryosphaerioides</i> K1-1,	<i>Melia azedarach</i> Yangtling, Shaanxi Province, China.	Pyronorphin (73), strobilopyrenol (74), chastoglabosin C (68), dihydrostrobilopyrenol (75), alantolactone (76), β -sitosterol glucoside (77), 5-hydroxyisoxynonylfuranol (78)	Compound 74 active against <i>A. solani</i> (MICs of 1.57 μ M) Compounds 68, 73, and 75–78 active against <i>A. solani</i> (MICs of 6.25–25 μ M)	[46]
27	<i>Mycosphaerella</i> sp.	<i>Eugenia binagatina</i> DC. Brazil (savannah), myracrodia (80)	2-amino-3,4-dihydro-2-(hydroxymethyl)-14-oxo-6,12-ecosenoic acid (79), myracrodia (80)	Compounds 79 active against several isolates of <i>C. neofumans</i> and <i>C. gattii</i> , with MIC values ranging from 1.3 to 2.50 μ g/mL and 0.5–14.6 μ M, for compound 80	[47]
28	<i>Guignardia</i> sp.	<i>Euphorbia sieboldiana</i> collected from the campus of China Pharmaceutical University, Nanjing, Jiangsu, China	Guignardone N (81), guignardic acid (82)	Compounds 79 active against several isolates of <i>C. neofumans</i> and <i>C. gattii</i> , with MIC values ranging from 0.49 to 7.82 μ M and 0.48–1.95 μ M for compound 80.	[48]
29	<i>Hyalodendriella</i> sp. Pomipodet 12	"Neva" hybrid of <i>Populus deltoides</i> Marsh \times <i>P. nigra</i> L., Longhua in Hebei Province of China	hyalodendriol C (83), rhizopycyan D (84), palmitoil B (85), TMC-264 (86), penicillamolide A (87) and alternanol 9-methyl ether (88)	At 6.3 μ g/mL combined with 0.031 μ g/mL of flutonazole, compounds 81 and 82 were found to have prominent inhibition on the growth of <i>C. albicans</i> with FC index values of 0.23 and 0.19, respectively. Combined with flutonazole, both of them (40 μ g/mL for (81) and 20 μ g/mL for (82)) could also inhibit <i>C. albicans</i> biofilms and reverse the tolerance of <i>C. albicans</i> biofilms to flutonazole	[49]
30	<i>Pericilia</i> sp.	<i>Forsythia viridissima</i> , collected from Zhejiang Province, Southeast China	Mellein (89)	Compound 88 exhibited spore germination of <i>M. oryzae</i> with [C ₅₀] value of 11.6 μ g/mL, positive control, carbendazim ([C ₅₀] 6.9 μ g/mL)	[50–52]
31	<i>Nodulisporium</i> sp. A21	Leaves of <i>Ginkgo biloba</i> , Nanjing in Jiangsu Province, China	Sporothriolide, (90)	Compounds 84–88 displayed antifungal effects against the spore germination of <i>M. oryzae</i>	[53]
32	<i>Lophodermium nitens</i> DAOM 250227	<i>Pinus strobus</i> Sussex, NB, Canada	Six phenolic bisabolane-type sesquiterpenoids (91–96), pyrenophorin (97)	Compound 89 active against <i>R. niveum</i> , <i>P. ultimum</i> , <i>F. oxysporum</i> , <i>f. sp. cucumerinum</i> , <i>C. oryzae</i> , <i>C. luteae</i> , <i>C. oryzae</i> , <i>P. thrysostachys</i> , <i>S. sclerotiorum</i> and <i>F. fuliginea</i> , especially <i>B. cinerea</i> and <i>F. fujikuroi</i> with EC ₅₀ values below 50–149 μ M.	[54]
33	<i>Erysimum</i> sp.	<i>Acer truncatum</i> Beijing, China.	Esterolide C (98), (12R)-12-hydroxymonocerin (99)	The EC ₅₀ of compound 90 against <i>A. solani</i> was 30.4 μ g/mL (11.6 μ M), while the EC ₅₀ of carbendazim was 18.4 μ g/mL (9.6 μ M).	[55]
34	<i>Bisogniauxia mediterranea</i> EFU38CA	<i>Echinacea purpurea</i> Missouri, USA, Canada.	(\sim)-5-methylinnelen (100) and (\sim)-(3R)-8-hydroxy-6-methoxy-3,5-dimethyl-3,4-dihydroisoquinolin (101)	Compound 97 significantly reduced the growth of <i>M. violaceum</i> and <i>S. cerevisiae</i> at 5 μ M whereas sesquiterpenoids 91–96 active at 50 μ M to both species tested	[56]
35	<i>Hypoxylon subanticulatum</i>	<i>Rubus idaeus</i> collected from Jordan Station, ON, Canada.	Trienylfuranol A (102)	Compounds 98 and 99 active against <i>F. oxysporum</i> , both showing a MIC value of 20 μ g/mL. Amphotericin B (MIC, 0.63 μ g/mL)	[57]
36	<i>Phialophoromuska</i>	<i>Creos sinensis</i> .	Phialomustin C-D (104) (105)	Compound 100 active against <i>P. heveae</i> , <i>P. nificola</i> , and <i>Erysiphe</i> , and caused growth stimulation of <i>C. fageliae</i> , <i>C. luteae</i> , <i>C. quercivorus</i> , and <i>B. cinerea</i> . Compound 101 was found to be slightly more active in the microtiter environment than 5-methylimidazol	[58]
37	unidentified Ascomycete,	<i>Melilotus officinalis</i> .	<i>cis</i> -4-acetoxyisomellitin (106) and 8-deoxy-6-hydroxy- <i>cis</i> -acetoxymellitin (107)	THF 7 (108) inhibited the growth of <i>S. cerevisiae</i> (74 \pm 4 % inhibition) at a concentration of 250 μ g/mL as compared with complete inhibition by nystatin at 10 μ g/mL	[59]
38	<i>Plectophanella</i> sp.		(\sim)Mycothiazin A (108)	Compounds 104–108 active against <i>C. albicans</i> (IC ₅₀ , 14.3 and 73.6 μ M)	[60]
39	<i>Physosphaera</i> sp.		Cytochalasin E (109) and C (110)	Compounds 106 and 107 displayed activities toward <i>M. violaceum</i> , <i>B. cinerea</i> , with 8 mm zone of inhibition for both fungi.	[61]
				Compound 108 active against <i>U. violaceo</i> and <i>E. repens</i> .	
				Compound 109–110 active against <i>E. repens</i> and <i>M. microspora</i>	[61]

Table 1. Cont.

Sr. No.	Fungus	Plantsource	Compounds Isolated	Biological activity *	Ref.s.
40	<i>Critartus monogyna</i> .	Radicinmin (111)	Radicinmin (111) active against <i>E. repens</i> and <i>M. microspora</i>	Radicinmin (111) active against <i>E. repens</i> and <i>M. microspora</i>	[61]
41	<i>Berkelasmium</i> sp.,	<i>Dioscorea zingiberensis</i> , Hubei Province, China.	Deposin ζ (112), Palmarumycin C1 (113), palmarumycin C12 (114), chadospine B (115), palmarumycin C6 (116).	Compounds 112–118 inhibited spore germination of <i>M. oryzae</i> (IC_{50} values in the range 9.1–124.5 μ g/mL). Compound 115 showed the best inhibitory activity (9.1 μ g/mL) among the compounds tested. Carbendazim (IC_{50} 6.3 μ g/mL)	[62]
42	<i>Bipolaris</i> sp. MU34	<i>Gynura hispida</i> Bangkok, Thailand.	Bipolamide B (119)	Bipolamide B (119) active against <i>C. cladosporioides</i> , <i>C. cucumerinum</i> , <i>S. cerevisiae</i> , <i>A. niger</i> and <i>R. oryzae</i> , with MIC values of 16, 32, 64 and 64 μ g/mL, respectively	[63]
43	<i>Alternaria alternata</i> Tche153	Termitaria chedula Rezt, Suanluang Rama IX Public Park, Bangkok, Thailand.	Alternisin (120)	Alternisin (120) in combination with each of three azole drugs, ketoconazole, fluconazole or triconazole at their low sub-inhibitory concentrations exhibited potent synergistic activity against <i>C. albicans</i> with the FIC index range of 0.078 to 0.188	[64]
44	<i>Alternaria</i> sp. UFMGCB 55,	Leaves of <i>Trixis multiflora</i> DC (Asteraceae).	Alternisin (120)	The Alternisin (120) exhibited strong activity against T1 strains <i>P. brasiliensis</i> with MIC values ranging between 1.9 and 312.1 μ g/mL. MIC values found for amphotericin B were between 0.031 and 0.12 μ g/mL. Additionally, 5, pombe cells treated with alternisin were more rounded in shape than untreated cells suggesting alternisin could act through the inhibition of cell wall synthesis or assembly in <i>P. brasiliensis</i> and <i>S. pombe</i>	[65]
45	<i>Cladosprium</i> <i>cladosporioides</i>	Cladosporin (121), Isocladosporin (122)	Cladosporin (121), Isocladosporin (122)	At 30 μ M compound 121 exhibited 92.7, 90.1, 95.4, and 79.9% growth inhibition against <i>C. acutatum</i> , <i>C. fagacearum</i> , <i>C. geotrichoides</i> and <i>A. nivalis</i> . Compound 122 showed 50.4, 60.2, and 83.0% growth inhibition at 30 μ M against <i>C. fagacearum</i> , <i>C. geotrichoides</i> , and <i>P. viticola</i> , respectively	[66]
46	<i>Epicoccum</i> sp. CAFTBO Olu 19B	<i>Theobroma cacao</i> (Sterculiaceae) Mount Kala, near Yaoundé, Centro Province, Republic of Cameroon	Epicoccolone (123), Epicoccolide A (124) and B (125)	Compounds 123–125 showed inhibitory effects on the mycelial growth of <i>P. ultimum</i> and <i>A. confalidans</i> and <i>R. solani</i> (MIC in the range of 20–80 μ g per paper disc)	[67]
47	<i>Biscogniauxia mediterranea</i>	<i>Opuntia humifusa</i> (Cactaceae) from the United States	5-methylmelokin (100)	Compound 100-5-methylmelokin was evaluated for antifungal activity against seven plant pathogens (<i>C. acutatum</i> , <i>C. fagacearum</i> , <i>C. geotrichoides</i> , <i>F. oxysporum</i> , <i>F. cincinna</i> , <i>B. dothidea</i> , <i>ana</i> , <i>A. fagaeae</i> , <i>C. gloeosporioides</i> , <i>F. oxysporum</i> , <i>F. tritici</i>) and <i>P. viticola</i> using an in vitro microdilution broth assay.	[68]
48	<i>Emerita</i> sp. XLD29	Leaves of <i>Panax notoginseng</i> Shijiazhuang, Hebei Province, China.	5,(undeca-3',7'-triene-1',3')furan-2-ol (126) and 5,(undeca-3',5',7'-triene-1',3')furan-2-carbonate (127)	Compound 126 active against <i>R. solani</i> , <i>V. dahliae</i> , <i>H. magnis</i> , <i>F. oxysporum</i> , <i>F. tritici</i> , <i>B. dothidea</i> , <i>ana</i> , <i>A. fagaeae</i> , <i>C. gloeosporioides</i> , <i>F. oxysporum</i> , <i>F. cincinna</i> , <i>B. dothidea</i> , <i>ana</i> , <i>A. fagaeae</i> (MIC values from 25 to 31.1 μ g/mL), while compound 127 displayed activity against <i>V. dahliae</i> , <i>H. magnis</i> , <i>F. tritici</i> , <i>B. dothidea</i> , <i>ana</i> , <i>A. fagaeae</i> (MIC values from 50 to 12.5 μ g/mL)	[69]
Compounds Produced by Hyphomycetes					
49	<i>Fusarium fujikumii</i> (WF5), <i>Poeciliella dryogenes</i> WF7, W16 and <i>P. expersum</i> WF7	Finger millet Plants Arkell Field Station, Arkell, ON, Canada.	5-hydroxy 2(3H)-benzofuranone (128), dehydrocostus lactone (129) and harpagoside (130)	Compounds 128–130 active against <i>F. gramineum</i> with MIC of 31.25, 250.00 and 31.25 μ g/mL, respectively.	[70]
50	<i>Trichoderma koningiopsis</i> YIM PH3002	<i>Panax notoginseng</i> , Wenshan, Yunnan Province, China.	Koniginin O (131), koniginin Q (132), 2-O-methylkoniginin D (133)	Compounds 131–132 active against <i>F. oxysporum</i> and <i>P. cinnamomea</i> , with an MIC of 128 μ g/mL. Compound 133 showed activity against <i>F. cinnamomea</i> with an MIC of 128 μ g/mL. Nystatin was active with MICs at 32 μ g/mL.	[71]
51	<i>Trichoderma koningiopsis</i> YIM PH3002	<i>Panax notoginseng</i> , Wenshan, Yunnan Province, China.	Koniginopin C (134)	Compound 134 exhibited in vitro antifungal activity against <i>F. oxysporum</i> , <i>A. paras</i> , <i>F. solani</i> and <i>P. cinnamomea</i> with MICs at 32, 64, 32, and 16 μ g/mL, respectively	[72]

Table 1. Cont.

Sr. No.	Fungus	Plantsource	Compounds Isolated	Biological activity *	Ref.s
52	<i>Trichoderma</i> sp. 09	<i>Mycoporum bentivides</i>	Dichlorodipropionate (135), dichlorodiaporthin (136)	Compounds 135–136 active against <i>C. musae</i> and <i>Rhizoctonafuscani</i> (MIC values from 6.25 to 50 µg/ml.)	[73]
53	<i>Trichoderma brevicompactum</i> 0248	<i>Allium sativum</i>	Trichodermin (137)	Compound 137 active against <i>R. solani</i> , <i>B. cinereol</i> , <i>C. lindemuthianum</i> with an MIC of 0.25, 2.02 and 25.0 µg/ml., respectively.	[74]
54	<i>Trichoderma boningiiopsis</i> YIM PHB0002	Wenshan, Yunnan Province of China.	Koninginin R (138) and S (139)	Carbofuranin showed antifungal activity against <i>R. solani</i> , <i>B. cinereol</i> , with an EC ₅₀ of 0.36 and 10.35 µg/ml., respectively.	[75]
55	<i>Aspergillus terreus</i>	<i>Carthamus lanatus</i> Al-Azhar University campus, Assuit Branch, Assuit, Egypt	(2Z,2E)-stigmasta-5,7(22)-triene-3-β-ol (140), aspernolides F (141)	Compound 138 active against <i>F. oxysporum</i> and <i>F. flaeciferum</i> with MICs at 128 µg/ml., while compound 139 displayed activity against <i>F. oxysporum</i> with MIC at 128 µg/ml.	[76]
56	<i>Aspergillus</i> sp. KJ-9,	<i>Melia azedarach</i> which was collected at Yangling, Shaanxi Province, China	Fonecinecone A (142), (R)-3-hydroxybutanoneitrile (143)	Compound 140 active against <i>G. strobiloides</i> , <i>M. siseia</i> , <i>B. cinerea</i> , <i>C. gloeosporioides</i> and <i>A. solani</i> with MIC range of 25–50 µM	[77]
57	<i>Aspergillus</i> sp.	Gloriosa superba Tripathi, India.	6-methyl-1,2,3-trihydroxy-7S-cyclohepta-9,12-diene-1-one-5,6Z,8-teralene-7-acetamide (KL-4) (144)	KL-4 (144) active against <i>S. cerevisiae</i> , <i>C. utilisans</i> and <i>C. gastricus</i> with MIC 25, 12.5, and 50 µg/ml., respectively.	[78]
58	<i>Penicillium</i> sp. R22	<i>Nerium indicum</i> collected from Qilong Mountain, Shanxi Province, China.	5-hydroxy-3-methoxy-4-phenylisoquinolin-1(2H)-one (145), 3-O-methylviridicatin (146) and viridicatin (147)	Compound 145 active against <i>A. brasiliense</i> , <i>A. nivalis</i> with MIC value of 31.2 µg/ml., compound 146 against <i>A. brasiliense</i> , <i>B. cinerea</i> and <i>V. male</i> and <i>B. cinerea</i> with MIC value of 31.2 µg/ml.	[79]
59	<i>Penicillium naticiborskii</i>	<i>Rhododendron lametensum</i> were collected at the test site of University of Oulu, Finland.	Cutovirin C (148)	Outovirin C (148) inhibited growth of <i>F. oxysporum</i> , <i>B. cinerea</i> , and <i>V. dahliae</i> at the concentration of 0.38 µM. Compound 148 is active against <i>V. dahliae</i> (5% inhibition) and slightly less effective against <i>B. cinerea</i> (5% inhibition).	[80]
60	<i>Fusarium</i> sp.	<i>Mentha longifolia</i> Saudi Arabia.	Fusirupeptide A (149)	Compound 149 active against <i>C. utilisans</i> , <i>C. gloeosporioides</i> , <i>C. krestae</i> , and <i>A. fumigatus</i> with IC ₅₀ values of 0.1, 0.124, 0.1, and 0.14 µM, respectively.	[81]
61	<i>Fusarium chlamydosporium</i>	<i>Arvilia gencinii</i> Al Madinah Al Munawwarah, Saudi Arabia.	Fusarithioamide A (150)	Compound 150 active with inhibition zone-diameters 1.62 mm and MIC 2.6 µg/ml. towards <i>C. utilisans</i> . Clotrimazole (inhibition zone-diameters 1.85 mm and MIC 3.7 µg/ml.)	[82]
62	<i>Fusarium</i> sp.	Fjella caricae Qilong Mountain, Shaanxi Province, China	Hebolic acid Methyl ester (151), hebolic acid (152) and hydrohebolic acid (153)	Compounds 151–153 active against <i>B. cinerea</i> , <i>C. gloeosporioides</i> , <i>F. oxysporum</i> , <i>F. graminearum</i> and <i>P. capsici</i> (MIC in the range of 12.5–25 µg/ml.), Carbendazim (MIC in the range of 32.2–62.5 µg/ml.)	[83]
63	<i>Fusarium</i> sp.		Colleopterin B (154), colleochochlorin B (155), 1L-Z1272β (illucicolin B) (156) and 4,5-dihydrodeltachlorochlorin (157)	Compounds 154–156 showed antifungal against <i>U. radicans</i> and <i>F. oxysporum</i> . Compound 157 showed antifungal activity towards <i>E. repens</i> .	[84]
64	<i>Curvularia</i> sp., strain M12.	<i>Murraya koenigii</i> Rajshahi University, Bangladesh	Murranolide A (158), murranolide B (159), Curvalin (160), (S)-dehydrocurvalin (161), Pyrenolide A (162), modiolide A (163), and 8-hydroxy-6-enethoxy-3-methylisocoumarin (164)	Pyrenolide A (162) showed a strong motility impairing activity against <i>Phytophthora capsici</i> zoospores at a low concentration (100%, at 5.14 µg/ml.) in a short-time (30 min). Compounds 158–161 and 163–164 exhibited zoospore motility impairment activity at higher concentrations (IC ₅₀ : 50–100 µg/ml)	[85]

Table 1. Cont.

Sr. No.	Fungus	Plant source	Compounds isolated	Biological activity *	Ref.s
65	<i>Phaeacrenatum</i> sp.,	<i>Senna spectabilis</i> , Aracuan-Cerrado area, Ataraqueta, São Paulo State, Brazil.	Isoaligalone B (165), and C (166), aigalone (167)	Compounds 165 and 167 exhibited antifungal activity, with a detection limit of 5 µg, for <i>C. falagrioides</i> and <i>C. sphaerogenum</i> , compound 166 exhibited weak activity (detection limit > 5 µg), with a detection limit of 25 µg. Nystatin, positive control showing a detection limit of 1 µg	[86]
66	<i>Trichothecium</i> sp.	<i>Phyllanthus amarus</i> Pune India	Trichothecinol A (168)	Compound 168 active against <i>C. albidus</i> up to 20 µg/mL	[87]
67	<i>Trichothecium</i> sp.	<i>Phyllanthus</i> sp., Pune India.	Trichothecin (169)	Trichothecin (169) active against <i>S. cerevisiae</i> , <i>C. albicans</i> var <i>diffluentes</i> (NCIM 3372), <i>F. oxysporum</i> , <i>P. expansum</i> , <i>T. viride</i> , <i>P. oryzae</i> and <i>A. niger</i> with MIC of 6.0, 20.0, 12.0, 100, 30.0, 40.0, 20.0 and 12.0 µg/mL, respectively	[88]
Compounds Produced by Basidiomycetes					
68	<i>Scleroderma</i> UFSM Sc (Persoon) Fries	<i>Eucalyptus gunnii</i> .	Sclerodol A (170) and B (171) and related lanostane triterpenoid (172)	Compound 170 active against <i>C. albinus</i> , <i>C. ketti</i> , <i>C. krusei</i> , <i>C. parapsoriasis</i> (MIC of 25.0, 25.0, 6.25 and 12.5 µM/C 25.0, 25.0, 12.5, 25.0 /48 h). Compound 171 and 172 were active against test strain (MIC in the range of 12.5–100 µg/mL). Nystatin active against test strains (MIC in the range of 0.77–1.52 µg/mL).	[89]

3. Antifungal Potential of Volatile Organic Compounds (VOCs) from Endophytic Fungi

Volatile organic compounds (VOCs) are generally carbon compounds which exist in the gaseous phase at normal/ambient temperatures and pressures. Over 250 different VOCs produced by fungi comprising different chemical classes such as aldehydes, ketones, alcohols, phenols, thioesters, and so forth, have been identified in the context of the deterioration of fruits, vegetables, indoor environments (sick building syndrome); as chemotaxonomic markers; and in the morphogenesis and development of fungi.

However, bioprospecting fungal endophytes for the production of volatile antimicrobials came into the limelight with the discovery of *Muscodor albus* from the plant *Cinnamomum zeylanicum*, from Honduras. *M. albus* was found to produce an admixture of VOCs which could effectively kill a variety of pathogenic bacteria and fungi associated with plants and animals. This research garnered much attention and drove people to explore the volatile antibiotic properties of endophytic fungi for varied applications [90,91].

The genus *Muscodor* comprises of an endophytic fungi which is predominantly sterile, does not possess true reproductive structures like other fungi, and emanates a characteristic smell which is largely attribute to its VOC composition [92]. Since the report of *M. albus* in the late 1990s, to date, 20 species have been added to this genus, which have largely been identified based on their volatile signatures, molecular phylogeny, and morphological characteristics (Table 2). The characteristic VOC profile, therefore, is helpful in delineating the species, as well as playing a significant role in its anti-fungal and anti-bacterial properties. In this section, we only be highlight the anti-fungal potential of VOCs produced by these endophytic fungi.

The majority of the VOCs produced by the endophytic fungi comprises of a mixture of volatile components which generally has either a synergistic effect or an additive effect that enhances their bioactivity against pathogenic microbes. However, in a couple of studies, the major components of the volatile mixture were independently evaluated to understand their true antimicrobial/anti-fungal potential. These are generally synthetically generated and converted into a volatile form and subsequently evaluated for their bioactivity against the test microorganisms. For instance, *Sclerotina sclerotiorum* was completely inhibited by 2-methyl-1-butanol and 3-methyl-1-butanol with an EC₅₀ value of 0.8 µL/mL. 2-methyl-1-butanol also inhibited *Penicillium digitatum* with an EC₅₀ value of 0.48 µL/mL and *B. cinerea* with a value of 1.38 µL/mL. However, the volatile admixture of the *M. albus* VOC exhibited an IC₅₀ range between 0.08 and 1.13 µL/mL, which clearly confirms the hypothesis of the synergistic/additive effects of the volatile components [93].

Recently, ethyl acetate has been reported to be the main VOC of yeasts *Wickerhamomyces anomalus*, *Metschnikowia pulcherrima*, and *Saccharomyces cerevisiae*, which inhibit the decay causing mold, as well as *B. cinerea*. All three yeasts exhibit excellent biological control properties and were used for checking the mold and pathogenic attack in sweet cherries and strawberries. *W. anomalus* induced the highest killing activity amongst the three which was attributed to the higher production of Ethyl acetate. The role of the ethyl acetate was re-affirmed by using synthetic ethyl acetate from strawberry fruits to affirm the anti-fungal action [94].

Similarly, *Phaeosphaeria nodorum*, which existed as an endophyte in plum leaves (*Prunus domestica*) was found to inhibit the pathogen *Monilinia fruticola*. The major component of the VOC produced by *Phaeosphaeran odoratum* comprised of 3-methyl-1-butanol, acetic acid, 2-propyn-1-ol, and 2-propenenitril [95]. Similarly, six VOCs from the endophytic fungus *Hypoxyylon anthochroum* (that is, phenylethyl alcohol), 2-methyl-butanol and 3-methyl-1-butanol, eucalyptol, ocimene, and terpenoline were tested against *Fusarium oxysporum*. The results indicated that these compounds exhibited concentration-dependent anti-fungal activity individually but have better action and control synergistically. Thus, the mixture of six VOCs may be used for the control of *Fusarium oxysporum* in tomatoes [96].

The genus *Muscodor* is one of the best studied endophytic fungus which produces a synergistic mixture of VOCs having lethal effects against a wide variety of plant and human pathogenic fungi, nematodes, and bacteria as well as certain insects [97–100]. The volatility of the *Muscodor* species

has been used to replace methyl bromide (MeBr)—a traditional soil fumigant—which has been globally banned as it causes the depletion of ozone layer. Different species of *Muscodor*, their major VOCs, and their anti-fungal spectrum are given in Table 2. Geographically, each *Muscodor* species has a characteristic signature volatility. For instance, the Indian *Muscodor* species invariably has 4-Octadecylmorpholine as a marker compound while 2-methyl propanoic acid is generally found in *Muscodor* isolated from North and South America.

The majority of the VOCs from the fungal endophytic fungi are used as biological control agents to prevent the fungal deterioration of crops, fruits, and vegetable, under both pre- and post-harvest conditions. However, the exploitation of these fungally volatile organic compounds (FVOCs) from endophytic fungi are not being actively applied to humans for the prevention of fungal infections.

There exists a huge scope in evaluating these FVOCs from endophytic fungi since they could be helpful in curing superficial skin infections, the sanitization of public toilets, and in night soil. They can also find applications in personal care products such as for the aroma/fragrance in deodorants and sprays. They could presumably be helpful in the development of sprays for inhalation to treat fungal diseases like Aspergillosis in lungs.

Table 2. Spectrum of VOCs emitted by *Muscodorus* species and their anti-fungal activity predominantly against plant pathogenic fungi.

No.	Name of the Endophytic Fungi	Geographic Area of Isolation	Major VOCs Produced	Anti-Fungal Activity	Refs.
1	<i>Muscodorus albus</i>	Central America (Honduras)	2-methylpropanoic acid; 2-methyl-1-butanol; ethanol; acetic acid (methyl ester)	<i>Rhizoctonia solani</i> ; <i>Phytophthora cinnamomi</i> ; <i>Sclerotinia sclerotiorum</i> ; <i>Fusarium solani</i> ; <i>Verticillium dahliae</i>	[91]
2	<i>M. vitigenus</i>	South America (Peru)	naphthalene; caryophyllene; azulene	<i>R. solani</i> ; <i>Phoma</i> sp.; <i>C. coifficola</i>	[98]
3	<i>M. roseus</i>	Australia	2-hydroxyacrylic acid (ethyl ester); 1,2,4-tri-methyl-benzene; 2-nonalene	Antifungal spectrum not reported	[101]
4	<i>M. yunnanensis</i>	South America (Mexico)	caryophyllene; aromadendrene	<i>Botryotis cinerea</i> ; <i>R. solani</i> ; <i>C. coifficola</i> ; <i>Phoma</i> sp.	[102]
5	<i>M. fenggangensis</i>	China	2-methylpropanoic acid; β-phellandrene	<i>B. cinerea</i> ; <i>Aspergillus clavatus</i> ; <i>Colletotrichum fragariae</i> ; <i>Sclerotium rolfsii</i>	[90]
6	<i>M. crispa</i>	South America (Bolivia)	2-methylpropanoic acid; ethanol; ethyl acetate	<i>B. cinerea</i> ; <i>Curvularia lunata</i> ; <i>P. cinnamomi</i> ; <i>S. sclerotiorum</i>	[103]
7	<i>M. sitara</i>	USA (Columbia)	butylated hydroxytoluene; octaceanoic acid; thujopsene; 2-methylpropanoic acid; naphthalene	<i>Aspergillus fungatus</i> ; <i>Colletotrichum lagunarium</i> ; <i>B. cinerea</i> ; <i>Cercospora beticola</i> ; <i>Phytophthora palmivora</i> ; <i>Fusarium solani</i>	[104]
8	<i>M. mungae</i>	Thailand	3-methylbutanol acetate	<i>Alternaria porri</i> ; <i>Alternaria solani</i> ; <i>Colletotrichum gloeosporioides</i> ; <i>Nigrospora oryzae</i>	[105]
9	<i>M. oryzae</i>	Thailand	2-methylpropanoic acid	<i>A. porri</i> ; <i>A. solani</i> ; <i>Aspergillus flavus</i> ; <i>B. cinerea</i> ; <i>C. gloeosporioides</i> ; <i>N. oryzae</i>	[105]
10	<i>M. suthepensis</i>	Thailand	3-methylpropanoic acid	<i>A. porri</i> ; <i>Alternaria alternata</i> ; <i>Aspergillus flavus</i> ; <i>B. cinerea</i> ; <i>C. gloeosporioides</i> ; <i>Fusarium oxysporum</i> ; <i>Fusarium solani</i> ; <i>N. oryzae</i>	[105]
11	<i>M. equiseti</i>	Thailand	3-methylbutan-1-ol; 3-methylbutanoyl acetate; 2-methylpropanoic acid	<i>A. porri</i> ; <i>A. solani</i> ; <i>B. cinerea</i> ; <i>C. gloeosporioides</i> ; <i>F. oxysporum</i> ; <i>F. solani</i> ; <i>N. oryzae</i>	[105]
12	<i>M. cinnamomi</i>	Thailand	2-methylpropanoic acid; 2-methyl butanoic acid; azulene	<i>Rhizoctonia solani</i>	[105]
13	<i>M. kashiyam</i>	India	1-methyl-4-(1-methylethylidene)-cyclohexane; 2-(4-anorpholinyl)ethylamine; 9-octadecanoic acid (methyl ester); 4-octadecylmorpholine	<i>Bianertia ochroleuca</i> ; <i>Cercospora beticola</i> ; <i>Chaitomium heteroporan</i> ; <i>C. gloeosporioides</i> ; <i>F. oxysporum</i> ; <i>Fusarium equiseti</i> ; <i>Curvularia lunata</i>	[106]
14	<i>M. darjeelingensis</i>	India	4-octadecylmorpholine; 2,6-di(1-dimethyllethyl)-4-(1-oxopropyl)phenol; beta-aminoethyl-morpholine	<i>Lasiobolus phaeobromae</i> ; <i>A. alternata</i> ; <i>Rizoctonia solani</i> ; <i>Cercospora beticola</i>	[107]
15	<i>M. strobelii</i>	India	4-octadecylmorpholine; tetraoxeprolan; aspidofractinane-3-methanol; viridiflorol	<i>Rhizoctonia solani</i> ; <i>Colletotrichum gloeosporioides</i> ; <i>Fusarium oxysporum</i> ; <i>Lasiobolus phaeobromae</i>	[108]

Table 2. Cont.

No.	Name of the Endophytic Fungi	Geographic Area of Isolation	Major VOCs Produced	Anti-Fungal Activity	Ref.s.
15	<i>M. strobelii</i>	India	4-octadecylmorpholine; tetraoxeprolane; aspidofractamine-3-methanol; viridiflorol	<i>Rhizoctonia solani</i> ; <i>Colletotrichum gaeumannoides</i> ; <i>Fusarium oxysporum</i> ; <i>Lasiodiplodia theobromae</i>	[108]
16	<i>M. ligrii</i>	India	4-octadecylmorpholine; 1-tetradecanamine; n,n-dimethyl 12-benzidicarboxylic acid mono(2-ethylhexoxy)ester	<i>Alternaria alternata</i> <i>Cercospora beticola</i>	[109]
17	<i>M. heiae</i>	Thailand	2-phenylethanol; azulene;	<i>Aspergillus niger</i> ; <i>Phellinus noxius</i> ; <i>Rigidoporus microporus</i>	[110]
18	<i>M. shoenensis</i>	India	n,n-dimethyl-1-pentadecamine;	<i>Cercospora beticola</i>	[111]
19	<i>M. indica</i>	India	n,n-dimethyl-1-pentadecamine; 4-morpholinorethanamine;	<i>Cercospora beticola</i> <i>Penicillium marneffei</i>	[111]
20	<i>M. camphora</i>	India	tetracontane; 4-octadecylmorpholine; n,n-dimethyl-1-pentadecamine	<i>Colletotrichum gaeumannoides</i> ; <i>Lasiodiplodia theobromae</i>	[112]

4. Methods Used for Activation of Silent Biosynthetic Genes

Several research studies confirm that most of the biosynthetic gene clusters are observed to be silent or expressed at a low (minimal) level upon employing conventional culturing conditions for growth/propagation of microorganisms [113]. To activate such silent biosynthetic genes, numerous strategies have been employed, such as the one strain many compounds (OSMAC) approach (activation mediated through modification in composition of medium, aeration, temperature or shape of culturing flask), co-culturing method (facilitating activation through interspecies crosstalk) and genomics based approaches (expression of orphan biosynthesis genes in a heterologous host). In recent times, the use of chemicals as modifiers to alter the epigenetic makeup/constitution of a microorganism to improve its biosynthetic potential has become a beneficial tool. The method uses a chemical that acts as DNA methyltransferase inhibitors (DNMTi) or histone deacetylase inhibitors (HDACi), thereby stimulating the transcription previously silent gene clusters and fostering the production of a spectrum of natural products. A comprehensive description of some of these methods are given below/highlighted in the subsequent section.

4.1. Epigenetic Modification

Endophytes have proven to be the prolific source of bioactive metabolites and offer a substitute and untapped reserve for the discovery of novel metabolites. Studies have led to findings that tell biosynthetic gene clusters of microorganisms are mostly silent or expressed at very low levels under standard culture conditions and are least expressed, but under stress condition may it be biological, chemical or physical their expression takes place. Epigenetic modulators lead to the expression of these silent or cryptic genes. Epigenetic gene regulation is mediated by covalent histone modification, DNA methylation chromatin modeling basically induced by DNA methyl transferase inhibitors such as 5-aza-2-deoxycytidine, 5-azacytidine, hydralazine, procaine and histone deacetylase [114]. Chromatic modification in fungi to enhance gene transcription has led to secondary metabolite production of anthraquinones, cladochromes, lunalides, mycotoxins, and nygerones [115]. Structural genes that control transcriptional factor regulates the synthesis of secondary metabolites in fungi, these genes mediate factors occupied in environmental signals like pH, nitrogen and carbon sources, temperature, light, etc. [116]. In lab condition, these gene clusters are mostly silent. Under which natural conditions these clusters become activated is still unexplained. As per genetic sequencing studies carried so far, it is estimated that the clusters of genes responsible for secondary metabolites have not yet been deciphered completely [117].

From endophytic fungi *Aspergillus fumigatus* (GA-L7) obtained from *Grewia asiatica* led to identification of seven metabolites namely pseurotin A (173), pseurotin D (174), pseurotin F2 (175), fumagillin (176), tryprostatin C (177), gliotoxin (178) and bis(methylthio)gliotoxin (179) (Figure 9). On addition of the valproic acid, increase in the production of fumiquinazoline C (180) up to 10 times was noticed along with a shift in the pattern of metabolite production. It was also observed that all the genes, i.e., Afua_6g 12040, Afua_6g 12050, Afua_6g 12060, Afua_6g 12070 and Afua_6g 12080, tangled in the biosynthesis of fumiquinazoline C (180), were upregulated significantly by 7.5, 8.8, 3.4, 5.6 and 2.1 folds, respectively [118].

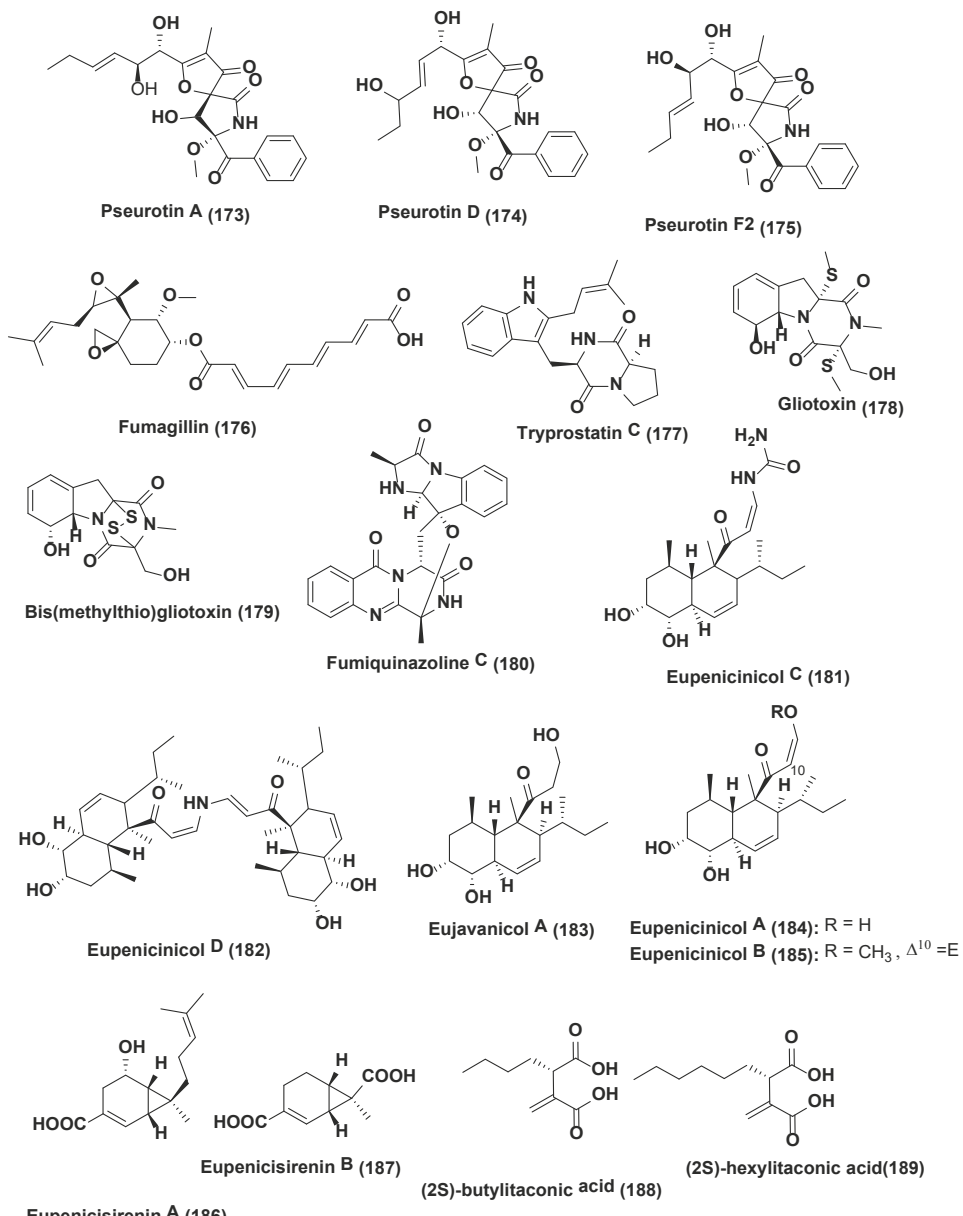


Figure 9. Structures of metabolites isolated from Epigenetic modification in endophytic fungi (173–189).

The NAD+-dependent HDAC inhibitor, nicotinamide, enhanced the yield of eupenicinicol C (181), and D (182) (Figure 9), decalin containing metabolites together with eujavanicol A (183), and eupenicinicol A (184) (Figure 9), biosynthetically related compounds by endophytic *Eupenicillium* sp. LG41, identified from the *Xanthium sibiricum*, a Chinese medicinal plant [119]. Under a similar condition without HDAC inhibitor, nicotinamide *Eupenicillium* sp. LG41 produce different decalin-containing compounds:

eupenicinolics A (184), and B (185); two new sirenin derivatives, eupenicisirenins A (186) and B (187); and other four known compounds, (2S)-butylitaconic acid (188), (2S)-hexylitaconic acid (189) (Figure 9), xanthomegnin (190), and viridicatumtoxin (191) (Figure 10) [120].

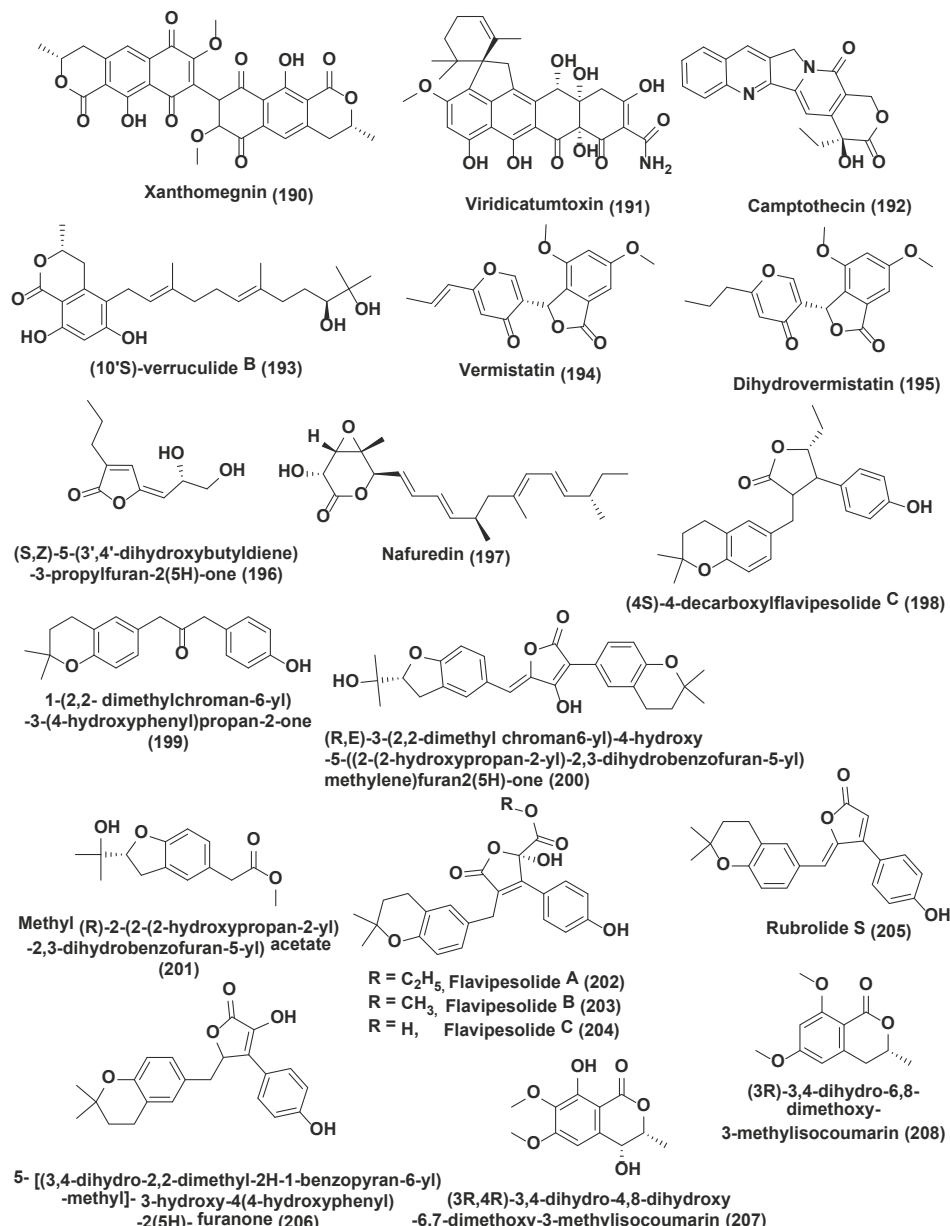


Figure 10. Structures of metabolites obtained from Epigenetic modification in endophytic fungi (190–208).

Vasanthakumari et al. [121] reported the attenuation of camptothecin (192) (Figure 10) yield in endophytic fungi obtained from camptothecin fabricating plants, *Nothapodytes nimmoniana* and *Miquelia*

dentata incorporated with 5-azacytidine, a DNA methyltransferase inhibitor. It was indicated that the mode behind decrease of camptothecin yield in endophytic fungi could in principle be inverted by stimulating some signals from the tissue of the plant, probably the methylation or silencing of the genes liable for camptothecin production.

Metabolites such as (10'S)-verruculide B (193), vermistatin (194), and dihydrovermistonatin (195) (Figure 10) were identified owing to the supplementation of HDAC inhibitor, suberoylanilide hydroxamic acid (SAHA) to endophytic fungi culture *Phoma* sp. nov. LG0217 isolated from *Parkinsonia microphylla*. However, in absence of SAHA, a novel metabolite (S,Z)-5-(3',4'-dihydroxybutyldiene)-3-propylfuran-2(5H)-one (196), along with nafturedin (197) (Figure 10), was produced [122].

Four new meroterpenoids identified as (4S)-4-decarboxylflavipesolide C (198), 1-(2,2-dimethylchroman-6-yl)-3-(4-hydroxyphenyl)propan-2-one (199), (R,E)-3-(2,2-dimethyl chroman-6-yl)-4-hydroxy-5-((2-(2-hydroxypropan-2-yl)-2,3-dihydrobenzofuran-5-yl)methylene)furan2(5H)-one (200), methyl (R)-2-(2-(2-hydroxypropan-2-yl)-2,3-dihydrobenzofuran-5-yl) acetate (201), along with nine known compounds flavipesolides A–C (202–204), rubrolide S (205), 5-[(3,4-dihydro-2,2-dimethyl-2H-1-benzopyran-6-yl)-methyl]-3-hydroxy-4(4-hydroxyphenyl)-2(5H)-furanone (206), (3R,4R)-3,4-dihydro-4,8-dihydroxy-6,7-dimethoxy-3-methylisocoumarin (207), (3R)-3,4-dihydro-6,8-dimethoxy-3-methylisocoumarin (208) (Figure 10), terretonin C (209), and ergosterol (210) (Figure 11) were obtained using chemically modified epigenetic culture of *Aspergillus terreus* OUCMDZ-2739 with 10 μM trichostatin A (TSA). Under the similar parameters without TSA, *A. terreus* OUCMDZ-2739 yielded many compounds, i.e., aspernolide B (211), butyrolactone II (212), butyrolactone IV (213), butyrolactone I (214), aspernolide A (215), asterrelenin (216) and (+)-terrein (217) (Figure 11), supporting that fungal metabolite enrichment and chemodiversity using epigenetic modifiers can be done to obtain new products [123]. Induction of isosulochrin (218) (Figure 11) was also witnessed when *Chaetomium* sp. was supplemented with 5-azacytidine or SAHA on solid rice medium [124].

Asai et al. [125] reported six new benzophenones, cephalanones A–F (219–224), and 2-(2,6-dihydroxy-4-methylbenzoyl)-6-hydroxybenzoic acid (225) (Figure 11) from culture of *Graphiopsis chlorocephala*, from *Paeonia lactiflora* in presence of HDAC inhibitors nicotinamide (10 μM) which resulted in significant increase in secondary metabolite production.

To express silent biosynthetic pathways, molecules such as HDAC and DNMT are used to enhance the fungal metabolites production. Different studies suggest an increase in chemical diversity of metabolites by induction with these epigenetic modifiers. For growth and acclimatization with the environment fungus are known to produce diverse secondary metabolites. Cross talk between microbes and plant lead to the expression of these pathways which stays silent in *in vitro* conditions. Metabolic profiles shift led by SMs induced modifier is due to expression of cryptic genes [126].

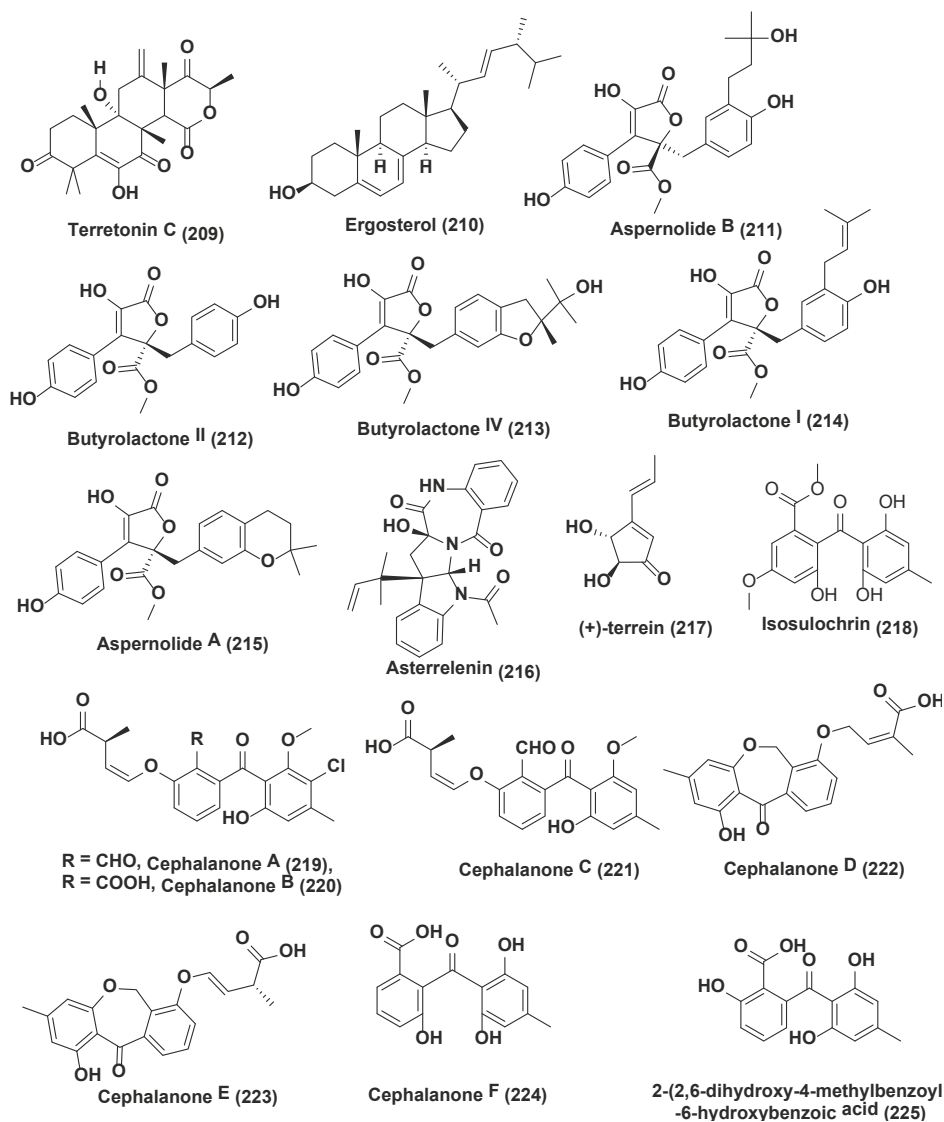


Figure 11. Structures of metabolites obtained from Epigenetic modification in endophytic fungi (209–225).

4.2. The Co-Culture Strategy

Interspecific interaction among different species leads to evolution and biodiversity, organism combines their genetic information for better adaptability. The cohabitation of different microorganisms that share similar niches competes with growth, morphology, adaptation, and development patterns [127,128]. The increased productions of metabolites in co-culture which are not produced in axenic culture are the result of competition or antagonism faced by the microorganism that leads to activation of cryptic genes. [129]. Co-cultivation is a way to provide natural habitat to fungi so that gene clusters become activated. In *Aspergillus nidulans*, the cryptic gene has been successfully activated leading to isolation of novel compounds [130].

In a study carried by Ola et al. [131], accumulation of secondary metabolites, i.e., lateropyrone (226), cyclic depsipeptides of the enniatin type (227–229), and the lipopeptide fusaristatin A (230) (Figure 12), was found to be enhanced by 78 folds by co-culturing *B. subtilis* 168 trpC2 with *Fusarium tricinctum* isolated from the *Aristolochia paucinervis*. This led to the identification of three new compounds macrocarpon C (231), 2-(carboxymethylamino)benzoic acid (232) and (−)-citreoisocoumarin (233), and a known compound, (−)-citreoisocoumarin (234), which was absent in axenic culture of bacterial or fungal control. On coculturing *Alternaria* sp. and *Phomopsis* sp. there was the enhancement of taxane (235) production by eight fold [132].

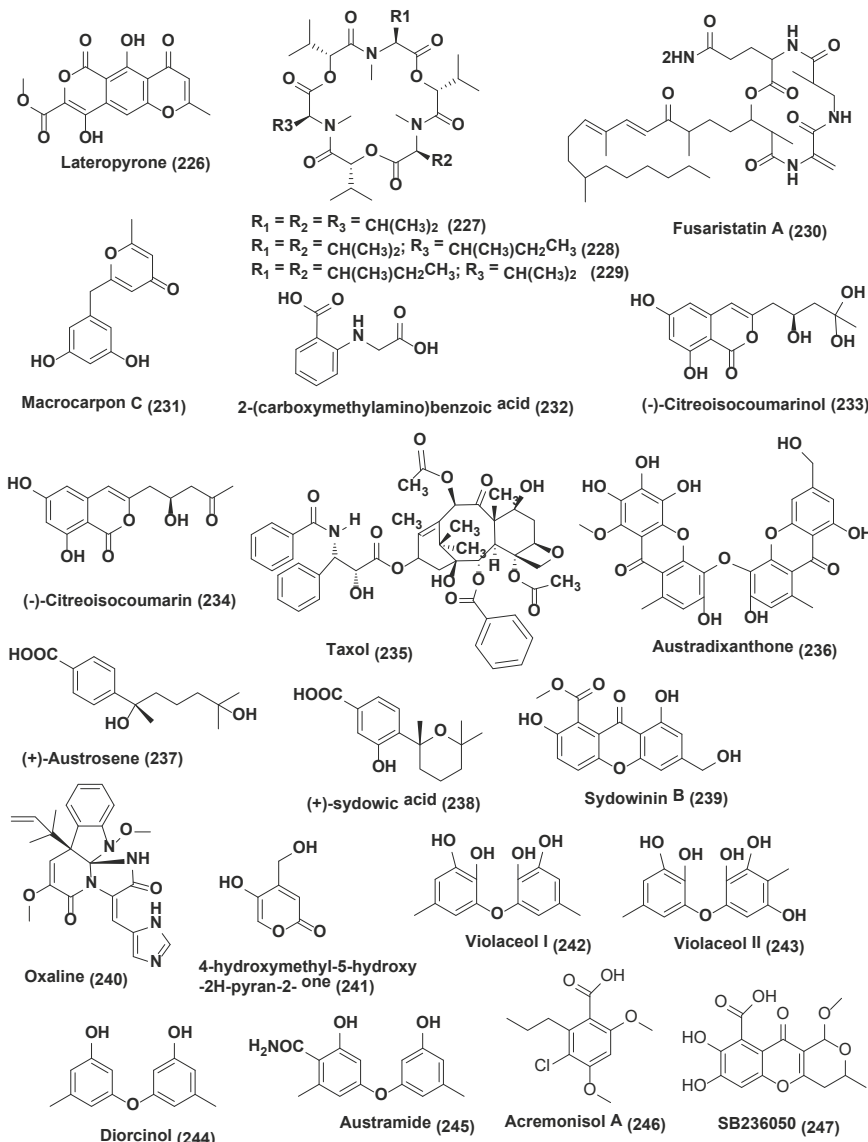


Figure 12. Structures of metabolites obtained from co-culture in endophytic fungi (226–247).

Aspergillus austroafricanus endophyte residing inside the leaves of *Eichhornia crassipes* was the source of two new metabolites, namely, xanthone dimer austradixanthone (236) and sesquiterpene (+)-austrosene (237), and five known compounds, (+)-sydowic acid (238), sydowinin B (239), oxaline (240), 4-hydroxymethyl-5-hydroxy-2H-pyran-2-one (241) (Figure 12), ergosterol (210) (Figure 13). However, the same endophyte grown in mixed cultures with *Bacillus subtilis* or *Streptomyces lividans* led to the identification of many diphenyl ethers i.e., violaceol I (242), violaceol II (243), and diorcinol (244) (Figure 12) along with new austramide (245) (Figure 12), increased up to 29 times [133].

Chaetomium sp. was isolated from *Sapium ellipticum* the Cameroonian medicinal plant. When *Chaetomium* sp. was cultured axenically on solid rice medium, average yields per culture flask were 2.8, 13.9, 132.7 and 14.6 mg of acremonisol A (246), SB236050 (247) (Figure 12), and SB238569 (248), respectively, and 1:1 mixture of 3- and 4-hydroxybenzoic acid methyl esters (249–250), respectively, (Figure 13) was observed. When Co-cultivation of *Chaetomium* sp. was undertaken with viable or autoclaved cultures of *Bacillus subtilis* there was a strong accumulation of the 1:1 mixture of (249), and (250), was observed, accounting for an 8.3 and 7.4-fold increase, respectively, compared to axenic fungal controls in both cases. SB236050 (247) and SB238569 (248), two major polyketides of *Chaetomium* sp., were not detected in co-cultures. Five new compounds, Shikimeran A (251), Bipherin A (252), Chorismeron (253), Quinomeran (254), and Serkydayn (255), and two known compounds, isosulochrin (218) and protocatechuic acid methyl ester (256) (Figure 12), were only detected in co-cultures of *Chaetomium* sp. with viable or autoclaved *B. subtilis* cultures, but were lacking in both fungal or bacterial controls when cultured axenically [124].

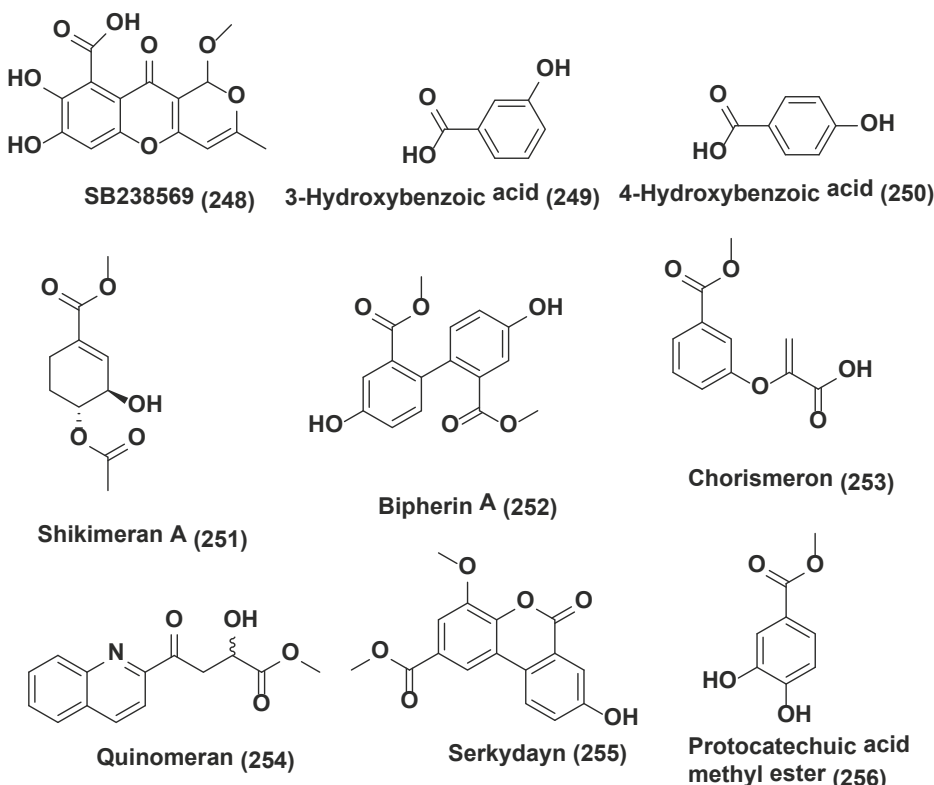


Figure 13. Structures of metabolites obtained from co-culture in endophytic fungi (248–256).

These studies indicate that co-culture generates a complex and promising environment to obtain new secondary metabolites as a response to the interaction between endophytic fungi. The above also indicates that the production of new natural products depends on stimuli.

5. Conclusions

Endophytic fungi are the ubiquitous source of novel chemical compounds having the potential to display antifungal activities. Interestingly, the active metabolites from endophytic fungi possess excellent antifungal activity not only against human fungal pathogens but also on plant fungal pathogens. In addition, the volatile organic compounds (VOCs) from genus *Muscodor* displayed significant antifungal as well as antibacterial properties and, therefore, they are used to prevent fungal deterioration of crops, fruits and vegetables. However, their application to control human fungal infection has not been explored. Fungal VOCs can be investigated for the development of sprays for inhalation to treat fungal diseases such as Aspergillosis in lungs, curing superficial skin infections and sanitization. Endophytic fungi are being studied to produce natural compounds which are originally produced from their host plants and, thus, emerging as an alternative and sustainable source of valuable natural products. It is important to investigate the interactions between endophytic fungi with the host plant and other endophytes which are very sensitive to the culture conditions and hence, provide an opportunity to tune the in vitro culture conditions to produce the desired range of secondary metabolites. It is possible to produce a compound of interest by varying the culture conditions such as media composition, aeration rate and temperature. In addition, cultivation of endophytic fungi in presence of bacteria or other fungi (co-cultivation) yield novel compounds which otherwise do not appear when fungi or bacteria are cultivated alone. Therefore, considerable research on endophytic fungi is required for the development of suitable co-culture system for the sustained production of the desired secondary metabolite.

Author Contributions: S.K.D., V.P., S.S. and M.K.G. reviewed the contents critically. V.P. and M.K.G. drew chemical structures and assisted in the preparation of Table 1. S.S. wrote the Antifungal potential of Volatile organic compounds of review. The manuscript has been read and approved by all named authors.

Funding: This research received no external funding.

Acknowledgments: The authors are thankful to Alok Adholeya, Senior Director, Sustainable Agriculture Division, The Energy and Resources Institute (India) for continuous support.

Conflicts of Interest: The authors declare no conflict of interest.

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Review

Endophytic Fungi in Species of *Artemisia*

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Received: 8 March 2018; Accepted: 28 April 2018; Published: 1 May 2018

Abstract: The genus *Artemisia*, a collection of ~400 hardy herbaceous plant and shrub species, is an important resource contributing to chemistry, medicine, agriculture, industry, and ecology. Its communities of endophytic fungi have only recently begun to be explored. Summarized from studies conducted on the fungal endophytes in *Artemisia* species, both fungal phylogenetic diversity and the associated bioactivity was examined. Isolations from 14 species of *Artemisia* have led to 51 genera of fungal endophytes, 28 families, and 18 orders. Endophytes belonged mainly to *Ascomycota*, except for two taxa of *Cantharellales* and *Sporidiobolales*, one taxon of *Mucoromycota*, and one species of *Oomycota*. The mostly common families were *Pleosporaceae*, *Trichocomaceae*, *Leptosphaeriaceae*, and *Botryosphaeriaceae* (relative abundance = 14.89, 8.51, 7.14 and 6.38, respectively). In the search for bioactive metabolites, 27 novel compounds were characterized and 22 metabolites were isolated between 2006 and 2017. The first study on endophytic fungi isolated from species of *Artemisia* was published but 18 years ago. This summary of recently acquired data illustrates the considerable diversity of biological purposes addressed by fungal endophytes of *Artemisia* spp.

Keywords: medicinal plants microbiome; phylogeny; bioactivity; rare fungal species

1. State of the Art

Artemisia is a genus of plants highly valued as a source of metabolites useful in, for example, medicine and biopesticides. Phytochemical analyses showed the main compounds in *A. vulgaris* to be flavones, flavone glycosides, flavanones, flavonols, flavonol glycosides, and volatile compounds such as α -pinene, camphor, camphene, germacrene D, 1,8-cineole, β -caryophyllene, α -thujone, 1,8-cineole, sabinene, β -thujone, β -caryophyllene oxide, neryl 2-methylbutanoate, β -eudesmol, and bornyl 3-methylbutanoate [1–4]. A review on the chemistry of 15 species of *Artemisia* resulted in 839 compounds with mainly terpenoids, flavonoids, coumarins, caffeoylquinic acids, sterols, and acetylenes [5]. Biological activity was detected in a series of compounds such as eupatilin (anticancer) [5]; artemisolide (anti-inflammatory) [6]; α -thujone (toxicity against adult mites) [7]; chamazulene, 1,8-cineole, and β -caryophyllene (toxicity against the cigarette beetle) [8]; luteolin (anti-inflammatory, anticancer, antimicrobial, antioxidant) [9]; and eriodictyol (anti-inflammatory properties, beneficial effect in treatment of diabetic retinopathy, emmenagogue) [10–12]. Also, *Artemisia annua* remains the main commercial source for artemisinin [13]. Nevertheless, essential oils; organic fractions of hydrolate byproducts; and extracts of various species of *Artemisia*, wild and domesticated (*A. absinthium*, [®]Cvar. Cancial, Spain), proved to have potential in further commercial use with effects such as anti-toxoplasmosis [14], anti-mosquitoes [15], nematicidal [16,17], larvicidal against *Pieris brassicae* [18] and reduction of the longevity and fecundity of *Tetranychus urticae* [19]. Extensive reviews describe (i) the conservation status, phytochemistry and biological activities of the *Artemisia* genus in the Iberian Peninsula and two Macaronesian archipelagos [20]; (ii) the chemical composition and biological activity of essential oils in various

species of *Artemisia* [6] and (iii) the phytochemistry and pharmacological and biotechnological potential of *A. vulgaris* [4].

A survey of the literature shows that this genus has engaged many researchers, with 12,300 publications in the Scopus library, almost double that of the genus *Taxus* (Scopus library—6593 entries). Although *Taxus* is associated with US\$1B per annum worth of sales of the cytostatic drug taxol, the fact that *Artemisia* is a medicinal plant genus with 474 accepted species [21] and multiple uses may explain its relative dominance of the literature. The present survey also reveals the dominance of the plant-focused research rather than that devoted to the plant–endophyte associations. As may be observed in Figure 1, whether we consider one or all databases, the numbers show that up to 0.9% of the number of the publications containing “*Artemisia*” as a keyword include “endophyte” as a keyword (Scopus—88 entries, 0.7%; PubMed—34 entries, 0.9%; and CAB Direct—two entries, 0.45%). For *Taxus* and endophyte, the comparative number of publications was up to 4.28% (Scopus—160 entries, 2.42%; PubMed—88 entries, 4.28%; and CAB Direct—three entries, 3.37%).

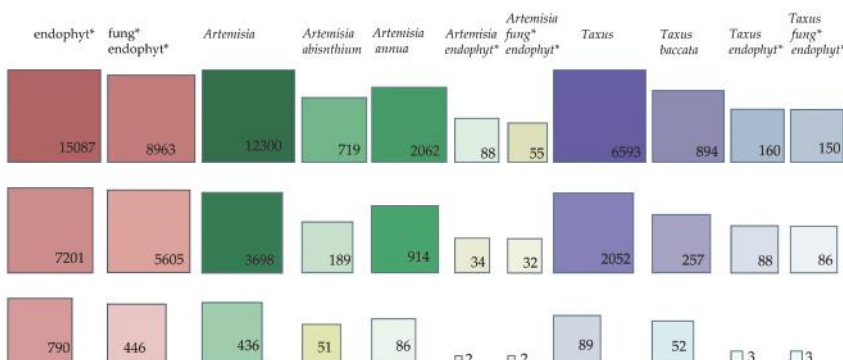


Figure 1. Survey of the literature in databases (Scopus—first upper line of squares; PubMed—middle line with squares; and CAB Direct—bottom line with squares) with indexed keywords related to host plants and endophytes. * = Keywords were used as shown in the figure except for the PubMed database, where the use of wildcards is irrelevant in a standard search as it uses only the root of the term for the generating hits. The search was performed for the title, abstract and keywords or descriptors, where possible (i.e., in Scopus and CAB Direct). Values in squares are the number of publications. Squares are proportional to the number of publications. Interpretation of squares scaling was calculated per group of data as follows: cluster 1—endophytes and fungal endophytes, cluster 2—*Artemisia* and related keywords, cluster 3—*Taxus* and related keywords; the highest number of publications per each cluster was considered 100%. Transformation to \log_{10} was performed to reduce the skewness and facilitate interpretation of the squares.

Comparing the keywords “*Artemisia* endophytes” and “*Artemisia* fungal endophytes” in Scopus, the number of publications containing “*Artemisia* fungal endophytes” represents 62.5% of the number of publications containing “*Artemisia* endophytes”, except in CAB Direct and PubMed where no relevant difference was observed. We might extrapolate that the rest of 37.5% of the publications regard bacterial endophytic communities in *Artemisia*. Similar numbers are shown for the comparison within the *Taxus* genus.

Artemisia annua was highly rated with three times more publications than for *Artemisia absinthium* (Scopus library—2062 hits versus 719, respectively). These values reflect the great interest in artemisinin, produced by *Artemisia annua*, which is an important drug used to treat malaria and cancer. Various fungi have been investigated as elicitors or direct producers of artemisinin. Up-to-date information in Table 2 summarises the findings from research into endophytes of *Artemisia* endophytic communities.

2. *Artemisia* Fungal Endophyte Identification and Diversity

Overall, the identification of the fungal endophytes in *Artemisia* spp. is made based on morphological characterization and molecular analysis using (i) 28S ribosomal DNA spanning domains D1 and D2 and (ii) nuclear ribosomal DNA sequences, including both the internal transcribed spacers (ITS1 and ITS2) and the 5.8S gene region. Three studies investigated the phylogenetic analysis of the *Artemisia* spp. fungal endophytes [22–24]. Although the number of studies is small, interesting facts are brought to light in terms of diversity and plant colonization. For *Artemisia annua*, the endophytic fungal infection frequency was slightly higher in the roots of cultivated plants (20.9%) than in the roots of wild plants (16.7%) [25]. Moreover, the authors showed that the latter roots harboured more rich fungal taxa, which supports the hypothesis that wild plant species are predisposed to be hosts for rich and novel mycoflora [26,27]. The differences can be extended to fungal pathogens, where 55% of the fungal communities in wild fruits were not present in their homologous cultivated species [28].

Strains of *Aplopsorella prunicola*, *Chaetomium* sp., *Macrophomina phaseolina*, *Nectria mauritiicola*, *Neofusicoccum australe*, *Pestalotiopsis* sp., and *Stachybotrys longispora* were isolated only from one nutrient medium and only from *Artemisia thuscula* collected in Tenerife island, in a study comparing La Palma and Tenerife in the Canary Islands [24]. Diversity indices showed that plant individuals collected in La Palma had higher diversity values in terms of species richness, diversity of taxa, abundance of rare species, and species evenness. The study indicates the relevance of the exploration of different plant tissues and usage of wider number of growth media to obtain diversity. Comparing arbuscular mycorrhizal fungi (AMF) with dark septate endophytes (DSE) and other types of endophytic fungi (EF), a low colonization frequency of AMF was detected in roots, while EF were more frequently found in *Artemisia annua* [25].

Endophytic fungi were isolated from *Artemisia argyi*, and *Pleosporales* was found to be the most represented group, with three species of *Alternaria* present [29]. It is worth mentioning that Qian et al. [29] reported the presence of *Rhodotorula* sp. and *Fusarium* sp. in *Artemisia argyi*. Myrchiang et al. [30] investigated the endophytic fungi associated with *Artemisia nilagirica*, and isolated among the dominant clade of *Ascomycota*, one strain of *Pythium intermedium* (Oomycota) and one strain of *Rhizopus oryzae* (Zygomycota). Comparing the colonization of three plant parts (i.e., root, stem, and leaf), the authors found that the highest diversity was in the roots (i.e., 14 species), followed by stems (i.e., 10 species), and least in the leaves (i.e., six species). Similarly, in *Artemisia thuscula* were isolated 29 distinct morphotypes: 20 from roots, seven from stems, and two from leaves [31]. In addition, Myrchiang et al. [30] observed that from all fungal endophytic species, only *Phoma eupryrena* was found to be most dominant in all plant samples; the other species having a certain preference for one or a maximum of two plant parts. Huang et al. [22] classified 108 fungal isolates obtained from three medicinal plant species, *Artemisia capillaris*, *Artemisia indica*, and *Artemisia lactiflora*, using morphological identification. In total, 42 isolates were obtained from the host *Artemisia capillaris* (five isolates from leaves, 15 isolates from stems, and 22 isolates from inflorescences). In total, 39 strains were isolated from the host *Artemisia indica* (18 isolates from leaves and 21 from stems). Twenty-seven fungal strains were isolated from the host *Artemisia lactiflora* (16 isolates from leaves and 11 from stems). Among them, *Alternaria*, *Colletotrichum*, and *Phomopsis* were common, especially *Alternaria* spp. (25% relative isolation frequency) and *Colletotrichum* spp. (20.4% relative isolation frequency). One strain was isolated from the stems of *Artemisia capillaris* and was similar to *Drechslera* sp. Among the three plant hosts, the highest endophytic colonization rate occurred in *Artemisia capillaris*, which had the highest fungal diversity. Five fungal isolates belonging to *Aureobasidium pullulans*, *Ephelis*, *Pestalotiopsis*, and *Pleosporaceae* were only obtained from *Artemisia capillaris*. *Xylaria* species appeared to be the most common endophytic fungi in *Artemisia indica*.

Seven *Artemisia* species were sampled in two locations (Qichun and Wuhan), and 21 fungal endophytic species were isolated from stems and identified as *Diaporthe*, *Colletotrichum*, *Nigrospora*, *Botryosphaeria*, *Aspergillus*, *Penicillium*, *Neofusicoccum*, *Cercospora*, *Rhizoctonia*, *Alternaria*, and *Curvularia* [23]. The highest incidences of colonization frequency per plant were attributed

to *Nigrospora sphaerica* in *Artemisia* sp., *Nigrospora oryzae* in *Artemisia argyi*, *Alternaria alternata* in *Artemisia subulata* and *Artemisia tangutica*, and *Botryosphaeria dothidea* in *Artemisia lavandulifolia*. This was the first report of *Nigrospora*, *Neofusicoccum* and *Curvularia* species in *Artemisia* spp. As for plant species specificity, only *Nigrospora sphaerica*, *Nigrospora oryzae* and *Alternaria alternata* were present in various plants; except for *Cochliobolus geniculatus*, isolated from *Artemisia brachyloba* and *Artemisia argyi*. Thus, few endophytic fungi were found to be entirely restricted to particular plant species, but significant differences were found in the frequency of colonization of individual morphotaxa. The study highlighted the *Alternaria* genus as the dominant one, having the reputation of being one of the cosmopolitan endophytes reported. *Nigrospora* displayed the second highest rate of occurrence in this analysis, followed by *Botryosphaeria* whose incidence has been described more in woody plants.

A heat map table (Table 1) describes the relative abundance of genera, families and orders of the endophytic mycopopulation of *Artemisia* species. The most isolated taxa belong to *Pleosporales*, *Hypocreales*, *Xylariales*, *Botryosphaerales*, and *Eurotiales* (relative abundance % = 25, 17.31, 9.62, 7.69, and 7.69, respectively). Yet these numbers do not reflect the diversity of the families included, as *Pleosporales* and *Hypocreales* have been found each with taxa belonging to four families while *Hypocreales* and *Xylariales* were found with four and two families, respectively. *Phomopsis*, *Alternaria* and *Penicillium* were the most isolated genera (relative abundance % = 36.36, 36.36 and 33.33, respectively). While many of the genera (Table 1) are frequently found as endophytic strains in medicinal plants [32–35], temperate grasses [36], tropical and subtropical rainforests [37] and also as fungi associated with *Vitis vinifera* [38], some are quite rare: *Aplopsorella*, *Ramichloridium*, *Ephelis*, *Stachybotrys*, *Drechslera*, *Biscogniauxia*, *Edenia* and *Thielavia*.

Table 1. Endophytic fungi order, family, genus and host plants with values of relative abundance (RA) for genus, family, and order. Values are represented using a color scale with green representing the smallest value and red the highest value.

Order RA %	Family RA %	Genus	Census RA % Per Host Plant/Plants	Host Plants
Amphisphaerales 1.92	Pestalotiopeltidaceae 2.13	<i>Pestalotiopsis</i>	0.9 1.19 11.76	<i>A. capillaris*</i> + <i>A. indica*</i> + <i>A. lacithophora*</i> (HK); <i>A. thyscula</i> (TF)
	Aphyllonellaceae 2.13	<i>Aphyllonella</i>	1.22	<i>A. thyscula</i> (TF)
Botryosphaerales 7.69	Ranquephoraceae 2.13	<i>Ranquephaea</i>	41.67	<i>A. lacaudifolia</i> (C-)
	Botryosphaeriaceae 6.38	<i>Macropythoma</i>	2.44	<i>A. thyscula</i> (TF)
	Nectriaceae 2.13	<i>Nectriococcum</i>	2.78 30.3 10.98	<i>A. lacaudifolia</i> ; <i>A. thyscula</i> (LP); <i>A. thyscula</i> (TF)
Cercophiales 1.92	Cercobasidiaceae 2.13	<i>Rhizoctonia</i>	16.67	<i>A. brachyloba</i> (C-H)
	Cladospoidiaceae 2.13	<i>Cladospodium</i>	2.44 2.77 5.68 5.69	<i>A. thyscula</i> (TF); <i>A. annua</i> * (CH); <i>A. argy*</i> ; <i>A. nilagirica*</i> (IN)
Copromiales 3.85	Dissoscydaceae 2.13	<i>Ramichordium</i>	2.77	<i>A. annua</i> *
	Diaporthaceae 4.26	<i>Diaporthie</i>	5.86 16.67 8.33 9.09 5.88	<i>A. lacaudifolia</i> ; <i>A. brachyloba</i> ; <i>A. scoraria</i> (CH); <i>A. thyscula</i> (LP); <i>A. argy*</i>
	Phomopsidae	<i>Phomopsis</i>	6.3 5.88 36.36	<i>A. capillaris</i> + <i>A. indica</i> + <i>A. lacithophora*</i> + <i>A. argy*</i>
Dothideales 1.92	Dothioraceae 2.13	<i>Aureobasidium</i>	0.9 3.66	<i>A. lacithophora</i> (CH)
		<i>Aspergillus</i>	8.33 4.55 8.69	<i>A. capillaris</i> + <i>A. indica</i> + <i>A. lacithophora*</i> ; <i>A. thyscula</i> (TF)
		<i>Eupenicillium</i>	5.85	<i>A. lacaudifolia</i> ; <i>A. thyscula</i> (LP); <i>A. nilagirica*</i>
		<i>Paecilomyces</i>	2.77	<i>A. annua</i> *
Eurotiales 7.69	Trichocomaceae 8.51	<i>Penicillium</i>	2.78 1.52 13.8 3.88 33.33 13.4	<i>A. lacaudifolia</i> ; <i>A. thyscula</i> (LP); <i>A. annua</i> *; <i>A. argy*</i> ;
		<i>Colletotrichum</i>	19.81 11.1 25 5.55 5.88 10.11 9.09	<i>A. capillaris</i> + <i>A. indica</i> + <i>A. lacithophora*</i> ; <i>A. lacaudifolia</i> ; <i>A. brachyloba</i> ; <i>A. annua</i> *; <i>A. argy*</i> ; <i>A. annua</i> ; <i>A. lacithophora</i> (CH)
Gloomerellales 1.92	Gloomerellaceae 2.13	<i>Ephedis</i>	1.8	<i>A. capillaris</i> + <i>A. indica</i> + <i>A. lacithophora*</i>
		<i>Hypocreaceae 4.26</i>	5.88 9.09	<i>A. argy*</i> ; <i>A. lacithophora*</i>
		<i>Trichocomella</i>	4.34	<i>A. nilagirica*</i>
		<i>Fusarium</i>	5.55 5.88 4.34	<i>A. annua</i> *; <i>A. argy*</i> ; <i>A. nilagirica*</i>
Hypoconiales 17.31	Nectriaceae 4.26	<i>Nectria</i>	2.44	<i>A. thyscula</i> (TF)
	Stachybotryaceae 4.26	<i>Stachybotrys</i>	1.22	<i>A. thyscula</i> (TF)
		<i>Mycotilicum</i>	N.A.	<i>A. nilagirica*</i>
		<i>Armeniaca</i>	13	<i>A. annua</i>
		<i>Cephalosporium</i>	N.A.	<i>A. annua</i>
Mucorales 3.85	Mucoraceae 2.26	<i>Rhizopus</i>	4.34	<i>A. nilagirica*</i>
Oomycetales 1.92	Arthrobotryidae 2.13	<i>Micor</i>	N.A.	<i>A. annua</i>
		<i>Arthrobotryma</i>	4.34	<i>A. nilagirica*</i>
	Sporangiaceae 2.13	<i>Pressia</i>	10.61 10.98	<i>A. thyscula</i> (LP); <i>A. thyscula</i> (TF)
	Didymosphaeraceae 2.13	<i>Tremellia</i>	1.52	<i>A. thyscula</i> (LP)
	Leptosphaeriaceae 4.26	<i>Coniothyrium</i>	1.52	<i>A. thyscula</i> (LP)
		<i>Lepidosphaeria</i>	N.A.	<i>A. annua</i>
Phasmatopores 25.00	Pleosporaceae 14.85	<i>Alternaria</i>	24.30 13.89 75 25 66.67 33.33 23.5 36.36 28.79 41.46	<i>A. capillaris</i> + <i>A. indica</i> + <i>A. lacithophora*</i> ; <i>A. lacaudifolia</i> ; <i>A. tangutica</i> ; <i>A. brachyloba</i> ; <i>A. subtilis</i> ; <i>A. argy</i> ; <i>A. scoparia</i> ; <i>A. argy*</i> ; <i>A. lacithophora*</i> ; <i>A. thyscula</i> (LP); <i>A. thyscula</i> (TF)

Table 1. *Cont.*

RA % for genus—calculated as the number of isolates belonging to a genus divided by the total number of individuals. Each value of RA was calculated for a host (except *A. capillaris* + *A. indica* + *A. lactiflora*, where all three species were considered as one pool as the study did not allow other analysis). For the column RA %—genus, each cell represents a host species in the right column, separated by “.”. Each cell corresponds to a host species in the right column. RA % for family and order—the number of families that one taxa was found in this survey divided by the total number of taxa (No. of families = 47, No. of order = 52); although no quantitative data was found for genera such as *Hippophae*, *Lepidophyllum*, *Mucor* and *Myrothecium*; these were used only to calculate the RA of the order/family. *—calculation of relative abundance made by the present authors using the data provided in the reviewed studies. For host plant species the geographical location was provided only at the first time of appearance in the text: CH = China, HK = Hong Kong, island, IN = India, IP = La Palma island, TF = Tenerife island; except LP and TF to distinguish two locations for the same host species in the same study. Following studies were used for the table: *A. capillaris* + *A. indica* + *A. lactiflora**—[133]; *A. millefolium**—[301]; *A. monostachya**—[57, 40, 43]; *A. nigrum**—[39]; *A. officinale**—[29]; *A. officinale**—[124]; *A. thlaspi**—[29]; *A. thlaspi**—[124]; *A. thlaspi**—[29]. A. *innocua**—[133]; *A. subulata*, *A. tenuis*, *A. villosa**—[301]

Rare and widely abundant species inhabiting *Artemisia* host plants are different from each other in distribution–abundance relationship (Figure 2). Some of the considered shortcomings of the study are: (i) data is based on purely culture-dependent strains; (ii) the sampling method differs among studies (for instance, in terms of considering two different strains of the same species, there is no standardized method or at least an extended discussion in the literature); and (iii) the lack of information hidden behind the studies which are interested in showing the bioactivity of certain species (or else) without mention on the abundance of strains.

The plotted data shows that the theoretical assumption of a log-normal distribution of the abundance in nature, that is, endophytic fungal communities, where few common taxa have a high abundance and also few taxa are scarcely found, does not fit to the collected data; to name relevant examples: (i) abundant genera (between three and 11 mentions) *Alternaria*, *Nigrospora*, *Penicillium*, *Phomopsis*, and *Neofusicoccum* (highest RA % = 75, 75, 36.4, 33.3, 41.7 and 30, respectively); (ii) least abundant (between two and seven mentions) *Pestalotiopsis*, *Colletotrichum*, *Diaporthe*, *Acremonium*, *Preussia*, and *Chaetomium* (highest RA % = 11.8, 19.8, 16.7, 13, 11 and 11.8, respectively); and (iii) singleton (with one mention) taxa such as *Edenia*, *Drechslera*, *Stachybotrys*, and *Ephelis* (RA % = 2.7, 0.9, 1.2 and 1.8, respectively). Studies show that the extension of data sampling is directly proportional to the unveiling of the missed taxa [44]. So, the deviations from the log-normal distribution can be used to estimate the inventory size and assume that more rare species would be identified or fall into common categories had there been more sampling and computation.

In this data collection, 71% of the species were less abundant (between one and eight mentions of strains) with 37.77% of the species being singleton. The cosmopolitan theory has been previously argued and demonstrations against its postulates (i.e., everything is everywhere) have shown patterns of biogeography for the rare marine microbial biosphere [45]. Furthermore, 59% of the endophytic fungal species found in tropical forest of Panama were singletons [46] while incidences of endophytic colonization depend on environmental and geographical factors [33]. Isolated ecosystems are known for their specific biodiversity. In this dataset, 28% of the taxa were isolated from *A. thyscula* in the Canary Islands while the rest of the genera were found among all regions (China, Hong Kong, India, La Palma, and Tenerife).

Studies on endophytic communities isolated from a plant genus host are encouraged, as besides the implications of the ecosystem, the relations with the host are considered to be specific (chemical, biochemical and physiological levels as well as the biological life cycle). Furthermore, studies have shown a presumable specificity or preference of various endophytic taxa for their hosts [47–49].

This difference in the distribution–abundance relationship between the rare and cosmopolitan taxa raises a question: what determines the exclusive occurrence of singletons; is it sampling size, host, or site?

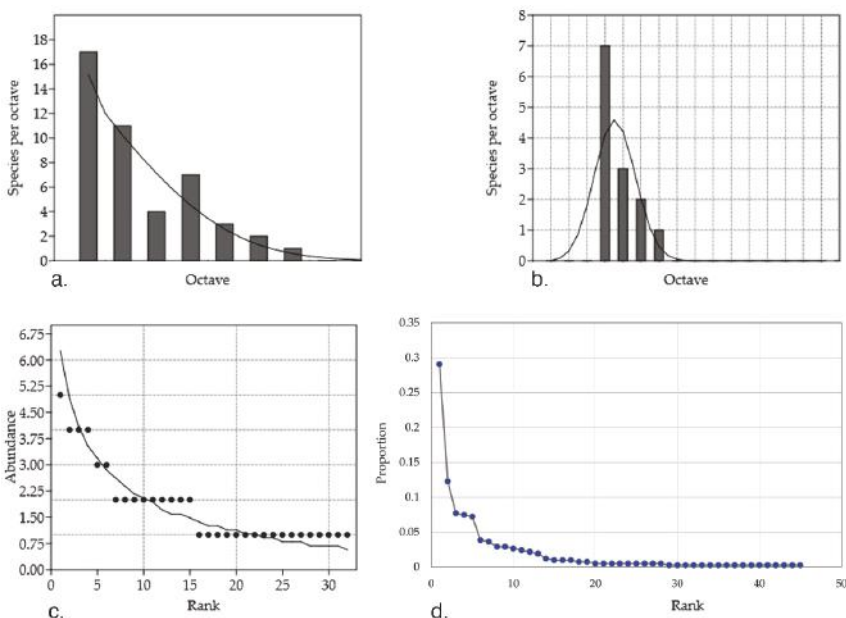


Figure 2. (a) The number of all endophytic fungi (EF) genera plotted in a log-normal distribution (fitted mean = 0.05, variance = 0.68, $\chi^2 = 2.81, p = 0.42$) for different abundance octaves. The octaves refer to power-of-two abundance classes. The species which theoretically are expected to be present are veiled, as their low abundance prevents them for being represented in the sample. The log-normal distribution was a poor fit to this data due to an overabundance of rare species compared with that expected from the log-normal curve; (b) Abundant EF genera (>eight individuals) predicted by a log-normal distribution (mean = 1.28, variance = 0.12, $\chi^2 = 3.9, p = 0.04$); a poor fit, as the first interval is dominated by a genus, namely *Alternaria*; (c) Rare EF species (< eight individuals) predicted by a log series distribution ($\alpha = 29.32, x = 0.66, \chi^2 = 0.72, p = 0.99$), where many rare genera (17 genera with one strain) are represented by a jagged line because predicted values are rounded to the nearest number; (d) Rank abundance curve of the endophytic fungal genera. The slope is extremely steep at the beginning at the curve, meaning that evenness of the community is low, where few of the genera are highly abundant. For (a–c), only absolute counts, as opposed to relative proportions, were considered, while for (d), relative proportions were considered. Abundance model were calculated with PAST 3.18 software (Øyvind Hammer, Natural History Museum, University of Oslo, Norway) [50].

3. Biological Activity of *Artemisia* Endophytes

Extremely diverse biological activity demonstrated by endophytic fungi associated with *Artemisia* spp. has been a driving force in the search for active endophytic strains in *Artemisia annua* [41,51–54]. Two new compounds have been discovered, namely leptosphaerone [43] and leptosphaeric acid [55] which were obtained from *Leptosphaeria* sp. (Table 2). Fungal endophytes in this plant species were mainly evaluated for their elicitor role in artemisinin production, for their direct production of the molecule or as their role in planta enhancement of artemisinin. Likewise, Lu et al. [40] isolated *Colletotrichum* sp., an endophytic fungus producing the plant hormone indole-3-acetic acid (IAA). Plant growth-promoting traits for example increase in plant biomass and chlorophyll content in leaves was demonstrated using a dual symbiotic culture of *Piriformospora indica* and *Azotobacter chroococcum* [56]. In addition, a positive influence of *P. indica* filtrate on overall growth and development of micro-propagated plants was observed [57]. Chlorophyll a, chlorophyll b, and carotenoid content were found at the highest rates in *Piriformospora indica*—associated plants

compared to the control and autoclaved *Piriformospora indica* treated plants [58]. Substantial increase in height and leaf area of *Artemisia annua* was observed in plants treated with fungal elicitor extracts obtained from *Cochliobolus lunatus*, *Colletotrichum gloeosporioides*, *Acremonium persicinum* and *Curvularia pallescens* [59].

Monitoring the developmental effects of the *P. indica* endophytic fungus on its *Artemisia annua* host plant, researchers studied the effect on artemisinin production [56,57]. This was proved not only to be an antimalarial drug for human health, but also known to be very effective against a wide spectrum of microorganisms including protozoa, bacteria, fungi and viruses as well as serving as a selective insecticide and phytoalexin [60] and also being used in cancer treatment [61]. In addition, oligo- or polysaccharides derived from endophytic *Colletotrichum* sp. were used successfully as elicitors to stimulate artemisinin production in *Artemisia annua* hairy root cultures [53,54,62]. *Artemisia annua* plants resulted in a 3.47-times increase ($p < 0.01$) in artemisinin production after being treated with *Curvularia pallescens* fungal elicitor extract [59]. Also, mycelium extracts of *Penicillium oxalicum* promoted artemisinin biosynthesis in *Artemisia annua* seedlings [25,63]. Furthermore, the induction of artemisinin biosynthesis by *Penicillium oxalicum* B4 in *A. annua* was shown to be strongly dependent on the induced reactive oxygen species (ROS) production [63].

Biological activity against bacteria and fungi was also sought, following the “medicinal plant harbouring active fungal endophytes” model; this time spreading the screening across various species of *Artemisia*. Lu et al. [40] isolated and characterised three new metabolites, 6-isoprenylindole-3-carboxylic acid, 3b,5a-dihydroxy-6b-acetoxy-ergosta-7,22-diene, and 3b,5a-dihydroxy-6b-phenylacetoxy-ergosta-7,22-diene, from the culture of the endophyte *Colletotrichum* sp. (host plant: *Artemisia annua*). Various inhibitory activities were observed against *Candida albicans*, *Aspergillus niger*, and phytopathogenic fungi *Gaeumannomyces graminis* pvar. *tritici*, *Rhizoctonia cerealis*, and *Helminthosporium sativum*, as well as against the bacteria *Bacillus subtilis*, *Staphylococcus aureus*, *Micrococcus luteus*, and *Pseudomonas* sp. A strain of *Colletotrichum gloeosporioides* from *Artemisia mongolica* produces a novel secondary metabolite named colletotric acid inhibiting growth of *Bacillus subtilis*, *Staphylococcus aureus*, and *Micrococcus luteus* [64]. Three strains of *Aspergillus* spp. (i.e., SPS-02, SPS-04, and SPS-01) were isolated from *Artemisia annua* and were found to possess strong antimicrobial activities against the human pathogens *Escherichia coli*, *Staphylococcus aureus* and *Trichophyton rubrum*, and cytotoxic activities [41]. The authors also found an inhibitory effect on *Rhizoctonia cerealis* with a strain of *Mucor* sp. SPS-11, and the strongest antimicrobial activities observed against *Magnaporthe grisea* were exhibited by two strains of *Aspergillus fumigatus* (SPS-02) and *Cephalosporium* sp. (SPS-08). The same strain of *A. fumigatus* produced four ardeemin derivatives with various activities of reversing the multidrug-resistant phenotype in three cancer cell lines [65].

Seven endophytic fungal strains isolated from leaves of *Artemisia annua* were solvent-extracted and tested against the bacteria *Staphylococcus aureus*, *Streptococcus mutans*, *Salmonella enterica* serotype Typhi, and *Bacillus subtilis*, and the fungi *Malassezia furfur* and *Candida albicans* [66]. Three strains had the broadest range of antimicrobial activity (i.e., inhibition of all tested strains), one strain inhibited only *E. coli* and *S. aureus*, and three endophytes were active against three bacterial strains and *M. furfur*. None were active against *C. albicans*. Extracts obtained from cultures of endophytic fungi (i.e., *Rhodotorula* sp. and *Fusarium* sp.) isolated from *Artemisia argyi* proved cytotoxic against liver cancer cell lines (Hep G2) [67]. Similarly, *Pestalotiopsis hainanensis* was active against all tested cell lines (MCF-7 breast, COLO205 colon, and HL-60 leukemia), while *Phomopsis mali* and *Rhodotorula glutinis* inhibited the HL-60 leukemia cell line, *Alternaria* sp. was active only against the MCF-7 breast cell line, and *Colletotrichum gloeosporioides* inhibited growth of the COLO205 colon cell line [29].

(i) Five new fungal polyketides and (ii) two known analogues were isolated from EtOAc-extracted biomass of *Trichoderma koningiopsis* (strain QA-3): (i) *ent*-koninginin A, 1,6-di-*epi*-koninginin A, 15-hydroxykoninginin A, 10-deacetylkoningsin D and koninginin T; and (ii) koninginin L and trichoketide A [68]. Both *ent*-koninginin A and trichoketide A inhibited *E. coli*, and the second

compound also showed antifungal activity against *Bipolaris sorokiniana*, *Ceratobasidium cornigerum*, *Colletotrichum gloeosporioides*, *Fusarium graminearum*, *Fusarium oxysporum*, *Penicillium digitatum*, *Physalospora piricola* and *Valsa mali*. *Myrothecium rorideum* (strain IFB-E012) isolated from stems of *Artemisia annua* produced lumichrome, a compound with moderate cytotoxicity to the human cell line KB [51]. Furthermore, a new cytotoxic trichothecene macrolide dihydromyrothecine compound was isolated from the same strain [69] together with three new 10,13-cyclotrichothecane-based macrolides [70]. *Hypoxyylon truncatum*, an endophytic fungus isolated from *Artemisia annua*, yielded two novel cytotoxic benzofluoranthene-based metabolites named daldinone C and daldinone D [42].

Previous studies exhibited antibiotic and antitumor activities provoked by secondary metabolites such as azaphilones, which were demonstrated to be of chemotaxonomic significance in the classification of the *Hypoxyylon* species [71]. Antioxidant phenolic metabolites were also reported to be produced by fungal endophytes inhabiting *Artemisia capillaris*, *Artemisia indica* and *Artemisia lactiflora* [72].

Likewise, 12 endophytic fungi isolated from *Artemisia absinthium* were tested for their ability to infect grape berries and for their potential to reduce the infection with *Botrytis cinerea*. Three of the fungal strains reduced both the percentage of infected berries and lesion diameter [73]. Antagonism assays are the preliminary step in these kinds of studies. For instance, Myrchiang et al. [30] proposes as good biocontrol agents of *Phytophthora infestans* the use of *Trichoderma viride*, *Aspergillus fumigatus*, and *Penicillium atrovenetum*, isolated from *Artemisia nilagirica*. Liu et al. [52] found 14 fungal endophytes in *Artemisia annua* which produce antagonistic substances against four phytopathogenic fungi: *Fusarium graminearum*, *Rhizoctonia cerealis*, *Helminthosporium sativum*, and *Geotrichum candidum*. Furthermore, inhibitory activities of culture broths were evaluated. Similarly, a series of endophytic fungi from seven species of *Artemisia* (i.e., *Artemisia tangutica*, *Artemisia brachyloba*, *Artemisia subulata*, *Artemisia argyi*, *Artemisia scoparia*, *Artemisia lavandulifolia*, and *Artemisia* sp.) were evaluated [23]. It is worth mentioning that both studies found that antagonism assays (i.e., dual culture) do not converge with toxicity assays (i.e., nutrient media amended).

Research on strains isolated from *A. annua* to pursue their degradation of triclosan, the emerging contaminant in the aqueous and soil environment, resulted in the identification of an active strain, B4, of *Penicillium oxalicum* [74]. Triclosan was removed from the culture media by fungal degradation with a rate of 97.26% after one hour of incubation in fungal culture medium (5 mg L^{-1}). Effective results were displayed in nonsterile synthetic wastewater and completion of degradation of triclosan was obtained in two hours. It represents a pioneering study that demonstrated the degradation of the notorious antimicrobial pollutant triclosan.

During the last decade, limited studies have been conducted on bioprospection of the microbiome of *Artemisia* sp. Nevertheless, its endophytic fungal communities have been a source of 27 novel compounds and 22 already characterised molecules of which 35 were shown to have at least one interesting trait to be further pursued in biotechnology.

Table 2. Literature review on bioactive endophytic fungi (EF) isolated from species of *Artemisia*: host species, plant part of isolation, fungal taxa, endophyte isolation method, bioactive compounds/extracts, class of compounds, bioactivity, and references.

Artemisia Species	Plant Part	EF Taxa	EF Identification Method	Compound/Extract	Class of Compounds	Bioactivity	Reference
<i>A. absinthium</i>	root	<i>Penicillium</i> spp. (strains B6C18 and B14C36)	N.A.	spore solution (10^6 μL^{-1})	N.A.	Antifungal inhibition of diameter of lesions on vine berries; 75% and 91%	[73]
<i>Aspergillus</i> spp.							
<i>Mucor</i> spp.		morphology		ethyl acetate crude extract	N.A.	Antibacterial, antifungal	[41]
<i>Cephalosporium</i> spp.							
<i>Fusarium</i> spp.							
<i>Aspergillus terreus</i> (strain IFB-E030)			N.A.	10-Phenyl-[12]-cytochalasins Z17	Cytochalasins, alkaloids	Cytotoxic [$\text{IC}_{50} = 26.2 \mu\text{M}$]	[75]
<i>Aspergillus</i> spp.							
<i>Colletotrichum</i> sp.	morphology			3 β -hydroxy-ergosta-5-ene	Ergosterol derivatives	Antifungal % ($200 \mu\text{g mL}^{-1}$): 77–85%	[40]
<i>Hypoxyylon truncatum</i>	stem			3-oxo-ergosta-4,6(1),22-tetraene	Triunsaturated steroids	Antimicrobial MIC ($\mu\text{g mL}^{-1}$): 50–75%	[69,76]
<i>Myrothecium roridum</i> (strain IFB-E012)	morphology and 18S rDNA sequence			3 β -hydroxy-5 α ,8 α -epidioxy-ergosta-5,22-diene	Ergosterol peroxides	Antimicrobial MIC ($\mu\text{g mL}^{-1}$): 50–75%	[42]
<i>Aspergillus fumigatus</i>				6- α -soprenylindole-3-carboxylic acid	indoles	Antimicrobial MIC ($\mu\text{g mL}^{-1}$): 55–77%	[40]
<i>Aspergillus</i> spp.				3 β ,5 α -dihydroxy-6 β -acetoxy-ergosta-7,22-diene	sterols	Antifungal % ($200 \mu\text{g mL}^{-1}$): 75–77%	[69,76]
<i>Aspergillus</i> spp.				3 β ,5 α -dihydroxy-6 β -phenylacetyl-oxo- α -ergosta-7,22-diene		Antimicrobial MIC ($\mu\text{g mL}^{-1}$): 50–100%	[40]
<i>Aspergillus</i> spp.				daldinone C	Benzoflouranthenes (polycyclic aromatic hydrocarbons)	Cytotoxic [$\text{IC}_{50} = 49.5 \mu\text{M}$]	[65]
<i>Aspergillus</i> spp.				daldinone D		Cytotoxic [$\text{IC}_{50} = 41 \mu\text{M}$]	[65]
<i>Aspergillus</i> spp.				myrothecine A	Sesquiterpene-based trichothecenes	Cytotoxic [$\text{IC}_{50} = 8.5 \mu\text{g mL}^{-1}$]	[65]
<i>Aspergillus</i> spp.				myrothecine B		Cytotoxic [$\text{IC}_{50} = 0.76 \mu\text{g mL}^{-1}$]	[65]
<i>Aspergillus</i> spp.				myrothecine C		Cytotoxic [$\text{IC}_{50} = 32.21 \mu\text{g mL}^{-1}$]	[65]
<i>Aspergillus</i> spp.				roridin E	Macrocyclic trichothecenes	Cytotoxic [$\text{IC}_{50} = 0.03 \mu\text{g mL}^{-1}$]	[69,76]
<i>Aspergillus</i> spp.				mytoxin B		Cytotoxic [$\text{IC}_{50} = 0.002 \mu\text{g mL}^{-1}$]	[65]
<i>Aspergillus</i> spp.				dihydromyrothecine C	Trychothecene macrolides	Cytotoxic [$\text{IC}_{50} = 44.48 \mu\text{M}$]	[65]
<i>Aspergillus</i> spp.				roritoxin E		Cytotoxic [$\text{IC}_{50} = 0.26 \mu\text{g mL}^{-1}$, $10.54 \mu\text{g mL}^{-1}$]	[65]
<i>Aspergillus</i> spp.				5-N-acetyl-1- β -hydroxydearmeen		Cytotoxic RF = 1.5–6 μM	[65]
<i>Aspergillus</i> spp.				5-N-acetyl-1- β -dihydroxydearmeen		Cytotoxic RF = 2.9–5.1 μM	[65]
<i>Aspergillus</i> spp.				5-N-acetyl-1- α -hydroxydearmeen		Cytotoxic RF = 2.5–8 μM	[65]
<i>Aspergillus</i> spp.				5-N-acetyl-1- α -hydroxydearmeen		Cytotoxic RF = 1.5–7 μM	[65]

Table 2. Cont.

	<i>Coleotrichum gaeppertiae</i>	morphology	oligosaccharides extract elicitor for artemisinin	increment of artemisinin = 64.29% [62]
	<i>Penicillium coeruleum</i>	ITS1, 5.8S, and ITS2	heat-killed mycelium degrading tricosan (TCS)	max. adsorbing capacity of TCS = 17.60 mg g ⁻¹ [74]
leaves	<i>Curvularia pallens</i>		elicitor extract for artemisinin	2–6% elicitor extract (w/v); artemisinin content = 1.21–3.47-times more than control [59]
	<i>Cladosprium sp.</i>	morphology	ethyl acetate extract	Antibacterial [66]
	<i>Paraphloeosphaeria moliniae</i> (strain IB-E011)	N.A.	paranolin	N.A.
stem	<i>Phomopsis sp.</i>		crude EOAc extract = BB1; fraction Hex/EDAc/MeOH = BB4 fractions MeOH = BB8, BB9, BB10	Cytotoxic IC ₅₀ values > 50 µg mL ⁻¹ [77]
			tyrosol (purified from BB10)	Cytotoxic: IC ₅₀ (µg mL ⁻¹) = 17.11, BB4 = 0.7%; % (20 µg/mL) BB8 = 1.97, BB9 = 0.53, BB10 = 53.98 [78]
	<i>Phomopsis sp.</i> (strain H31)			Antifungal % (1 mg mL ⁻¹) = 34.57; Cytotoxic % (20 µg mL ⁻¹) = 31.69 [66]
	<i>Cladosporium sp.</i> (strain H23)			Antifungal % (1 mg mL ⁻¹) = 23.46 [66]
	<i>Chaetomium sp.</i> (strain H40)			Antibacterial [29]
	<i>Penicillium sp.</i> (strain H9)			Antibacterial/Cytotoxic % (20 µg mL ⁻¹) = 53.47; 71.43, 97.03 [29]
	<i>Pestalotiopsis sp.</i> (strain HI4)	ITS1, 5.8S, and ITS2	ethyl acetate + ethyl acetate/water extract	Antibacterial [29]
	<i>Diaporthella sp.</i> (strain H26)			Antibacterial [29]
	<i>Trichoderma sp.</i> (strain HT7)			Cytotoxic % (20 µg mL ⁻¹) = 16.41, 23.56, 78.28 [29]
	<i>Rhodotorula sp.</i> (strain H13)			Cytotoxic % (20 µg mL ⁻¹) = 11.22, 26.95, 71.99 [29]
	<i>Coleotrichum sp.</i> (strain H42)			
A. argyi	<i>Nigrospora sphaerica</i>	morphology and ITS1, 5.8S, and ITS2	ethyl acetate crude extract	N.A.
				Antifungal % (1 mg mL ⁻¹) = 48.37, 11.94, 24.92 [23]
				Antifungal MIC (µg mL ⁻¹) = 8–64 [23]
stem	<i>Trichoderma koningiopsis</i> (strain QA-3)	ITS1, 5.8S, and ITS2	<i>ent</i> -konkinginin A 1,6-di- <i>epi</i> -konkinginin A 15-hydroxykonkinginin A	Antibacterial MIC (µg mL ⁻¹) = 8–64 [68]
				Antifungal polyketides [68]
				Tricyclic polyketides [68]
				Antifungal MIC (µg mL ⁻¹) = 64 [68]
				Antibacterial MIC (µg mL ⁻¹) = 16–64 [68]

Table 2. Cont.

<i>A. brachyloba</i>		<i>Alternaria alternata</i>	morphology and ITS1, 5'S, and ITS2	ethyl acetate crude extract	N.A.	Antifungal MIC ($\mu\text{g mL}^{-1}$): 8–64 Antibacterial MIC ($\mu\text{g mL}^{-1}$): 32–64
				Bicyclic polyketides		Antifungal MIC ($\mu\text{g mL}^{-1}$): 16 Antibacterial MIC ($\mu\text{g mL}^{-1}$): 32–64
<i>A. capillaris</i>	flower	<i>Alternaria</i> sp. (strain Acap F1)		Tricyclic polyketides		Antifungal MIC ($\mu\text{g mL}^{-1}$): 16 Antibacterial MIC ($\mu\text{g mL}^{-1}$): 32–64
		<i>Phomopsis</i> sp. (strain Acap, F3)	ITS1, 5'S, and ITS2	ethyl acetate extract of mycelia; Trolox equivalent antioxidant content (TEAC), Total phenolic content (TPC)		Antifungal % (1 mg mL^{-1}): 29.8, 33.2, 42.4 Antibacterial MIC ($\mu\text{g mL}^{-1}$): 16–64 [23]
<i>A. italicina</i>	leaves	<i>Alternaria</i> sp. (strain Ail3)		phenolic acids, flavonoids		TEAC (μM trolox equivalent/100 mL culture) = 17/41; TPC (mg gallic acid equivalent/100 mL culture) = 13.71
		<i>Phomopsis</i> sp. (strain Ail4)		phenolic acids, flavonoids		TEAC (μM trolox equivalent/100 mL culture) = 52/63; TPC (mg gallic acid equivalent/100 mL culture) = 49/22
<i>A. laetiflora</i>	N.A.	<i>Alternaria</i> sp. (strain CYBH42)				TEAC (μM trolox equivalent/100 mL culture) = 17/12; TPC (mg gallic acid equivalent/100 mL culture) = 8.43 [22,72]
		<i>Alternaria</i> sp. (strain CYBH47)		ethyl acetate crude extract	N.A.	TEAC (μM trolox equivalent/100 mL culture) = 10/21; TPC (mg gallic acid equivalent/100 mL culture) = 8.69
<i>A. laevigata</i>	stem	<i>Cochliobolus gentianalis</i>	morphology and ITS1, 5'S, and ITS2			TEAC (μM trolox equivalent/100 mL culture) = 21/76; TPC (mg gallic acid equivalent/100 mL culture) = 21.27 [23]
		<i>Botryosphaeria dothidea</i>				Cytotoxic, antioxidant [39]
<i>A. mangolica</i>		<i>Colletotrichum gloeosporioides</i>	morphology	colletotric acid	phenolic acids	Antifungal EC ₅₀ = 0.03 % (1 mg mL^{-1}) = 31.5 and 100 [23]
		<i>Curvularia spicifera</i>	morphology and ITS1, 5'S, and ITS2	ethyl acetate crude extract	N.A.	Antifungal % (1 mg mL^{-1}): 85/79, 22.4 Antibacterial MIC ($\mu\text{g mL}^{-1}$): 25–50 [64]
<i>A. subulata</i>	N.A.	<i>Chalara</i> sp. (strain 6661)	N.A.	isofusidienol A, B, C, and D	chromones (benzopyran derivatives)	Antifungal % (1 mg mL^{-1}): 8/95, 52/09, 77.9 Antibacterial [23]
		<i>Curvularia spicifera</i>				[79]
EF—endophytic fungi; MIC—minimum inhibitory concentration; %I—percentage of inhibition, TEAC—trolox equivalent antioxidant capacity; TPC—total phenolic content; where both %I and EC ₅₀ are presented, it means that results were expressed as such depending on the pathogenic strains tested and conditions employed. InChI keys are provided as supplementary materials.						

4. Conclusions

Artemisia is a large genus of perennial and annual plants with species producing a variety of interesting and active compounds. The early exploration of its endophytic communities indicates diversity with promise of further significant biological activity. Research of the endophytes associated with the genus has revealed both cosmopolitan and rare endophytic species with highly studied chemical profiles, active molecules, and developed in vitro culture techniques.

Supplementary Materials: The following are available online at <http://www.mdpi.com/2309-608X/4/2/53/s1>.

Author Contributions: A.C. and R.C. conceived, designed, and wrote the manuscript.

Acknowledgments: We thank Kanika Chowdhary (TERI Institute, New Delhi) for reviewing the English language of the manuscript. We deeply appreciate Matias Reina Artiles (IPNA—CSIC, Tenerife, Spain) and two unknown reviewers whose valuable considerations have improved this survey. Also, we are grateful to Alin Gayraud who helped with the graphical abstract.

Conflicts of Interest: The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; and in the decision to publish the results.

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Review

Endophytic Mycoflora and Their Bioactive Compounds from *Azadirachta Indica*: A Comprehensive Review

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Received: 3 February 2018; Accepted: 22 March 2018; Published: 24 March 2018

Abstract: Plants are all inhabited by endophytic fungi in the interior of their tissues. The neem tree *Azadirachta* is an Indian lilac used for various therapeutic purposes in different forms of preparations. This plant hosts different types of endophytic fungi. In some cases, different tissues of a given plant are inhabited by different endophytic fungi which are discussed in this paper. Recently, there have been new reports on endophytic fungi and their bioactive compounds from *Azadirachta indica*. The biological function of bioactive compounds was discussed in view of their future industrial prospects. There are a number of different research investigations that examine the endophytes isolated and screened for their potential bioactive secondary metabolites from neem, but there is no comprehensive review on neem endophytes and their secondary metabolites to bring all trends from different researchers together. Therefore, in this review, we have discussed the endophytic fungi from the different tissues of neem, in view of the latest understandings of antimicrobial, antioxidant, and pathogenicity target compounds. Importantly, tracing the previous findings would pave the way to forecast the missing link for future work by researchers.

Keywords: endophytic fungi; plant tissues; bioactive compounds; biological activities

1. Introduction

The term “endophyte” refers to all microorganisms that colonize internal plant tissues for all or part of their lifetime [1]. They cause unapparent and asymptomatic infection and live entirely within plant tissues. They cause no symptoms of disease [2]. Endophytes are the chemical synthesizers inside plants and plants have been extensively investigated for their endophytic microbial complement [3].

Endophytes are an under-investigated group of microorganisms that represent a plentiful and renewable source of bioactive and chemically new compounds with potential for exploitation in a wide variety of medical, agricultural, and industrial realms [4]. They are a taxonomically and ecologically heterogeneous group of organisms; mainly belonging to Ascomycota, coelomycetes, and hyphomycetes [5–8].

All plants are inhabited internally by diverse microbial communities comprising bacteria, archaea, fungi, and protista [1,9]. The association between fungal endophytes and their host plant is due to the result of unique adaptations which enable the endophytes to harmonize their growth with that of their host [10]. These endophytic communities are accountable to either partial or complete biosynthesis of host plant secondary metabolites [8,11].

Endophytic biotechnology can be used for the efficient production of agriculturally, industrially and economically important plants and plant products. The rational application of endophytes to manipulate the microbiota, intimately associated with plants, can help in enhancement of production of

the agricultural product, increased production of key metabolites in medicinal and aromatic plants, as well as adaption to new bio-geographic regions through tolerance to various biotic and abiotic conditions [12].

Endophytes have recently generated significant interest in the microbial chemistry community due to their great potential to contribute to the discovery of new bioactive compounds. It has been suggested that the close biological association between endophytes and their plant host results in the production of a great number and diversity of biologically active molecules compared to epiphytes or soil-related microbes [13]. Moreover, the symbiotic nature of this relationship indicates that endophytic bioactive compounds are less toxic to the cell, as these chemicals do not kill the eukaryotic host system. This is particularly important to the medical community as potential drugs may not adversely affect human cells [14].

Azadirachta indica is widely used for the investigation of endophytes and their secondary metabolites, but it remains crucial to further extend the study as it is one of the major traditional medicinal plants used by about 80% of developing nations. This is because the discovery of new bioactive compounds as well as new endophytic fungi from *Azadirachta indica* has been reported recently. Hence, this review discusses the latest research advances on endophytic fungi from the neem tree, including types of endophytic fungi identified, described, bioactive compounds, and biological activities of endophytes to trace the current scope of knowledge of endophytes. It gives collective recent knowledge on antimicrobial activities of endophytic fungi of neem and on recent advances concerning their efficacy as potential antimicrobials to fight the current health problems due to emerging multidrug-resistant pathogens. It also gives focus to the well-studied endophytic sources of potential antimicrobials with emphasis on their future prospects of endophytes on human pathogens based on the recent research outputs.

The chance of getting new bioactive compounds from endophytes depends on several factors, for example, culture condition, tissue type, and age of host, the degree of interaction of endophytes with their photobionts in nature, the genetic basis of mycobionts and photobionts. Leaving aside most of these factors, by modifying the culture condition different investigators are reporting new endophytic isolates [15] and new bioactive compounds in recent studies. This signifies the continuous nature of exploiting the endophytes even from largely exploited hosts as well. Importantly, to our knowledge, there is no review on the endophytic fungi and their bioactive compounds from neem plants. Therefore, the need to review the endophytic fungi and their bioactive compound from *Azadirachta indica* is thus apparent. Ultimately, we believe that this review provides relevant research trends on the subject.

1.1. Plant-Endophyte Metabolism

Plant and endophyte are under continuous interaction in nature in their symbiotic existence. Their metabolism can interact on many levels: (a) the endophyte induces host metabolism, (b) the host induces endophyte metabolism, (c) host and endophyte share parts of a specific pathway and contribute partially, (d) the host can metabolize products from the endophyte and vice versa, (e) the endophyte can metabolize secondary compounds from the host. The two latter possibilities can be accomplished by only one, several or all enzymatic steps for biochemical transformation [11].

Endophytes can influence their host plant's metabolism, but one can speculate that the established host range could also alter or influence the pattern of secondary metabolites in endophytic fungi. Moreover, the host plants can influence the metabolite pattern in pathogenic fungi [11].

1.2. Effects of Endophytes on Neem Plants

Endophytes form a symbiotic relationship with their plant host. It is believed that in many cases the microbes function as the biological defense for the plant against foreign phytopathogens due to the fact that the majority of secondary metabolites occurring in endophytic microorganisms have antimicrobial activity and these have been implicated in protecting the host plant against [16]. The protection mechanism of the endophytes are exerted directly, by releasing metabolites to attack any antagonists or lyse affected cells, and indirectly, by either inducing host defense mechanisms or

promoting growth [14]. Antibiotics or hydrolytic enzymes can be released by endophytes to prevent colonization of microbial plant pathogens [13,17].

2. Endophytic Fungi from *Azadirachta indica*

Different tissue samples of a given plant in several locations yield the greatest species diversity of endophytes. *Periconia*, *Stenella*, and *Drechslera* were endophytes reported from neem plant [6]. Moreover, the authors of [6] reported *Phomopsis oblonga*, *Cladosporium cladosporioides*, *Pestalotiopsis* sp., *Trichoderma* sp., and *Aspergillus* sp. as the dominant endophytic fungal isolates. The huge and substantial collection of endophytic fungi from neem may represent a peculiar source of the interesting and useful bioactive compounds associated with *Azadirachta indica* such as the azadirachtins and related tetranortriterpenoids [6].

The diversity of endophytic fungi is likely determined by the age of the plant parts [18]. Some endophytes are isolated only from certain specific locations irrespective of the difference in tissue specificity [19]. Therefore, in this section, endophytic fungi from different tissues of *Azadirachta indica* are reviewed and presented in Table 1.

2.1. Endophytic Fungi of Leaf

The leaf is an ideal tissue for exploring the endophytes and their secondary metabolites. The leaf has shown the highest report on species richness and colonization of endophytes [6]. The authors of [6] reported the occurrence of five fungal endophytes from asymptomatic green and senescent leaves of neem plant. From these, four were sterile forms and one was *Fusarium avenaceum*. The frequency of occurrence of endophytes was significantly higher in the basal leaflets than in the apical or middle leaflets and in the main vein of the leaflet than in the lamina tissue. The colonization frequency of green leaves by endophytes increased during the rainy season although no new endophyte species could be recovered by their study. The antifungal metabolites present in the leaves could be accounted for by the restricted number of endophytic fungal genera and the absence of common endophytic fungi in the neem leaves. The occurrence of foliar endophytes in tropical trees is influenced by environment, types and chemistry of the host tissue [20], and the nature of interaction between the mycobiont and phycobiont. They have isolated and recorded *Fusarium avenaceum* with some other sterile mycelia. Another study conducted by [21] isolated four endophytic fungal species (*Trichoderma* sp., *Colletotrichum* sp., *Curvularia* sp. and *Chaetomium* sp.) and *Alternaria alternata* [22] from the leaves of *Azadirachta indica* A. Juss.

In most studies of endophyte fungi, the sterile mycelia fungi are reported as mycelia-sterile. Unlike those, the three mycelia-sterile isolated were identified as *Fusarium solani* and *Chaetomium globosum*. Additionally, leaf isolated mycelia-sterile was identified as *F. solani* (soil fungus) by Verma [7], and it shows the vertical traveling nature of this fungus from root to upper tissues of the host by the same study.

Tenguria and Khan [23] have isolated 85 endophytic fungi of 10 genera from 200 segments of fresh *Azadirachta indica* leaves. These are *Chaetomium globosum*, *Pestalotiopsis* spp., *Phoma* sp., *Aspergillus flavus*, *Aspergillus niger*, *Alternaria alternata* (Fr.) Keissl., *Fusarium* spp., *Penicillium* spp., *Trichoderma* spp., and Sterile mycelia. Their isolates are almost similar to fungi reported by Taware & Rajurkar [24]. The endophyte *Geotrichum* sp. AL4 was cultivated from the leaf of *Azadirachta indica* and produced novel active components with nematicidal activities [25]. Abubakar and Ndana [26] have isolated five species of endophytic fungi from leaf tissue of *Azadirachta indica* with *Cladosporium* spp. dominated the leaves with the frequency of colonization of 11.3%.

2.2. Endophytic Fungi of Stem

The endophytic fungus *Xylaria* sp. YM 311647 was isolated from stems of healthy *Azadirachta indica* collected in China [27]. Another endophytic fungus *Phomopsis* sp. YM 311483 was isolated from surface-sterilized fresh stems of an apparently healthy *Azadirachta indica* specimen collected in China.

This fungus produced two new ten-membered lactones which have antifungal potential against plant pathogenic fungi [28].

Wu et al. [29] have isolated and reported the fungal genus *Xylaria* sp. YC-10 from stems of *Azadirachta indica* A. Juss. Abubakar and Ndana [26] have isolated seven species of endophytic fungi from stems (twig) tissue of *A. indica* with *Cryptococcus* spp. reported being the most dominant isolate with high percentage frequency of colonization 13.9%.

2.3. Endophytes of Fruit

The endophytic fungi from *A. indica* are mostly known from leaves, bark, and stems but it is uncommon from fruits of this host. Verma et al. [7] have isolated endophytic fungi from unripe fruit and roots. They have isolated 105 endophytic fungi from these unripe fruits, from 29 taxa, at a rate of 68.0%. Their isolates were composed of 11.06% mycelia sterilia, 7.25% coelomycetes, and 81.69% hyphomycetes, while *Humicola*, *Drechslera*, and *Colletotrichum* sp. were obtained exclusively from fruit samples.

2.4. Endophytic Fungi of Bark

Mahesh et al. [5] have investigated the endophytic fungi from the inner bark of *A. indica*. They have isolated 77 endophytic fungal isolates belonging to 15 genera composed of hyphomycetes, coelomycetes, ascomycetes and sterile mycelia (Table 1). The endophytes *Curvularia*, *Cochlonema*, *Gliomastix* and *Verticillium* spp. were reported in their study. Most importantly, *Trichoderma*, *Penicillium* and *Pestalotiopsis* spp. were the most dominant endophytes recorded. Verma et al. [6] have isolated and characterized 18 different taxa of endophytic fungi from leaf, stem, and bark of *Azadirachta indica* A. Juss. Their isolates are composed of 62.2% hyphomycetes, coelomycetes (27.4%) and mycelia sterilia (7.7%). Importantly, the endophytic fungi occur more often in leaf segments than stem and bark tissues [6].

Tejesvi et al. [19] have isolated the endophytic fungi *Gliomastix*, *Curvularia*, *Phoma eupyrina*, and *Phyllosticta* from the inner bark of *A. indica*. In another investigation, they have reported three isolates of *Pestalotiopsis microspora* and two isolates of *Bartalinia robillardooides* from the inner bark of *A. indica* [30].

2.5. Endophytic Fungi of Root

The endophytic fungus *Chloridium* sp. was isolated from the roots of *A. indica* A. Juss. in Varanasi, India. The fungus produces a highly functionalized javanicin, with promising antibacterial activity [31] that might prevent its host from invasion by pathogenic microorganisms.

Verma et al. [7] have isolated 167 endophytic fungi from roots, at a rate of 68.0%. Their isolates are identified into 29 taxa; those composed of mycelia sterilia accounted for 11.06%, coelomycetes 7.25%, while hyphomycetes showed the maximum number of representative isolates (81.69%). They have recorded species such as *Chaetomium globosum*, *Chloridium*, *Scytalidium*, *Nigrospora* and *Verticillium* exclusively from the root. Kusari et al. [32] have reported the isolation and characterization of a novel endophytic fungus *Eupenicillium parvum* in their study.

2.6. Endophytic Fungi of Seed and Twigs

Although endophytic fungi isolated from different tissue of plants in several investigations, the reports of endophytic fungi from twigs of neem plant, unlike from other plants, are very few. Ashkezari and Fotouhifar [15] have isolated nineteen new endophytic fungi from a common yew plant. The pathogenic endophyte *Phomopsis azadirachiae*, the cause of devastating die-back disease, were reported from seeds and twigs of neem plants [33]. It is important to note that the twigs of *Azadirachta indica* are a poorly exploited surface for investigation of endophytes. Therefore, it is imperative to conduct further study on the endophytic fungi and their bioactive prospects in the search for potential applications.

Table 1. Endophytic fungi from different parts of neem plant.

Division (Subdivision)	Class	Subclass	Order	Family	Genus (Former Name)	Identified by	Plant/Tissue	References
Aскомицота (Pezizomycotina)	Sordariomycetes	Hypoconymycetidae	Hypocreales	Nectriaceae	<i>Fusarium</i> <i>Trichoderma</i> <i>Gliomastix</i>	ITS-5.8S rDNA analysis LSU, SSU, TEF and RPB2 sequence data Phylogeny and DNA-based identification	R,L,S,B,F R,L,S,B,F L,B	[6,7,19,34–36] [7,36] [5,6,19,36]
	Glomerellales	Plectosphaerellaceae			<i>Acremonium</i>	DNA-based identification	R,B,F	[7,19,37]
Sordariomycetidae	Sordariales	Glomerellaceae			<i>Verticillium</i>	LSU, SSU, TEF and RPB2 sequence data	R	[7,36]
	Chaetomiaceae	Chaetomiidae			<i>Colletotrichum</i> <i>Chaetomium</i> <i>Humicola</i>	ITS-5.8S rDNA analysis	R,L,F	[7,21]
Xylariomycetidae	Xylariales	Chaetosphaeriaceae			<i>Chloridium</i> <i>Xylaria</i>	ITS-5.8S rDNA analysis	R,L,B	[7,19,23]
		Xylariaceae			<i>Pestalotiopsis</i>	ITS	F	[7,36]
		Bartaliniaeae			<i>Bartalina</i>	Phylogenetic analysis; Molecular clock analysis	R	[7]
Dothideomycetes	Diaporthales	Trichocomphorales	Trichocomphoraceae	Diaporthaceae	<i>Nigrospora</i> <i>Diaporthe</i> (<i>Phomopsis</i>) <i>Scutellatium</i>	Molecular clock analysis; Molecular clock analysis; ITS rDNA analysis	S	[29,34–36]
Leotiomycetes					<i>Aspergillus</i> <i>Penicillium</i> <i>Eupenicillium</i>	ITS-5.8S rDNA analysis	B,L,S	[5,6,23,30,36]
Eurotiomycetes					<i>Curularia</i> <i>Alternaria</i> <i>Drechslera</i> <i>Phoma</i>	six-gene phylogeny six-gene phylogeny five-gene phylogeny ITS-5.8S rDNA analysis	R,S,L,B S,L,B	[6,7,36] [6,34,36,40]
Pleosporales					<i>Peronospora</i>	ITS-5.8S rDNA analysis	R	[7]
Dothideomycetes					<i>Phoma</i>	ITS-5.8S rDNA analysis	R,S,L,B	[34]
					<i>Phoma</i>	ITS-5.8S rDNA analysis	R,F	[7]
					<i>Peronospora</i>	ITS-5.8S rDNA analysis	R,S,L,B	[41,42]
Capnodiales	Davalliales				<i>Cladosporium</i> <i>Senella</i> <i>Phyllosticta</i> <i>Cryptococcus</i> <i>Cochlonema</i> <i>Cercinella</i>	ITS-5.8S rDNA analysis	R,L,B,F	[7,43,44]
		Boryosphaeriales	Botryosphaeriaceae			ITS-5.8S rDNA analysis	B	[6]
Zygomycota (Zoopagomycotina)	Tremellomycetes	Tremellomycetidae	Filiobasidiidae	Zoopagales	<i>Cochlonemataceae</i> <i>Mucoraceae</i>	six-gene phylogeny Phylogeny	R,F	[7,39–45]
						ITS-5.8S rDNA analysis	S	[34,46]
							B	[5,19,46]
							R	[7]

Note: "R": root; "L": Leaf; "B": Bark; "S": Stem; "F": Fruit. The endophytes are isolated from the tissue types.

3. Current Trends on Bioactive Metabolites from Neem Endophytic Fungi

The production of antimicrobial bioactive compounds by endophytes is currently receiving urgent concern due to the emergence of multidrug-resistant pathogens in chemotherapy. Unlike the chemical synthetics, due to health and environmental issues, there is a strong need of endophytic sources of antimicrobials for their biological and chemical safety effects. There exist many appreciable possibilities for the exploitation of endophytic fungi for the generation of abundant novel biologically active secondary metabolites. Secondary metabolites from microorganisms have proven to be tremendous and enduring sources [47]. The endophytes isolated from the medicinal plants have the potential as an alternative source of these bioactive secondary metabolites [48]. Some of the bioactive compounds produced by endophytic fungi of the neem tree and their biological activities are discussed hereunder.

3.1. Melanin Pigment

Diaporthe (former name: *phomopsis*) is a phellophytic fungus that lives in the outer layer of the bark [49]. Found in *Azadirachta indica*, it synthesized and deposited DOPA (3,4- dihydroxyphenylalanine) type of melanin on their hyphae. Melanin produced by the endophyte, *Phomopsis*, provides adaptations, which include mechanisms for overcoming host barriers, successful competition with other phylloplane fungi and surviving harsh environmental conditions [50] and in the widespread occurrence of the endophyte [51].

Melanin pigment production is catalyzed by the enzyme tyrosinase produced by endophytic fungi *Fusarium* spp. from *Azadirachta indica*. The fungi produced 2.8 U/mol (L-DOPA) extracellular tyrosinase activity. This enzyme is essential for pigmentation, an important factor in wound healing and primary immune response. It has a great application in environmental, pharmaceuticals, cosmetics and food industries [52].

Even though endophytes are known for their antibacterial, antitumor, immunomodulatory, anti-inflammatory and antiviral activities, there are few reports on endophytes for tyrosinase production from neem plants [52]. However, extracellular production of fungal tyrosinase was reported, firstly, from a filamentous fungus, *Trichoderma reesei* [53]. Therefore, future research should focus on investigating tyrosinase from neem endophytes due to its higher industrial applications.

3.2. Antioxidant Metabolites

Kumaresan et al. [21] have screened endophytic fungi, from the leaf, for their antioxidant bioactive compounds. They have investigated the constituents of antioxidants like phenols, tannin, flavonoid, ascorbic acid, and β -carotene. All isolated endophytic fungi studied by their team possessed in vitro antioxidant activities. Similarly, evaluation of secondary metabolites from *Penicillium* species revealed the presence of saponins in addition to those mentioned above [54]. This is one of the very few reports from neem endophytes for their antioxidant activities. Another study by Zhao et al. [55] reported the natural antioxidant cajaninstilbene acid ($100.5 \pm 9.4 \mu\text{g/g}$ dry weight of mycelium) from endophytic fungi *F. solani* isolated from pigeon pea. This *Fusarium* was also found in neem tree as mentioned in Table 1.

3.3. Antimicrobial Secondary Metabolites

Wu et al. [56] reported two solanapyrone analogues (solanapyrones N and O) compounds isolated from the fermentation culture of *Nigrospora* sp. YB-141, an endophytic fungus isolated from *Azadirachta indica* A. Juss growing in the tropical region of Southwest China.

Verma et al. [57] have screened and evaluated the efficacy of six fungal endophytic strains from *Azadirachta indica* A. Juss against dermatophytic fungi *Trichophyton* and *Microsporum*. They have reported production of active compounds by their isolates but identification of the compound is not yet reported. Similarly, evaluation of secondary metabolites from *Penicillium* species isolated from the leaf of neem plant showed that they possessed some degree of antibacterial and antifungal activities [54].

Five 10-membered lactones were isolated from the endophytic fungus, *Phomopsis* sp. YM 311483, obtained from the stem of *Azadirachta indica*. Of these, two compounds (8R-acetoxy-5R-hydroxy-7-oxodecan-9-olide and 7R-acetoxymultiplolide A) were newly recorded by Wu et al. [28]. Two compounds identified as 7R,8R-dihydroxy-3,5-decadien-10-olide and 8R-acetoxymultiplolide A are identical to compounds reported by Tan et al. [58]. All the compounds were evaluated for their antifungal activity against seven plant pathogens, *Aspergillus niger*, *Botrytis cinerea*, *Fusarium avenaceum*, *Fusarium moniliforme*, *Helminthosporium maydis*, *Penicillium islandicum*, and *Ophiostoma minus*. Compound 8R-acetoxymultiplolide A showed the most potent antifungal activities with MIC values in the range of 31.25–500 µg/mL. Interestingly, 8R-acetoxymultiplolide A was more potent [58] than 7R-acetoxymultiplolide A even though their structures differed only in the position of the acetoxy substituent [40]. On the contrary, Tan et al. [58] have reported that compounds 7 α ,8 α -Dihydroxy-3,5-decadien-10-olide and 8 α -Acetoxymultiplolide A have not shown any antifungal activities against *Candida albicans*. Therefore, further antimicrobial activity bioassay is deemed necessary to reconcile the discrepancy of the report and to utilize the potential of the isolates and their compounds.

Five guaiane sesquiterpenes isolated from the culture broth of endophytic fungus *Xylaria* sp. YM 311647 showed moderate or weak antifungal activities in a broth microdilution assay against five pathogenic fungi, but no obvious inhibitory activities against *Fusarium avenaceum* [27]. Nine oxygenated guaiane-type sesquiterpenes and three isopimarane diterpenes were reported from this fungus, by Tejesvi et al. [30]. Suggested by Wu et al. [59], 18-norisopimarane diterpene and the diterpene sulfate are effective against human pathogenic fungi.

The extracts of *Pestalotiopsis microspora* isolated by Tejesvi et al. [30] demonstrated the potential of antibacterial activities as therapeutic agents against various pathogens tested. The same species from different isolates by the same work demonstrated antioxidant and antihypertensive properties. From this, it is possible to deduce that this isolate needs to be thoroughly investigated for its commercial use as an antioxidant or antihypertensive agent for the treatment of the respective diseases.

An investigation of the secondary metabolites from *Chloridium* sp. root isolates by Kharwar et al. [31] resulted in the identification of javanicin. This compound was identical in all respects to a previously described naphthaquinone, javanicin. However, its crystallized structure was confirmed by X-ray crystallography. Javanycin was tested for its antimicrobial potential against human and plant pathogens. Tests demonstrated that javanicin had antimicrobial potential against human and plant pathogens. It was either slightly active or not active against fungi such as *Pythium ultimum*, *Phytophthora infestans*, *Botrytis cinerea*, and *Ceratocystis ulmi*, whereas it was active against *C. albicans*, *Escherichia coli*, *Bacillus* sp., and *Fusarium oxysporum* at higher MIC values ranging from 20 and 40 µg/mL. Its activity was recorded against *Rhizoctonia solani* and *Verticillium dahliae* at 10 µg/mL while it was active at 5 µg/mL against *Cercospora arachidicola*. The bacteria that were the most sensitive to the javanicin (2 µg/mL) were *P. aeruginosa* and *P. fluorescens*. On account of this, it may be ideal for compounds, particularly for valuable selective antibiotic formulations for human and plant pathogens [31].

3.4. Natural Insecticides

The production of natural insecticides, azadirachtin A and B, from endophytic fungus *Eupenicillium parvum* was reported by Kusari et al. [42]. These compounds are exclusive to the neem tree, *Azadirachta indica* A. Juss, from where they are currently originated. The production of this compound by the fungus is not only in the mycelium but also released into the culture media. The discovery of this azadirachtin-producing endophytic fungus has vast implications for further research from the ecological and biochemical point of view. Another study conducted by Kaur et al. [22] reported antifeedant and toxic activity against tobacco caterpillar *Spodoptera litura* by *Alternaria alternata*. The larvae fed on the diet supplemented with fungal extract significantly reduced the survival and influence the development and reproduction of *S. litura* due to potent toxicity of *A. alternata*. Additionally, the fungus was known for its immunomodulatory effects on *Spodoptera litura* [60].

The genus *Xylaria* sp. YC-10 isolated from stems of *Azadirachta indica* A. Juss. produced eleven compounds from which eight compounds (5-Methylmellein, 5-Carboxymellein, Hymatoxin C, Hymatoxin D, Halorosellinic acid, Cerebroside C, Cerevisterol and (2S,3S,4R,2'R)-2-(2'-Hydroxytetracosanoylamino)-octadecane-1,3,4-triol) exhibited weak insecticidal activity against *Plutella xylostella* [29].

3.5. Nematicides

Li et al. [25] isolated four compounds from the endophytic fungal strain *Geotrichum* sp. AL4, cultivated from the leaves of the neem tree. They reported two compounds, chlorinated oxazinan derivate (1-[(2R*,4S*,5S*)-2-chloro-4-methyl-1,3-oxazinan-5-yl] ethenone) and an epimer of the former (1-[(2R*,4S*,5R*)-2-chloro-4-methyl-1,3-oxazinan-5-yl] ethanone) and two other known compounds. The two compounds reported by their team were assessed for nematicidal activities against the nematodes *Bursaphelenchus xylophilus* and *Panagrellus redivivus*, and showed noticeable bioactivities. Of known compounds, compound 1-(2,4-dihydroxyphenyl)-ethanone is commonly isolated from plants. However, they have reported, for the first time, this compound from a microbial source showing nematicidal activity. Therefore, the *Geotrichum* sp. AL4 can be an ideal source for the formulation of antinematicidal agents.

3.6. Antiparasites

Verma et al. [61] have investigated several endophytic fungal strains from *Azadirachta indica* A. Juss and came up with the endophytic fungal strain *Pestalotiopsis* sp. with a significant anticestodal potential against hydatid cysts *Echinococcus granulosus*. The anticestodal activity observed with *Pestalotiopsis* sp. showed promising scolicidal activity of up to 97% mortality within 30 min of incubation. Since this was the first report that dealt with fungal endophytes for the anticestodal potential activities, further studies to reaffirm its anticestodal biopotential activities through optimized culture condition are thus needed.

4. Conclusions

Currently, the demand for health services is growing dramatically, particularly in developing countries, due to the emergence of drug resistance by pathogenic microorganisms. Moreover, the highly-increased recurrence of cancer and other infectious diseases makes the situation a greater tragedy. Hence, it is urgently necessary to investigate new bioactive compounds effective against drug-resistant pathogens for the remedy of the aforementioned diseases. Therefore, endophytic fungi, producer of a wide array of secondary bioactive metabolites with their peculiar potential compounds, namely, melanin, antimicrobials, antioxidant, anti-inflammatory, insecticides, nematicides, etc., are the ideal targets. These bioactive compounds have a range of potential to combat etiologic agents of plants and animal's disease, prevent cell damage due to reactive oxygen species, crop pests, animal pests, and to target pathogenicity traits of pathogenic microbes for necessary remediation.

To date, very few structures of bioactive compounds are characterized and identified from endophytic fungi of neem, but most of the investigations are restricted to the level of fungal identification and bioactivities assay. The different types of Ascomycota, Basidiomycota and Zygomycota endophytes found in the neem tree are phylogenetically synthesized using modern taxonomy from different systematic reviews (Table 1). Some of the endophytic isolates which failed to sporulate, sterile mycelia, are placed to their natural group based on modern taxonomy. Most importantly, the uncultured endophytic fungi from the neem tree were not reported in the literature surveyed so far. As a result, non-culturable endophytes have not been looked for with a metagenomics method. Metagenomics approaches, if used for neem tree, will allow identifying uncultured endophyte fungi that are ignored when isolation and cultivation of endophytes are done before identification. Similarly, too many authors do not make sufficient use of the modern taxonomy based on phylogenetic analysis for identification of their endophytic isolates. The compounds are

identified by few investigators and this may be due to a lack of advanced equipment, but possible cooperation with the institutions with the tools are crucial for further discovery.

Even though there are several records on the antioxidant activities of endophytic fungi from different plant hosts, very few endophytes having these effects are known from *Azadirachta indica*. However, this host is widely used for its various biological activities, as antioxidant properties, in different forms of the formulation. Therefore, more work on the antioxidant activities of its endophytes can exert a profound effect on the search for novel antioxidant bioactive compounds.

Acknowledgments: The authors are thankful to Mangalore University for providing access to necessary resources. The departments of Biosciences and Applied Botany are acknowledged.

Author Contributions: Eyob Chukalo Chutulo—conceived of the presented idea. Wrote the manuscript with support from the idea of Raju Krishna Chalannavar. Eyob Chukalo Chutulo took the lead in writing the manuscript. Raju Krishna Chalannavar—Encouraged Eyob Chukalo Chutulo to investigate [a specific aspect] and supervised the findings of this work. Supervised development of work, helped in data interpretation and manuscript evaluation and acted as the corresponding author. Both authors discussed the results and contributed to the final manuscript.

Conflicts of Interest: The authors declare no conflict of interest.

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Review

From Concept to Commerce: Developing a Successful Fungal Endophyte Inoculant for Agricultural Crops

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Received: 10 January 2018; Accepted: 9 February 2018; Published: 11 February 2018

Abstract: The development of endophyte inoculants for agricultural crops has been bedevilled by the twin problems of a lack of reliability and consistency, with a consequent lack of belief among end users in the efficacy of such treatments. We have developed a successful research pipeline for the production of a reliable, consistent and environmentally targeted fungal endophyte seed-delivered inoculant for barley cultivars. Our approach was developed de novo from an initial concept to source candidate endophyte inoculants from a wild relative of barley, *Hordeum murinum* (wall barley). A careful screening and selection procedure and extensive controlled environment testing of fungal endophyte strains, followed by multi-year field trials has resulted in the validation of an endophyte consortium suitable for barley crops grown on relatively dry sites. Our approach can be adapted for any crop or environment, provided that the set of first principles we have developed is followed. Here, we report how we developed the successful pipeline for the production of an economically viable fungal endophyte inoculant for barley cultivars.

Keywords: barley; commercialisation; crop wild relatives; endophytic fungi; yield enhancement

1. Introduction

The use of microorganisms to improve agricultural crop performance has a long history, but only in the last few decades have we been able to describe in any detail how bacteria, fungi, algae and protozoans interact with plants to enhance desired traits [1–3]. Mankind has understood the benefits of various practices, often without conscious intent, that directly or indirectly involve the manipulation of the soil and plant microbiota [4,5]. Only in the last century have we incorporated the use of microorganisms in agriculture in a deliberate, focused and knowledgeable way [6,7]. However, it is the plant that is the ultimate arbiter of how and when these microorganisms are incorporated into the functional plant microbiome, often in an unpredictable fashion [8,9]. It is the goal of all plant-microbiome research to understand and use these relationships in the most effective way, but we are still some way from obtaining this full understanding [10,11]. Such a lack of informed and validated knowledge has resulted in inconsistent results when using microorganisms in the field situation, resulting in a reduced end-user confidence and uptake of the technology [12,13].

Endophytes are a class of plant-associated microorganisms that have shown particular promise in agriculture [14–21]. Endophytes (bacteria, fungi and unicellular eukaryotes) live at least part of their life cycle inter- or intracellularly inside plants, usually without inducing pathogenic symptoms. This can include competent, facultative, obligate, opportunistic and passenger endophytes (a passenger endophyte enters the plant by accident in the absence of selective forces maintaining it in the internal tissue of the plant [22]). Endophytes can have several effects on plants and/or may change function during their life-cycle [14]. Bacterial and fungal endophytes have shown promise as beneficial crop

inoculants, and many are known to enhance abiotic and biotic plant stress tolerance [23–28]. But a large proportion of the associated studies have been conducted in a controlled environment and do not translate successfully to the field [29].

The approach adopted by our research group has consciously focused on making this transition from ‘pot to plot’ more successful. We aimed to remove the inconsistency from endophyte application in field crops by targeting a single crop, barley, as our model plant, and testing the efficacy of endophyte strains under various environmental stresses, culminating in extensive field trials [30–35]. Only with this field validation can we say with any confidence that a microbial inoculant really “works” [36]. We have demonstrated the effectiveness of this approach by developing a fungal endophyte inoculum that consistently increases barley grain yield over several seasons under a variety of chemical fertilizer inputs in dry growing environments [31]. The outcomes from our research and field application have been so successful that we have been able to bring our endophyte technology into commerce, which will result in significant reductions in economic and environmental costs.

This article will review the development of this technology, outlining the concepts involved and the methods employed to create a viable, consistent and environmentally targeted commercial endophyte inoculant for barley crops. We will discuss a set of first principles that are important for success, and we will outline future research directions that will increase the portfolio of endophyte inoculants and expand the scope of target crop species.

2. Concept

Before any resources are allocated to an endophyte research project, it is important to understand the problem that is being addressed and the solution that is proposed. The problem must be a real commercial and/or functional need of the eventual user of the solution. Nearly half of all company start-ups fail because there was no market need for the product [37]. So it is essential that the proposed concept is going to provide a viable and needed solution. This solution should also attract potential commercial partners and collaborators on the project, thus increasing the resources available for successful implementation. For illustration, we focused on the need to reduce the high economic cost of chemical crop inputs; any product that successfully addresses this need would provide real and tangible benefits for the crop growers and also reduce environmental impact of the chemical inputs.

Crop wild relatives are an important genetic resource for breeding desired plant traits into related crop cultivars [38]. We suspected that the same was true for microbial resources, and particularly for fungal endophytes. Our hypothesis was that endophytes that were recoverable from crop wild relatives would be more compatible with the related crop, both for inoculation and colonization purposes and for *in planta* behaviour. We further hypothesized that endophytes recovered from a particular type of environment would be more compatible and effective on crops growing in a similar environment, and that endophyte crop wild relative interactions were more likely to have been maintained through time than endophyte-free elite crop cultivars.

It was important to retain this focus in all subsequent experiments so that we could produce a set of results that may or may not support our hypotheses. We were also aware that to produce a comprehensive and useful dataset, with the resources available, we would need to target only one important crop species. For our purposes, barley was the ideal crop as it is the most important arable crop grown in Ireland, where it is grown on over 200,000 ha, and there are several local wild barley relatives available for sampling.

The endophyte source we selected was the wild barley species *Hordeum murinum* ssp. *murinum* L. (wall barley), as it is relatively common and occurs in a wide variety of habitats on the east coast of Ireland, especially those subject to human disturbance. This enabled us to sample from diverse environments and to establish any significant correlations between endophyte recovery and alpha/beta diversity. It also gave us the largest possible set of endophyte strains from which to screen potential study taxa. The only other naturally occurring wild barley species in Ireland are *Hordeum jubatum* and *Hordeum secalinum*, and these only occur in rare and scattered populations [39].

By targeting endophytes recovered from natural and native plant populations we avoid introducing exotic microorganisms into the environment which may face regulatory and safety issues. They would also likely be more suited to local barley crop growing conditions, as they would have been selected by historical climatic conditions. Finally, as these host plants and geographical locations have never previously been sampled for fungal endophytes, we expected that the strains would be novel and previously uncharacterized, a real advantage when there is a need to protect intellectual property.

3. Developing the Concept

3.1. Endophyte Recovery

Having selected the barley wild relative to sample, it was important to ensure that we could recover the greatest variety of endophytes for subsequent biofertilisation and biocontrol screening in agricultural applications. Here, we followed a set of general principles derived from years of experience, and which we recommend for similar research.

The most important of these was the selection of the sampling sites, as this will determine the direction and success of all subsequent work:

- Sampling site should contain a minimum population of 10 genetically individual plants
- Plants on sampling site should be healthy and free of disease
- If selected for biotic tolerance, the site should preferably be in an area of potentially high disease pressure for the species
- There should be some form of abiotic stress present at the site, such as a very dry or nutrient-poor soil, which must be measureable
- The site should be relatively undisturbed and natural, i.e., not recently worked
- The site should not be exposed to contaminants, such as close to a busy roadside
- The site should preferably contain no alien plant species
- The host plant species should be identifiable using a recognised and reliable key
- The plants should be actively growing and not in a senescent phase
- The plants should be sampled when the site is in its ‘normal’ condition, i.e., not after any heavy rain, frost or unusual weather event, nor after any disturbance

While collecting the plant material it is vital to obtain as much environmental data as possible, as this information will be the major deciding factor when selecting individual strains for consortia application at the crop growing sites. For example, endophyte strains that are sourced from a particularly dry site may be more suitable for a relatively drought-stressed crop growing environment. Recording of environmental data should include variables such as GPS location, soil pH, soil moisture, soil nutrients, soil salinity, soil type, accompanying ground vegetation, tree cover, and exposure. We also recommend collecting a soil sample onto dry ice for later analysis of soil microbial community and elemental constituents. It is also important to ensure that plant sampling is aligned with principles outlined in The Nagoya Protocol on Access to Genetic Resources and the Fair and Equitable Sharing of Benefits Arising from their Utilization to the Convention on Biological Diversity, also known as the Nagoya Protocol on Access and Benefit Sharing (ABS) [40].

For our experimental work, we chose endophyte source sites that were particularly dry, with thin sandy soils and relatively high soil pH and salinity (Table 1). These sites were chosen as we ultimately wanted to test the effects of the endophytes on field barley crops growing in similar environments; so we had the experimental goal in mind at all times from initial concept to pre-planning to field sampling. This goal-oriented approach is at the heart of our methodology, and can be applied to the selection of endophytes for other environmental conditions.

Table 1. Soil variables for endophyte recovery sites.

Location	pH	Moisture Content %	Salinity *	Soil Type
1	7.8 ± 0.0	9.1 ± 0.0	1.18 ± 0.0	Loamy silt, stony
2	8.0 ± 0.0	13.4 ± 0.0	1.41 ± 0.0	Loamy silt, shallow
3	7.6 ± 0.0	0 ± 0.0	1.22 ± 0.0	Sandy silt, shallow
4	7.9 ± 0.0	4.4 ± 0.0	1.39 ± 0.0	Sandy silt, stony
5	7.9 ± 0.0	0 ± 0.0	1.45 ± 0.0	Sandy silt, shallow
6	7.7 ± 0.0	19.5 ± 0.0	1.26 ± 0.0	Clay loam, shallow
7	7.7 ± 0.0	0 ± 0.0	1.49 ± 0.0	Sandy silt, shallow
8	7.7 ± 0.0	0 ± 0.0	1.51 ± 0.0	Sandy silt, shallow
MEANS	7.8	8.3	1.37	

* Salinity is Osmotic Pressure in bars. pH, Moisture content % and Salinity are mean values ± standard error ($n = 10$).

The plant tissue that is chosen for sampling must be disease- and blemish-free, especially when sampling roots. The sampled tissue should be placed in sealed plastic bags immediately and processed in the laboratory as soon as possible, preferably within one day of sampling. Well-established protocols can then be followed for inoculation and incubation of the plant tissue to enable maximum recovery of endophyte isolates. A general principle to follow when preparing the culture media is to use the medium concentrate at only 50% of the manufacturers' recommendation, i.e., double the proportion of water. This will reduce the chances that the endophyte will experience any osmotic shock when emerging from the plant tissue and also make the culture medium more open and accessible for explorative hyphae. We also found that adding a high proportion (50%) of autoclaved plant extract increased the number of endophyte isolates recovered [41], an effect also reported by Prior et al., 2014 [42]. This plant extract should ideally be obtained from the same source species that is being sampled for endophytes. The endophytes are thus emerging into a chemically similar environment to that of the plant interior. Emerging hyphae need to be subcultured immediately, and sometimes single spore or hyphal culturing is required.

Identification of the isolates can be carried out using a combination of standard morphological and genetic barcoding techniques with analytical tools [43]. Establishing the endophyte strain identity is the most important process in screening the isolates for future experimental work.

3.2. Endophyte Screening

The initial recovery and isolation of endophytes from the host populations is only the first step in a comprehensive screening process to select the isolates with the greatest potential as crop inoculants. We took several factors into account when selecting strains for efficacy testing with barley crops:

1. The isolate should not be related to any known human or plant pathogen. This is a crucial factor which needs careful qualifying. The degree of DNA similarity between isolate and closest match is important in deciding the identity and whether to proceed with the isolate. Genetic similarity can best be judged by comparing a standard barcoding gene from the isolate with known accessions deposited in a genetic database such as GenBank (NCBI). For our purposes we compared the nuclear ribosomal Internal Transcribed Spacer gene (ITS), as it is sufficiently discriminatory to the species level when assigning taxonomy [44–49]. If the isolate is only distantly related to a known pathogenic strain, then it may be worth pursuing but is a personal judgement call depending on taxonomic group. Fungal isolate growth and proliferation at 37 °C would also be a contra-indicator for selection.
2. The isolate must show vigorous growth on a range of substrates.
3. The isolate should produce early and copious spores (endophyte cultures that do not easily produce spores are of limited use in large-scale agricultural applications).
4. The isolate should be pure; i.e., subcultured from a single spore.

It is likely that the sampling of host populations will result in a large collection of endophyte isolates and final selection of which isolates to use in subsequent experiments will depend largely on the resources available—it may be unrealistic to test 100 s or 1000 s of individual isolates. Careful screening and selection at this point will enable a more focused approach, and the controlled environment experiments will enable a further reduction of isolates selected for field trials. So we recommend that a set of criteria based on desired strain characteristics be used to select the final experimental subjects. For example, fungal endophytes should grow readily on a broad range of substrates, be of easy culture, retain competence over many cycles of sub-culturing, and should produce copious spores/conidia at a relatively early stage after initial recovery and culture. We have found that relatively few strains pass these criteria; from over 100 recovered strains, only 12 were finally selected for further experimentation [50].

4. Proof of Concept

With the basic ideas now in place we proceeded to obtain a proof of concept for our approach. At this stage, we have a selection of identified endophyte strains collected from a particular environment. Before using these strains in experimental work, the target agronomic traits must be clear from the very start, and the statistical analyses that will be used should be established; as the eminent statistician Sir Ronald Fisher once said: ‘To consult the statistician after an experiment is finished is often merely to ask him to conduct a post mortem examination. He can perhaps say what the experiment died of [51]. There is always a danger that any post-hoc selection of statistical methods may bias the analyses in favour of the desired results [52].

For many agricultural crops, yield is often the most important trait of interest and is related to a whole plethora of influences: pathogen and disease resistance, nutrient use efficiency, photosynthetic efficiency, hormone and metabolite balance, abiotic stress resistance, etc. So it makes sense to test the effect of the endophyte inoculant on as many of these factors as possible; there is no short cut to developing an effective endophyte inoculant. The crucial point here is to focus on the main influences on crop performance that will be experienced by the target crop species in a particular environment.

To achieve a comprehensive set of data for barley-endophyte responses under different conditions, we decided to carry out extensive controlled environment experiments under a variety of stresses using the fungal endophytes recovered from the sampled populations of *H. murinum*.

Prior to the experimental cycle, a range of commercial culture media combined with whole plant extract of *H. murinum* were tested for their effect on endophyte recovery from the roots of their host (*H. murinum*), and for their subsequent growth and sporulation [41]. It was found that there were significant differences between the media in endophyte recovery, endophyte mycelial growth and time to sporulation. A significantly greater number of different endophytes were recovered from roots on the malt extract plus whole plant extract medium than any of the other media tested. These results indicated that different media are suitable either for the initial recovery and isolation of fungal root endophytes or for increasing fungal biomass and inducing earlier sporulation.

The first controlled-environment experiment with barley examined the effects of ten endophyte strains on disease development on seeds of a barley cultivar using five artificial and one soil-based growth media [30]. A co-inoculant of all ten isolates, as well as two individual isolates, successfully suppressed the development of seed-borne fungal infections on germinated and ungerminated seed. We further reported that the ability to suppress the seed-borne infections was related to the soil properties of the isolate origin [33].

Secondly, we found that inoculation with six different individual endophytes increased grain yield in a nutrient-starved barley cultivar by up to 29% [32]. Furthermore, we also showed that inoculation with the isolates induced increases of up to 70% in shoot dry weight in the nutrient-starved spring barley; the nutrient input was only 10–15% of that recommended.

Thirdly, five individual endophyte strains induced significant improvements in agronomic traits for a severely drought-stressed barley cultivar, including the number of tillers, grain yield and shoot

biomass [35]. Soil moisture content was allowed to reach 10–15% of field capacity before watering. The trait that showed the greatest significant difference in the drought-stressed plants was the number of tillers, where all of the endophyte treatments induced a greater number of tillers per plant. However, except in one case, the mean dry root weight for all plants was greater in the control plants, indicating preferential allocation of resources to above-ground parts in the endophyte treatments.

Fourthly, five fungal root endophytes, either individually or combined, were inoculated onto seeds of a barley cultivar grown in optimal conditions and under a combined drought (10–15% soil moisture), heat (33 °C), nutrient (15% of recommended input) and pathogen (*Gaeumannomyces graminis*) stress (multiply-stressed). We found a greater endophyte-induced improvement in important agronomic traits in the multiply-stressed plants compared with the plants grown in optimal conditions [53]. For the multiply-stressed plants, only 13% of the controls survived to the end of the experiment compared with 80% of the endophyte treatments. In multiply-stressed plants, the endophytes induced increases in the number of tillers and root and shoot biomass. The improvements were most significant for barley inoculated with a combination of all five endophytes.

Finally, we tested a range of seed dressings to evaluate the effects of the endophytes on germination and early growth of barley. Seeds were either untreated, dressed with a fungicide (triticonazole and prochloraz), an endophyte spore solution or a combination of endophyte and fungicide. We found significant increases in mean barley seedling length induced by the endophytes after 28 days of seedling growth at time points of up to 6 months from seed dressing. The increases in mean seedling length were greatest for the combined fungal endophyte and fungicide treatment. These results indicated that the endophytes tested were persistent in a seed dressing, enhanced early seedling growth and were fungicide tolerant [34].

Results from these experiments showed that novel symbiotic associations between barley and fungal root endophytes significantly increased yield and biomass in barley grown under nutrient, drought, heat and pathogen stress, and also suppressed the development of seed-borne pathogenic infections.

5. Translation

Isolating, selecting and testing the endophytes in controlled environments to determine their potential as crop inoculants is only the first step towards future agricultural application. We now needed to see whether the positive results translated to field-grown barley. The four endophyte strains that induced the greatest significant yield increase in nutrient- and drought-stressed barley were selected as the consortium members. We reasoned that the different modes of action associated with each of the two stresses would allow a degree of compatibility as a consortium, with each endophyte bringing different functional mechanisms to the plant in a particular ecological niche space [54–58].

To obtain the maximum benefit from the endophytes as a crop treatment, it is essential to select the most effective inoculant delivery formulation. The formulation will depend to a large extent on the type of fungal material that is to be used as the active ingredient. The method of applying the treatment should be economically and ergonomically viable, and be easy to apply. The inoculant carrier substrate should provide a stable environment for the microbial fractions and prolong product shelf-life [59]. Some microbial products have a very short shelf-life [12], with a severely reduced window of effectiveness. Our decision to dress the barley seeds with environmentally stable fungal endophyte spores helps to provide robustness and preserve vitality of the treatment, even several months after dressing [34]. Dressing the seeds using standard seed dressing equipment means that the seed supplier does not need to purchase extra and possibly expensive equipment.

Dressing the seed also ensures that the endophyte inoculant will be deployed in the most efficient way, as the inoculant is delivered precisely to where it will be most effective, with no wastage. The germinated seedling root(s) will come into immediate contact with the fungal spores, which will then colonise the plant at the earliest possible stage. This will also have the added benefit of reducing the release of the strains into the environment. The endophyte may be more effective if it colonises

the plant tissue before any competitors, as the order of arrival of endophyte and pathogen may even change the effect on the plant from beneficial to detrimental [60].

This scenario depends on a productive relationship with industry partners, and this is often the point at which transition from promising laboratory results to real-world applications fails. Unless specific funding is available to support this process then the researcher must rely on the good will of the industry partner(s). Getting the right partners in the right place at the right time will have a big effect on the success of the translational research effort. We have found it helpful to include such funding in the associated research proposal. This is particularly important when trialing field crops, as multi-year validation is required before any robust claims can be made for the effectiveness or otherwise of the endophyte treatment.

The approach we have adopted in developing a crop inoculant, as outlined in this paper, has resulted in the successful translation of promising laboratory-based research to agriculture. In field trials with barley, we found that for an endophyte-associated increase in grain yield, a strong correlation was found between increased yield and low seasonal rainfall [31]. Furthermore, the endophytes were just as effective with regular foliar fungicidal treatment. Another recently completed field trial on dry sites also returned significant endophyte-induced yield increases (manuscript in preparation), providing strong validation for the value of our methodology.

6. Discussion

Many promising scientific discoveries do not make a successful transition to commerce, for any number of reasons, while some discoveries happen almost by accident [61]. The most common point of failure may be in the initial research project design, where a focused, achievable and outcome-based plan is not in place [62]. This is especially true of research into the potential of endophytes in agricultural crops, where many valuable discoveries languish in the limbo between the laboratory bench and the farmer's field. In this paper, we have shown that a properly focused endophyte project with a continuous emphasis on one particular outcome can result in a more successful transition from idea to application. We have described a set of best practices leading to the development of a successful endophyte field crop inoculant, and this approach can, in principle, be adapted for any crop or agricultural environment (including marginal lands) provided that a similar methodology is followed. The flowchart (Figure 1) shows the basic steps and decision points of our approach.

The prospecting and discovery of endophytes that have beneficial application to agricultural crops is an ongoing and potentially open-ended enterprise, with rates of successful application only limited by available resources and unproven methodologies. Resources are finite and competitive, but ensuring that the methods and principles applied are sound will maximize the chance of success. While the use of endophytes as beneficial crop inoculants has potential, the solution to a defined agricultural need is often complex and multi-factorial and it is unlikely that a single approach will provide the full answer. Future research directions should focus on combining different approaches that will provide more robust and persistent solutions for agriculture. For instance, an endophyte crop treatment combined with a change in agricultural practice can give extra benefits. For example, moving towards a 'no-till' method of cultivation along with longer rotations and the use of cover crops can drastically reduce chemical inputs and labour costs [63] and, if combined with an endophyte inoculant, may produce surprising synergistic results.

Global climate change will bring new and greater stresses to bear on crops, and future research efforts should focus on the solutions to the expected changes. It may be best to target expected conditions in particular geographic areas experiencing particular stresses, especially in poorer areas where farmers cannot afford to use expensive chemicals. Endophyte-based solutions to stress-related problems that minimize economic and environmental costs have the potential to bring real benefit to those who need it the most.

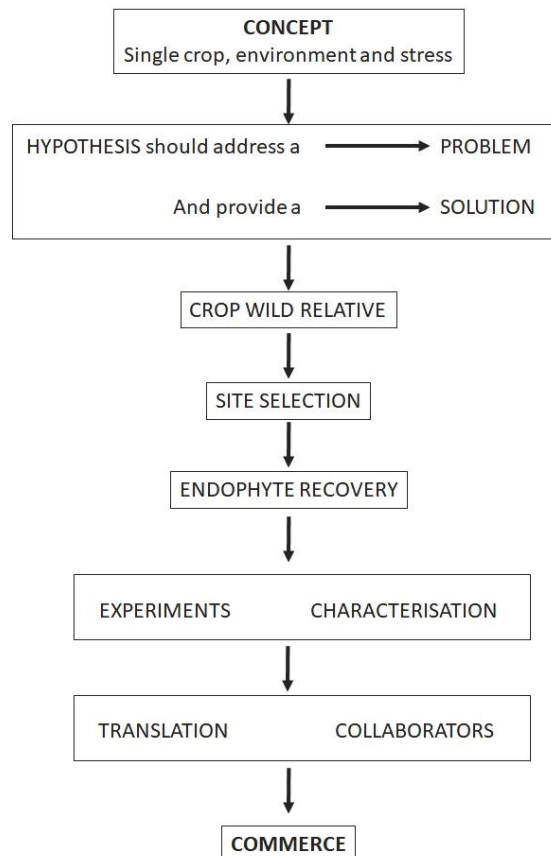


Figure 1. From Concept to Commerce: a schematic flowchart for the prospecting of endophytes for use in agriculture.

Acknowledgments: We thank: Goldcrop Limited, Cork, Ireland for the generous supply and dressing of barley seeds, and for advice on suitable cultivars to use; Crop Plot Trials and the Irish Department of Agriculture, Food and the Marine for carrying out the field trials; Helena Murphy for proof reading and the de-cluttering of technical terms; laboratory technicians at Trinity College Dublin and University College Dublin for providing supplies and technical support. Science Foundation Ireland funded the 2016 field trials through a Technology Innovation Development Award (Grant Number R15449). Enterprise Ireland are supporting further research and commercialisation through a Commercialisation Fund (Grant number CF 2017 0625-P). Trinity College Dublin provided financial support for the 2015 field trial through a PhD studentship grant.

Author Contributions: Brian R. Murphy, Fiona M. Doohan and Trevor R. Hodgkinson conceived and designed the experiments; Brian R. Murphy performed the experiments; Brian R. Murphy analyzed the data; Brian R. Murphy wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest.

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Opinion

The Emergence of Endophytic Microbes and Their Biological Promise

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Received: 20 April 2018; Accepted: 11 May 2018; Published: 16 May 2018

Abstract: As is true with animal species, plants also have an associated microflora including endophytes as well as microbes associated with the phyllosphere and rhizosphere (plant surfaces) and this is considered the plant microbiome. However, those organisms within virtually all tissues and organs of the plant are known as endophytes. Most often fungi are the most frequently recovered endophytes from plant tissues, but bacterial forms generally occur in greater numbers, but not in species varieties. The exact biological/biochemical role of the endophyte in the plant and how it interacts with the plant and other endophytes and plant associated organisms has not been intensely and carefully examined. However, this has not stopped investigators in exploring the direct utility of endophytes in boosting plant production, and discovering that endophytes can directly influence the plant to resist temperature extremes, drought, as well as the presence of disease causing organisms. Also, because of the relationships that endophytes seem to have with their host plants, they make a myriad of biologically active compounds some of which are classified as antibiotics, antioxidants, anticancer agents, volatile antimicrobial agents, immunosuppressive compounds, plant growth promoting agents, and insecticides. These endophytic compounds represent a wide range of organic molecules including terpenoids, peptides, carbohydrates, aromatics, hydrocarbons and others and it seems that these compounds may have a role in the host microbe relationship. Most recently and quite surprisingly, some endophytes have been discovered that make hydrocarbons of the types found in diesel and gasoline fuels. In addition, recently discovered are epigenetic factors relating to the biology and biochemistry of endophytes. Interestingly, only about 1–2% of the entire spectrum of 300,000 known plants have been studied for their endophyte composition. Additionally, only a few plants have ever been completely studied including all tissues for the microbes within them. Likewise, the vast majority of plants, including those in oceans and lower plant forms, have never been examined for their endophytes. Furthermore, endophytes representing the “microbiome” of world’s major food plants as they exist in their native “centers of origin” are largely unknown. This non-classical review is intended to provide background information on aspects of developments in endophyte biology and more importantly the identification of new questions in this field that need to be addressed. The review is primarily based on the author’s long held experience in this field.

Keywords: fungi; endophytes; natural products; plant microbiome; mycodiesel

1. Introduction

Endophyte biology is an emerging field. With many new developments in natural product characterization, along with their increasing use in biological control and the application of these organisms to sustain and assist crop production, the field has recently seen a huge jump in scientific attention and interest which has sprung from approaches and methods used in classical plant pathology [1]. These organisms are obtained from all types of plant tissues that seemed to bear no external evidence for the presence of any life forms within them. The advent of important and novel strides being made in the make-up and subsequently the importance of the human microbiome has

sparked a huge interest in the plant microbiome (endophytes) and how these organisms may influence the development and, the ability of a plant to resist disease, drought, heat, cold and other insults [2]. In addition, there has been renewed interest in endophytes since many of them have been shown to produce important compounds of pharmaceutical and commercial interest [3]. This list includes novel anticancer agents, antibiotics, novel immunosuppressive compounds, volatile antibiotic mixtures, antioxidants and most recently—fuel related hydrocarbons [3,4]. It appears that the field is ripe for scientific discovery and invention in a number of important scientific areas.

Since the early times, the definition of an endophyte has more or less remained the same—"a microorganism associated with living plant tissues that produces no apparent indication of its presence in the plant and seems not to cause harm to the host [5]." Endophytes have been isolated from virtually all plant organs (roots, stems, leaves, flowers, fruits, and seeds) and the most commonly observed endophytes are the fungi (usually Ascomyceteous fungi associated with Fungi Imperfecti), as well as bacteria and this includes the filamentous bacteria which are the Actinomycetes. Periodically, a Zygomycete, and a Basidiomycete and are also isolated. In the future, it may be the case that other life forms may be found that can be considered endophytes such as the mycoplasmas. There is some certainty that all plant forms, including those in the world's oceans, are hosts or are potential hosts for one or more endophytes [6]. Additionally, many of the lower plant forms such as mosses and liverworts are also hosts to endophytes [7]. This review will primarily focus on fungal endophytes of higher plant forms.

As with most biological sciences, the initial work on endophytes began as an observational science with investigators who were keenly interested in isolating, identifying every possible endophyte from a given plant and then moving on to the next plant species [5,8]. Much of this early work was being done by the Petrini's and their group in Switzerland [9]. However, in spite of all of the reports on endophytes from Europe, Canada and the USA, it appears that hundreds, if not thousands of novel endophytic species still remain to be discovered especially in those remote, unique and untouched forests, fields and even oceans of the world. Thus, recent efforts in India, Thailand, Brazil and China have resulted in a plethora of new information on fungal endophytes of tropical plant species. In addition, investigators, at the ecological level, have begun to study how endophytes may influence the biology of the plant enabling it to resist biological as well as environment stresses [2,10–12]. Collectively, these efforts have crystallized thinking about these organisms and how they represent the "plant microbiome." This switch in thinking from endophytes as strictly an academic subject to that of practical biology now has the potential for crop associated endophytes to be useful biological tools in promoting plant growth and assisting the plant to better cope with environmental and biological stresses.

On another note, studies on the distribution of endophytes quickly led to some attention be paid to the relationship of endophytes to their host plants. It has been learned that endophytes are mutualists and some, on occasion may become pathogenic when environmental and physiological conditions of the plant may allow [5,8]. However, the exact nature of the interaction with the plant and the products and processes involved in the interactions has gone missing from the literature. What has appeared however is conjecture on what might be happening in the microbe-plant interaction based on tangential observations and speculation.

The likelihood of finding taxonomic novel endophytic microbes is the greatest in the world's tropical and temperate rainforests and also with plants growing in peculiar and unexpected places such as harsh deserts, high alpine areas, or even in the world's oceans [3]. It is to be noted that novel endophytes probably do not commonly exist in botanical gardens in population dense areas because of the pollutants in the air that can and do affect microbial growth and activity. The same can be said of unusual plants artificially placed on university campuses where the same nasty pollution elements undoubtedly also exist and researchers can find easy access to plant material but take no thought of the microbes. Ultimately, with taxonomic novelty, it seems, the chances of finding chemical novelty greatly increases [3]. In addition, with the discovery of novel organisms possessing unique biochemical properties there are opportunities for the development of uses of such organisms.

This necessarily implies the extreme likelihood for patenting possibilities for new organism and new product development.

This review is certainly not intended to cover all aspects and all discoveries related to fungal endophytes, but it will cover the main recent focal points in this field from the perspective of work done in the author's own laboratory. Then, as part of the content of each focal point area the author has put forth his ideas and suggestions on what he sees as the potential for future studies in the given area and provides suggestions and guidance on how work on endophytes should be encouraged in some areas and discouraged in others. Specific examples that are given are used to illustrate important considerations when working with endophytes, their discovery, their biology, and their utility. Also discussed and referenced are certain critical techniques that apply to the science of endophyte biology and how they may be useful and applicable to those working in this field.

2. Endophytes, Isolation, Storage, Fundamental Biology

Many of the commonly isolated fungal genera associated with plants as pathogens can also be isolated as fungal endophytes and upon inoculation of the plant with the endophyte one does not witness the development of disease symptoms. Some of these fungal organisms may represent new endophytic species or biotypes. Some of the commonly fungal genera that we have commonly isolated include *Fusarium* sp., *Colletotrichum* sp., *Phoma* sp., *Pestalotiopsis* sp., *Xylaria* sp., *Cladosporium* sp., *Curvularia* sp., and a host of others. The most common represented group of fungi isolated as endophytes are the Fungi Imperfici whose perfect stage is commonly the Ascomycetes. Less frequently isolated members of the groups Basidiomycetes and Zygomycetes.

The methods of isolating these organisms are well known and usually involve the use of a surface treatment solution to rid the selected plant material of unwanted contaminants. This usually involves a thorough rinse in a 70% ethanolic solution followed by flaming the tissue or treatment with a Clorox solution. One must be careful as to not over treat the tissue to kill or harm the endophytes associated with epidermal tissues. Plant tissues are placed on water agar or other agars that may represent selective media containing certain antibiotics to discourage bacterial growth. Over the years we have realized that plant associated mites can cause major damage to endophyte based experimentation in the laboratory and can easily be handled by adding a small crystal or two taken from a common moth ball and placed into a plastic box containing Petri plates from which the endophytes will eventually be selected (Strobel—unpublished, 2018). Mite contamination of the laboratory can literally halt all scientific progress on endophytes until it is cleaned up. Once obtained in pure culture, we have learned that most tropical endophytes may be best preserved for future work by growing them on doubly autoclaved barley seed (hydrated), placed directly in small freezer vials and stored at -70°C [13].

In order to do taxonomic work on novel fungi it is important to have one or more spore stages of the organism available for description. Frequently endophytes show no sporulation on potato dextrose agar. While a choice of other agars may provide some help in encouraging sporulation, the best approach is to use gamma irradiated carnation leaves placed on water agar as a medium of choice [14]. In addition, as an alternative one may use pieces of the plant host material that has been autoclaved on a water medium that may encourage fructification of the organism. One could also even make a host based concoction agar medium from stem, leaf or root materials and eventually use more purified plant compounds that are critical for fructification [15]. Ideally, one can use the fungus growing and fructifying on these media as a source of biological material for scanning electron and light microscopy as aids in identifying the organism. Limited molecular techniques such sequencing targeting the ITS region are an important aid in identification and should be used to assist, but not used exclusively in trying to describe an organism. When no spore forms are evident under a multitude of growth conditions, one must rely on a host of other methods to assist in identification and description at the taxonomic level. This includes a complement of molecular gene sequence information starting at the ITS region and moving to specifically targeted gene sequence information involving certain enzymes, other analytical chemical data entries including an analysis of volatile chemicals, and scanning electron

microscopy of any unique hyphal/spore/fruiting body characteristics. Ultimately, any data gathered must be placed in context with data that exists in the mycological literature for other already known and previously described fungal species. It is to be noted that one must be prepared to face the taxonomic literature and scrutiny of the mycological community when working in the field of endophyte biology especially if one is intent on finding and thence describing novel microbes.

Microorganisms requiring plant tissues to complete their life cycle are classified as “obligate.” Well-documented examples of obligate endophytes are found among mycorrhizal fungi and members of the fungal genera *Balansia*, *Epichloë*, and *Neotyphodium*, from the family Clavicipitaceae (Ascomycota). Interestingly, there is little work demonstrating if there are other obligate endophytes, that are not specifically mycorrhizal fungi or members of the Clavicipitaceae. It is generally suspected that they do exist based on what is known and understood from obligate plant pathogens such as the rust and powdery mildew causing fungi and assuming that the number of host species is constant, the total number of host specific endophytic species can be extrapolated from the number of plant species [16]. Metagenomic analyses for the total population of endophytes in plants is a challenge since there is the technical limitation of separating fungal from host DNA. Basically, the plant DNA is much more abundant than fungal DNA which confounds the analysis. Additionally, we have observed that frequently slow growing unidentified endophytes do arise from many plant species. Upon the second transfer to a potato dextrose medium they again begin to grow very slowly and then gradually die as if some critically important nutrient was missing from the medium. Obviously, further work on such organisms is confounded by absence of the organism. We have seen hints of this phenomenon over the years in my laboratory but lack of available host plants (since many were obtained in the world’s jungles) to substantiate the initial observations made further work on the problem virtually impossible to study.

Also needed are studies showing the complete endophyte characterization of a given plant species. This should include mention of all organisms associated with all tissues and organs of the plant including flowers, fruits and roots which are sometimes difficult specimens to find and study. It is known that the make-up of the population of microbes in a plant will vary as a function of season, the organs being studied, and the host itself. It appears, however, that it is exceedingly interesting to find, identify, and describe totally novel fungal genera and species that are acting as endophytes. The biological potential of such studies is great given the fact, as previously mentioned, that novel taxonomy portends the presence of novel chemistry.

Specifically, I do not completely encourage studies describing the endophytes of a previously unstudied plant that show no novelty in the endophytic composition of the plant, any novel organism or other unique scientific contribution resulting from the study since so little new is to be learned. On the other hand, if a novel fungal genus is isolated from the plant then it is worthwhile to do the work that properly describes the new organism. In addition, if new uses and concepts are found with already known organisms this too can be valuable information. Furthermore, it must be recognized that the use of limited molecular genetic techniques to assist in the identification of the endophytic flora of a plant can be helpful but misleading if no attempts are made at the actual isolation and examination of the host endophytes. It is well understood that identification of fungi at the species level is really not possible with ITS sequence information alone. The use of classical mycological methodologies come in handy in helping with fungal species identification and this is not to be ignored.

3. Novel Endophytes

The opportunity to find novel endophytic fungi at the genus or species level can be done if one’s attention is properly directed. Many of the frequently appearing endophytic fungi have their counterpart relatives existing as plant pathogens. This is especially true of the genus *Fusarium* which is often isolated from numerous plant species as pathogens but it seems that they are even more common as endophytes. Most interesting however, are reports of truly novel fungi as endophytes that represent novel fungal genera. Such reports require all of the work necessary to list a valid new genus/species. This necessitates getting a pure culture, and then doing the comparative taxonomic work with other

related fungi at all levels including structural, developmental, and molecular as previously mentioned. Depositing the organism in a national, state or well-maintained university culture collection is also a requirement and one that is critically important. Novel endophytes can best be found in endemic plant species which are numerous in certain locations especially those hosting great biodiversity. Usually these places exist in temperate or tropical rainforests and maybe remote.

To illustrate the point of how to approach the discovery of novel endophytes I have shown below several examples of how we have solved the issue of biological discovery by visiting far reaching places in the world to find novel fungal genera. For instance, the fungus—*Seimatoantlerium tepuiense* was discovered as an endophyte of *Maguireothamnus speciosus* on top of the Roraima tepui on the border junction of Guyana, Brazil and Venezuela. The area is very remote and the host plant is endemic to this location and only two other small locations in tropical America. To get to this spot required the use of a rented helicopter and some knowledge of the local flora. The conidia of this endophyte resemble those of the commonly isolated endophyte—*Pestalotiopsis macrospora* however, the condiospore develops an appendage that is much more complex than the helicopter-like appendage in *P. macrospora* (Figure 1). In this case it forms an elk antler-like appendage and this structure easily dehisces from the spore (Figure 1) [17]. Presumably this allows for easy distribution by birds visiting the N.E. South American tepui plateaus since the spores can easily become attached to the fine structures of their feathers and eventually fall as the appendage becomes detached from the spore [17]. The fungus also makes small amounts of the anticancer drug-taxol. Thus, in this case not only was novel taxonomy discovered but so too was a product of extreme medical importance also found quite to our surprise.



Figure 1. Development of the conidiospore of *Seimatoantlerium tepuiense* a novel endophyte discovered on a tepui in the Brazil, Venezuela, Guyana area of South America. The actual spore is shown to the right as a scanning electron micrograph. The bar is equivalent to 20 microns.

One of the most important novel endophytes that we have ever isolated is *Muscador albus*. This whitish, non-spore producing endophyte was originally obtained from a cinnamon tree (*Cinnamom zeylanicum*) in Honduras [18]. This endophyte is relatively slow growing on potato dextrose agar and it makes a plethora of volatile organic compounds that possess strong antimicrobial activities [19]. In fact, the volatiles in split plate agar assays are not only inhibitory to other microbes but in most cases are lethal [19]. This makes for a likely prospect that the fungus itself might be useful as a biological control measure. In fact, in November of 2016, the US—Environmental Protection Agency approved the use of this organism for applications in US agriculture. Field testing by Marrone BioInnovations Co. in Davis, California, has shown efficacy of *M. albus* as a pre-plant soil additive to effectively eliminate root rot causing pathogens in fields of strawberry and celery. Numerous other applications of this fungus to control disease problems in various crops has been demonstrated by Mercier and his group [20,21]. The fungus has the potential to replace the use of methyl bromide as a

soil sterilant. It is the first broadly active antimicrobial producing endophyte to officially be on the market in 2017.

Since its discovery almost twenty years ago many other species of this genus have also been described and now number over twenty. All members of this group have several things in common—(1) they have close homology at the ITS r DNA level [22], (2) they each produce bioactive volatile substances and (3) they are all producing a sterile (non-spore) producing mycelium that is commonly convoluted, interwoven, and sometimes possess unique hyphal projections (Figure 2) [23]. One interesting example of a novel species of this fungus is *M. vitigenus*, which was obtained from *Paullinia paullinoides* in the Peruvian Amazon, and it makes only one volatile which is naphthalene, an insecticide [24]. Cultures of *M. vitigenus* were extremely active in repelling insects in Y-tube testing experiments [25].



Figure 2. The convoluted and interwoven hyphae of *Muscodor albus*, a volatile antibiotic producing fungus. The white bar is equivalent to 5 microns.

Another unique species is *Muscodor crispans* whose volatile composition is extremely antimicrobial, and the majority of its active compounds are on the FDA-GRAS list, that is Generally Recognized As Safe [26,27]. Thus, the bioactive volatiles can be either synthesized or made individually by fermentation processes using other microbes to make large quantities of the product in a liquid state (N. Gandhi unpublished, 2018). It has been learned that all of these volatile compounds in the fungal gas mixture are not needed for biological activity, but certain ones act in a synergistic manner to cause microbial inhibition and death of target microbes. Those that have such activity have been termed synergistans [28]. The products of *M. crispans* (Jeneil Biotech name of Flavorzon) at the 1% level has antimicrobial activity that parallels that of Clorox or quaternary ammonium [29]. This makes the solution usable on biological or industrial surfaces that may be contaminated or have the potential to be infected by or contaminated with organisms such as *Salmonella* sp. [29]. Since it is a safe product it has been used to decontaminate rooms where food-grade work is performed. Presently, another version of the Flavorzon formula is being tested for the prevention of decay in post-harvest fruits/vegetables. Still another version of the fungus gas formula is now being used to enhance the shelf life of certain products such as soy milk and is in the market place.

It turns out that each of these *Muscodor* spp. that were unique species, was patented by us at the time that they were discovered. Likewise, claims were made for the volatile chemistry that was discovered in each organism. In each case the US and many International patent offices approved the applications. It seems as if this intellectual property approach represents the best mechanism to make the discovery process useful to humankind. It is something that needs to be considered as one does all of the work to make important, valuable, and useful discoveries. Without intellectual property, a discovery such as the ones related to *Muscodor* are interesting but are rendered useless unless protection is sought and product development is pursued. Ultimately, the general public benefits from all of the efforts made in work of this type.

As a side light and most pleasing to me is the efforts of a group of Indian scientists who have recently described *M. strobilii* from *Cinnamomum zeylanicum*, the same species from which the original isolate of muscodor was obtained, only in this case the host tree was growing in southern India, which is most likely the same source area for the Honduran tree mentioned above. The organism was unique from other species in this genus by virtue of its production of very unique volatiles, its antimicrobial activity and its unique hyphal structure [30].

4. Bioactive Products of Endophytes and Their Utility

Most work on the secondary products of endophytes arose in the 1980s with a focus on the toxic compounds associated with diseases in livestock. These symptoms were associated with fescue grass that was infected with *Neotyphodium* sp. that produces toxic metabolites causing abortion and even death in animals grazing especially on this grass species [31]. Endophytes, during that era, received notoriety and the greatest interest in them was for the damage that they caused in livestock. The broader picture of their benefit to plants and mankind was not evident until later. Then with the advent of the discovery of *Taxomyces andreanae* in Montana in a yew tree, making the anticancer drug taxol, the concept of endophytes as a source of novel and useful drugs was introduced (Figure 3) [32,33]. At the outset, it seems only logical that such possibilities exist in nature if one assumes that the endophyte and the host plant are living in some sort of symbiotic relationship and horizontal gene exchange may occur between the symbionts. Thus, any contribution to the survival of a plant has the potential to be translated into products that also might be beneficial to mankind once isolation of the endophyte is achieved along with its biologically active products. As it turns out, taxol is a potent anti-oomyceteous compound and its mode of action on fungal pathogenic oomycetes is identical to its activity against rapidly dividing cancer cells [34]. It seems likely that taxol and its hundreds of related compounds are produced by *Taxus* sp. to protect the plant from attack by “water molds.”

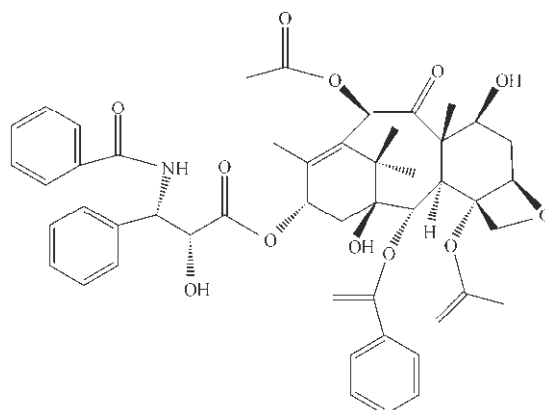


Figure 3. The structure of taxol.

In my experience, any search for a natural product (with a given bioactivity) is driven by what system of bioassays might be in place or available at the time of fermentation of the fungal culture and the purification procedures to which it may eventually be applied—bioassay guided fractionation. A good and effective bioassay can greatly assist in the eventual isolation and purification of a bioactive product by allowing an assessment of where on a TLC plate or a column eluent the product of interest may be located. To this end, over the past 20 years, a plethora of review papers have appeared on the secondary natural products of endophytes and their associated biological activities [35–39]. These reports nicely show that endophytes are a wonderfully unique and growing source of biologically active molecules and one must be prepared to work on virtually any class of biologically active molecules as one follows the bioactivity through the purification processes.

Thus, once an endophyte is isolated in pure culture some form of fermentation methodology is employed in order to find and characterize biologically active secondary products made by the organism. Normally, a more defined medium such as the M-I-D medium is used rather than more complex media [3]. The use of a more defined medium necessarily eliminates many medium products from the purification processes. Generally, we carry out fermentation in still culture in large flasks for a period of 2–3 weeks. Initial extraction is done with methylene chloride, or ethyl acetate or n—butanol. This step is followed by flash chromatography, then preparative thin layer chromatography and HPLC. After each step the various fractions are subjected to the biological assay that has been selected. The final and most desirable goal is to get a product that is crystalline so that the product may be subjected to X-ray crystallographic analysis. Ultimately, all other spectroscopic analytical data should be in agreement with the X-ray data.

A few of the products of endophytes on which we have worked are discussed and illustrated below. The first three products were all obtained from various endophytic isolates of *Pestalotiopsis microspora* that were found in various plants in various locations around the world. This fungus is one of the most frequently isolated endophytic fungi from tropical and temperate rainforests plants. It has been seen many times and assigned many different species names, usually solely based on the host plant from which it was isolated. Until recently little work has been done on the biological chemistry of this organism. It turns out to have a treasure trove of interesting novel and bio-active molecules. Each of the products, shown below, has a unique biological activity and the chemistry of the products is also unique. This nicely illustrates the point that one must be prepared to deal with a variety of techniques in separation science and chemical characterization when dealing with the bioactive products of endophytes.

4.1. Pesatin

This novel benzofuran was obtained from a culture of *Pestalotiopsis microspora*, an endophytic fungus obtained from *Taxus wallichiana* sampled in the Himalayan foothills at which time we were engaged in looking for other endophytes making taxol and were surprised in finding an endophyte making both taxol as well as two novel antioxidants. This fungal endophyte produces a new 1,3-dihydro-isobenzofuran which exhibits antioxidant activity eleven times greater than the vitamin E derivative troxol. The compound also has moderate antifungal activities [40] (Figure 4). Isolation of pestatin was achieved by extraction of culture fluid with methylene chloride followed by silica gel chromatography. Its structure was established by X-ray diffraction and ¹³C and ¹H NMR spectroscopies. The X-ray data demonstrated that pestatin occurs naturally as a racemic mixture. Mechanisms for antioxidant activity and post-biosynthetic racemization have been proposed. Isopestatin is also produced by this endophyte and it also possesses similar bioactivities as pestatin [41].

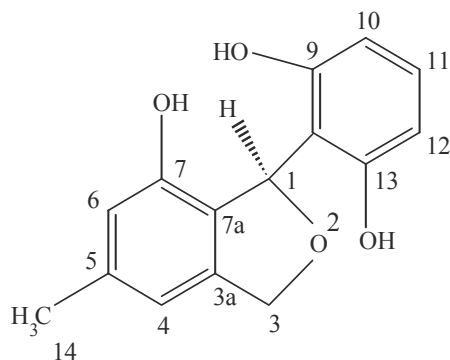


Figure 4. The structure of pestacin.

4.2. Ambuic Acid

We were greatly surprised to find a highly functionalized cyclohexenone being produced by an isolate of *Pestalotiopsis microspora* as an endophyte of *Fagraea bodenii* found in the highlands of Papua New Guinea [42] (Figure 5). The compound possesses weak antifungal properties. It was first natural product to have its absolute structure established by solid state NMR methods allowing a spatial assignment to the –OH group on carbon 7 [43,44] (Figure 5). Quite surprisingly, after the initial work on the isolation and structural determination of ambuic acid, it was later learned that it is one of the best compounds known for its anti-quorum sensing activity in Gram-positive bacteria [45]. Ambuic acid inhibits the biosynthesis of the cyclic peptide quormones of *Staphylococcus aureus* and *Listeria innocua*. Ambuic acid is a lead compound in the search for anti-pathogenic drugs that target quorum sensing—mediated virulence expression of Gram positive bacteria. Once again, what initially was supposed as a weak antifungal agent turned out to have a totally unsuspected biological activity as an inhibitor of quorum sensing in bacteria and this point was not established by us but by other investigators in Japan. An assay for anti-quorum sensing activity was not originally in our repertoire of bioassays.

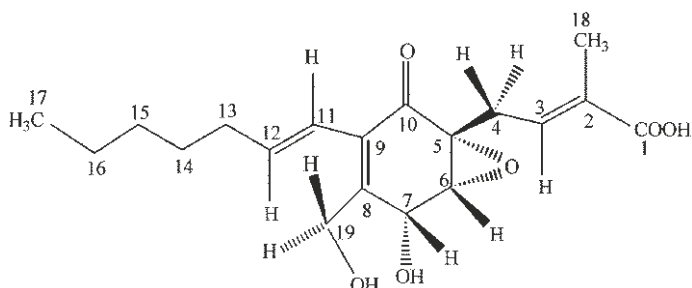


Figure 5. The structure of ambuic acid.

4.3. Torreyanic Acid

The Florida Torreya is a rare and threatened plant and for this reason it was sampled for the presence of unusual endophytes. Torreyanic acid was isolated and characterized as a dimeric quinone obtained from the endophyte, *Pestalotiopsis microspora* originally isolated from *Torreya taxifolia* in Northern Florida [46] (Figure 6). The compound was cytotoxic against 25 different human cancer cell lines with an average IC₅₀ value of 9.4 µg/mL, ranging from 3.5 (NEC) to 45 (A549) µg/mL. Torreyanic acid is 5–10 times more potent in cell lines sensitive to protein kinase C (PKC) agonists, 12-O-tetradecanoyl

phorbol-13-acetate (TPA), and was shown to cause cell death via apoptosis. Torreyanic acid also promoted G1 arrest of G0 synchronized cells at 1–5 µg/mL levels, depending on the cell line. It has been proposed that the eukaryotic translation initiation factor EIF-4a is a potential biochemical target for the natural compound. Additionally, it has been prepared by organic synthetic techniques.

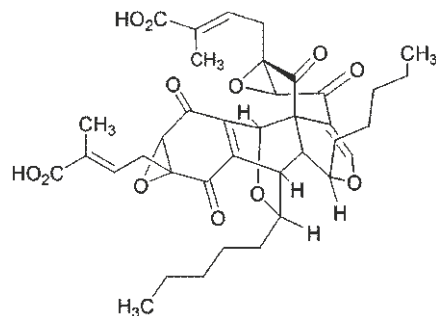


Figure 6. The structure of torreyanic acid.

4.4. Colutellin A

Colletotrichum dematium is an endophytic fungus recovered from a *Pteromischum* sp. growing in a tropical forest in Costa Rica [4]. Strangely enough this was the only endophyte isolated from samples of this plant and this is unusual circumstance especially in dealing with tropical plant species. This fungus makes a novel peptide antimycotic, Colutellin A, with minimum inhibitory concentrations of 3.6 µg/mL (48 h) against *Botrytis cinerea* and *Sclerotinia sclerotiorum*, respectively. This peptide has a mass of 1127.70 and contains residues of Ile, Val, Ser, N-methyl-Val, and β-amino-isobutyric acid in nominal molar ratios of 3:2:1:1:1, respectively. Independent lines of evidence suggest that the peptide is cyclic and sequences of val-ileu-ser-isoleu as well as ileu-pro-val have been deduced by MS/MS as well as Edman degradation methods. Colutellin A inhibited CD4—T cell activation of IL-2 production with an IC₅₀ of 167.3, whereas cyclosporine A, in the same test yielded a value of 61.8 nM. Since IL-2 production is inhibited by Colutellin A, at such a low concentration, this is an effective measure of the potential immunosuppressive activity of this compound. On the other hand, in repeated experiments, cyclosporin A at or above 8 µg/mL exhibited high levels of cytotoxicity on human peripheral blood mononuclear cells whereas, Colutellin A or DMSO alone, after 24 and 48 h of culture, exhibited no toxicity. Because of these properties Colutellin A has potential as a novel immunosuppressive drug [47].

4.5. Cryptocin

Tripterygium wilfordii is an Asiatic plant with strong immunosuppressive properties. It was targeted for study to learn if any of its endophytes might also make the same or related biologically active molecules. This rationale for this work follows that of the fungal taxol story. Instead, one of the endophytic fungi that was isolated was a unique species of *Cryptosporiopsis*. It was unique in that its large conidiospore was segmented quite unlike other members of this group [48]. This organism possessed unusual antifungal activities that were related to two novel compounds—cryptocandin (a novel lipopeptide) and cryptocin, an unusual tetramic acid. The acid was isolated and characterized by X-ray crystallography and other spectroscopic data (Figure 7) [48]. The compound was active against many plant pathogenic fungi (<1 µg per mL) and much less impressive activity against human pathogenic fungi (>50 µg per mL). However, to be noted is that the most sensitive fungus was *Pyricularia oryzae*, the causal agent of rice blast one of the most important pathogens in the world as it relates to food production.

The search for these valuable products has just begun. Usually, in my experience, there are several initial clues that make an organism a potential candidate for a producer of novel products. If it had been isolated from a unique environment or an endemic species, if it is a slow grower on common lab media and if it is coming from a totally unique fungal genus, the chances for finding something new are greatly improved. It is not, however, uncommon to work on an organism only to find that the bioactive product already been discovered in the same or related organism by someone else. Thus, novelty is a major goal! Repeated again, once the work is completed and a novel bioactive product has been discovered, one should make an effort to get patent coverage and find an interested commercial partner in order for some aspect of the general public to directly benefit from the discovery.

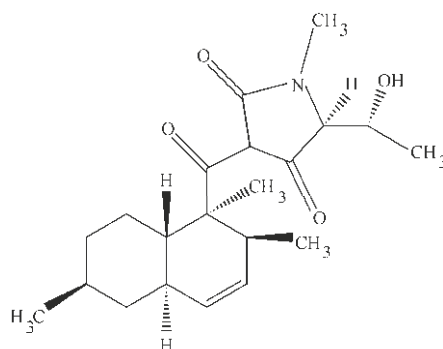


Figure 7. The structure of cryptocin.

5. Fungal Hydrocarbons and Fuels

We have relatively recently described endophytic fungi that make one or more hydrocarbons that have potential as fuels. Such products have been dubbed –mycodiesel [49]. This dramatically differs from standard yeast fermentation processes that utilizes sugars or starch to produce ethanol which is a proven and useful source of fuel, but by no means is it ideal. A number of endophytic fungi have been isolated and described that make compounds such as mono-terpenoids, alkanes, cyclohexanes, cyclopentanes, and alkyl alcohols/ketones, benzenes and polyaromatic hydrocarbons [4]. Many of these compounds are either identical to or are closely related to those specific classes of molecules that are found in diesel/gasoline [4,50]. Most importantly, these organisms make hydrocarbons while utilizing cellulose, hemicellulose and other polymers found in all plant-based agricultural wastes [51–53]. Endophytes are a prime source of hydrocarbon producers because they are the first microbes to begin the processes of tissue degradation, when a plant dies, to yield products with fuel-potential [4,53]. Examples of fungi that have been discovered that make fuel related hydrocarbons include—*Ascocoryne sarcoides*, *Gliocladium* sp., *Hypoxyylon* sp. (*Nodulisporium* sp.), *Annulohypoxylon* sp., *Phoma* sp., *Phomopsis* sp., and *Daldinia* sp. [13,50–55]. The volatiles produced by these organisms are usually possessing antibiotic activities which conceivably contribute to their role as symbionts. Finally, it seems possible that endophytic fungi may have an additional attribute of having contributed to the formation of crude oil in the first place and experiments have been done to demonstrate this phenomenon [53].

Muscodor albus, as described above, was used to discover other fungi making bioactive volatiles that turned out to be hydrocarbon-like molecules. Plant tissues of carefully selected plants were placed on half plates containing 10 day old cultures of *Muscodor albus* whose gases killed most fungi other than those also making bioactive volatiles. As an example of the use of this selection technique was the sole appearance of *Gliocladium roseum* (now classified as *Ascocoryne sarcoides*) [49,54] in Petri plates (PDA) supporting the growth of *Muscodor albus* showed that this fungus was able to survive and grow

in the presence of the inhibitory and lethal volatiles (VOCs) of *M. albus* [49]. Further testing of the recovered *A. sarcoïdes* culture revealed that its VOCs were active against other test fungi and that some of its volatile products had fuel potential [49]. Most interesting was the appearance of a series acetic acid esters of straight chained compounds including those of hexyl, heptyl, octyl, sec-octyl and decyl alcohols [49]. In addition, many other hydrocarbons were also noted in the GC/MS analyses of the VOCs of this fungus. The straight chained hydrocarbons (alcohols in the reduced form) are the backbone compounds found in all diesel fuels that we have investigated from widely differing parts of the world.

Since those observations were made it has become increasingly evident that many other endophytic fungi making volatile hydrocarbons are also resistant to the *M. albus* VOCs and their VOCs are biologically active [4]. Thus, this one selection technique has the potential of eliminating 80–90% of endophytic microbes that probably would not be of interest relative to VOC production since they would succumb to the VOCs of *M. albus* while growing out of the plant tissues on the Petri plates. This technique has merit if one does not wish to deal with a plethora of microbes that may not be of interest and waste time doing GC/MS analyses on organisms that would not be of interest. Nevertheless, it is to be noted that some important hydrocarbon producing fungi are sensitive to the VOCs of *M. albus* and would have been missed if this technique had been exclusively used for selection [55]. They can be usually selected on the basis of the biological activity of their VOCs.

An endophytic fungal strain of *Hypoxyylon* sp. was isolated from *Persea indica* an evergreen tree native to the Canary Islands where it grows not in abundance but is found on several islands including Tenerife in the Laurisilva [13]. This organism was isolated in its imperfect stage as *Nodulisporium* sp. from a small stem in the crown of the tree and it readily grows in the presence of the VOCs of *M. albus* which should facilitate its isolation from other plant sources. When grown on PDA—Petri plates, the VOCs produced by this fungus were primarily 1,8-cineole; 1-methyl-1,4-cyclohexadiene, and (+)-alpha.-methylene-.alpha.-fenchocamphorone (Figure 8). Not only these but many of the compounds made by this organism are of interest because of their high energy densities and thus the potential they might have as Mycodiesel fuels [13].

Six-day-old cultures of *Hypoxyylon* sp. (imperfect stage—*Nodulisporium* sp.) displayed maximal VOC-antimicrobial activity against *Botrytis cinerea*, *Phytophthora cinnamomi*, *Cercospora beticola*, and *Sclerotinia sclerotiorum* suggesting that the VOCs may play some role in the biology of the fungus and its survival in its host plant. In fact, this discovery has implications in developing methodology for strain improvement via mutation/selection techniques [13]. Media containing starch-or sugar related substrates best supported VOC production by the fungus. Continuous direct on-line quantification of VOCs was measured by proton transfer mass spectrometry (PTR-MS) covering a 12 day range with optimum VOC production occurring at 6 days at 145 ppmv with a rate of production of 7.65 ppmv/h [13]. The production of 1,8-cineole from a fungal source is of significant interest given the fact that this compound does not have any previously known biological sources aside from plant tissue, and has thus far limited the compound's availability for fuel purposes (Figure 8).

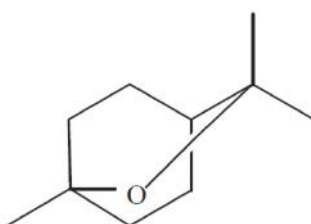


Figure 8. The structure of 1,8-cineole.

In a related *Hypoxylon* sp. a specific cineole synthase has been identified along with 11 new terpene synthase genes [56]. The discovery is an important first step in identifying a complete fungal pathway for the synthesis of 1,8-cineole and related monoterpenes. Currently, the yields of these terpenes and other hydrocarbons from fungal cultures does not allow for the production of these compounds at an economical level. These problems may eventually be overcome with conventional mutagenesis techniques and or genetic manipulation combined with complete sequencing and annotation of important fungal genomes such as *Ascocoryne sarcoides*, which is the best annotated endophyte thus far studied [57]. In combination with these approaches it will be necessary to develop non-conventional fermentation methods which may employ novel solid state methods allowing for a constant yield of volatile products via an airsteam and appropriate gas traps (Strobel unpublished, 2018).

Furthermore, as a practical matter cineole has been examined in multiple ways for its utility as a fuel. Studies on 1,8-cineole have shown prevention of phase separation when used as an additive in ethanol-gasoline fuel blends [58]. Furthermore, when fuels comprised of a gasoline/eucalyptus oil mixture (with 1,8-cineole as the major fuel component up to 80% or more) there was an improved octane number and reduced carbon monoxide exhaust [59]. Thus, 1,8-cineole is a worthy target molecule for study and scale up and it has amazing potential for replacing fossil-based hydrocarbons as a fuel additive.

Finally, it is to be noted that to do the best work on fungal volatiles does require the use of some unique methods and techniques in qualitative and quantitative analysis of these gaseous compounds. Standard practice is the use of the SPME fiber trapping technique combined with GC/MS to acquire an idea of the compounds being produced. The advent of PTR/MS has permitted the evaluation of on line-real time analysis and quantification of fungal gas production and now PTR/TOF-MS allows for this plus qualitative analysis of the gas stream [4]. Gas trapping in stainless steel columns as well as PTR /MS in line with a platinum catalysis will permit a total organic volatile compound production analysis—again in real time [60].

6. Control of Secondary Product Formation in Fungi

In many cases, once a product has been isolated and identified, the fungus may become attenuated in the production of the wanted compound (s). This is an event that has happened at least 50% of the time in my experience and can cause concern and self-doubt in the mind of the investigator. This is best exemplified by the attenuation of helminthosporide (a plant host specific toxin) produced by the plant pathogen *Helminthosporium sacchari*. When the organism was continuously cultured and transferred on a semisynthetic medium the production of this fungal toxin totally ceased [61]. However, when amounts of leaf exudate—concentrate were placed into the cultural medium, toxin production fully resumed. It turned out that several secondary products of sugarcane metabolism were responsible for the regulation of toxin production by this pathogenic fungus. One of these compounds was isolated and characterized as serinol (primary alcohol of serine) and the other was a sugarcane lipid-like compound [62]. It seems that the best place to look for answers to problems surrounding attenuation of secondary product formation in endophytic or plant pathogenic fungi is the host itself.

Similar events happened during the course of the discovery of taxol in an endophytic *Periconia* sp. which was isolated from *Torreya grandifolia* (a relative of yew that does not synthesize taxol) in China. This fungus, not previously known as a tree endophyte, was isolated from the inner bark of a small lower limb. When freshly isolated from the tree and placed in a semi-synthetic medium, the fungus produced readily detectable quantities of the anticancer drug taxol [63]. The production of taxol by *Periconia* sp. was demonstrated unequivocally via spectroscopic and immunological methods [63]. However, successive transfers of the fungus on a semi-synthetic medium resulted in the gradual attenuation of taxol production until minimal amounts of it were produced even though fungal growth was relatively unaffected. Several compounds, known previously as activators of microbial metabolism, including serinol, *p*-hydroxybenzoic acid, and a mixture of phenolic acids, were capable of fully or partially restoring taxol production to otherwise taxol-attenuated cultures. The compound

with the most impressive ability to activate taxol production was benzoic acid at 0.01 mM. To our knowledge, benzoic acid is not a direct taxol precursor [63].

The fungal fuel producing fungus—*Nodulisporium* sp. also falls into the category of an organism that will attenuate secondary product formation. It appears as if there are factors controlling 1,8-cineole and other hydrocarbon molecule production in this fungus based on the observation that serial transfer of the fungus on PDA resulted in a dramatic reduction of VOC production including 1,8-cineole [64]. Subsequently, when the attenuated organism was placed on certain plant parts and plant extracts the VOC production resumed to normal as measured by PTR-MS [64]. *Nodulisporium* sp. (Ti-13) in question had been isolated as an endophyte from *Cassia fistula* in Thailand. The fungus produces a spectrum of volatile organic compounds (VOCs) that includes ethanol, acetaldehyde, and 1,8-cineole as major components [64]. Initial observations of the fungal isolate suggested that reversible attenuation of the organism via removal from the host and at least 5 successive transfers in pure culture resulted in a 50% decrease in cineole production unrelated to an overall alteration in fungal growth. A compound (CPM₁) was obtained from *Betula pendula* (silver birch) that increases the production of 1,8-cineole by an attenuated Ti-13 strain to its original level as measured by a novel bioassay method employing a 1,8-cineole sensitive fungus—*Sclerotinia sclerotiorum* [64]. The host plant *Cassia fistula* also produces similar compounds possessing this activity. Bioactivity assays with structurally similar compounds such as ferulic acid, gallic acid and others suggested that the CPM₁ does not to act as a simple precursor to the biosynthesis of 1,8-cineole. Nuclear magnetic resonance spectroscopy and high-performance liquid chromatography electrospray ionization mass spectrometry indicated that the CPM₁ is a para-substituted benzene with alkyl and carboxyl substituents [64]. The VOCs of Ti-13, especially 1,8-cineole, have potential applications in the industrial, fuel, and medical fields.

These observations are suggestive of one of more regulatory mechanisms involved in hydrocarbon production that seems to be influenced by the host plant. Obviously, these considerations on the regulation of secondary product formation will be vital in making hydrocarbon—based fuel production from fungi a reality in the future. The work also has relevance to literally any plant microbe that is under the regulation of secondary product formation by host-related substances.

7. Endophytes and Epigenetics

We have recently learned that the regulation of the production of fuel –like compounds in fungi is under epigenetic control. *Hypoxylon* sp. (*Nodulisporium* sp.) is of interest because of its ability to make hydrocarbon-like compounds that may serve as fuels. Thus, experimental work was undertaken to affect the gene expression of this organism through compounds known to act as epigenetic modifiers and explore the possibility of the appearance of new products and/or the increased or decreased production of other VOCs already known from this organism [65]. For instance, it is realized that the complete analyses of fungal genomes in recent times has indicated that many putative biosynthetic gene clusters are located in the distal regions of the chromosomes and exist in a heterochromatin state with the constitutive genes often transcriptionally controlled by epigenetic regulation such as histone deacetylation and DNA methylation [66,67]. Several studies have demonstrated that the inhibition of histone deacetylase activity, through gene disruption or use of epigenetic modulators, leads to the transcriptional activation of gene clusters resulting in enhanced production of secondary metabolites [66,67]. Fungi treated with DNA methyl transferase- and histone deacetylase inhibitors exhibited natural product profiles with enhanced chemical diversity demonstrating that the small-molecule epigenetic modifiers are effective tools for rationally controlling the native expression of fungal biosynthetic pathways and generating biomolecules not previously associated with the organism [66,67]. Thus, the fungus was exposed to the epigenetic modulators—histone deacetylase- and a DNA methyltransferase inhibitor -SAHA (suberoylanilide hydroxamic acid) as well as 5-azacytidine (AZA), respectively. Subsequently, the organism displayed striking cultural changes including variations in pigmentation, growth rates and odor, in addition to significant differences in the bioactivities of its VOCs [65]. Analyses (GC/MS) of the VOCs produced by the

modulated fungus showed considerable variation with the emergence of several compounds not previously observed in this fungus, particularly an array of tentatively identified terpenes such as α -thujene, sabinene, δ -4-carene, γ -terpinene, α -terpinolene and β -selinene, in addition to several primary and secondary alkanes, alkenes, organic acids and derivatives of benzene. The spectrum of identifiable compounds, in this study, was greatly enhanced by virtue of the organism being altered with the epigenetic modulators [65]. It is likely that the same phenomenon would occur with other fungi but the result would be an entirely different set of secondary products.

8. Biology of Endophytes and Their Ultimate Utility

This review has considered only a limited range of topics concerning endophytes. Their vastness and diversity literally covers the entire planet since they are associated with all macroscopic plant life forms. Only a small portion of their range and diversity has been sampled with entire vast reaches of the world never having been sampled at all. This would include areas of central Africa, the lower reaches of the Amazon, many area of the rainforests of South Eastern Asia, as well as the expansive forests of Russia and central Asia. In addition to these sources are the macro algae of the world's oceans along with an abundance of other plant forms such as the ferns, mosses, lichens that also need attention. Only about 1% of the world's botanical sources of endophytes have ever sampled and incompletely at that since only a limb or leaf samples were the subject of study. It is well known that less accessible roots, fruits and the inner tissues of the stem each contains its own endophytic assemblage. The patterns of abundance and diversity of these organisms then becomes an issue for study as per Arnold et al. [68]. The biology of endophytes is also influenced by leaf or plant age, canopy cover, seasonal factors and even the presence of other endophytes [69].

Most interesting are observations that suggest that endophytes may play some role in the protection of the host plant from plant pathogens or other agents of destruction including insects and herbivores [11]. This brings into focus the potential roles of endophytes in enabling the host plant to be best fit for its survival and the potential that exists for harnessing these organisms for the benefit of agriculture [11,70]. Thus, managing the plant microbiome is a goal of several companies that wish to take advantage of the role that endophytes may play in the health, fitness, growth and development of the plant in its environment. Some of these are Indigo (Cambridge, MA, USA), Bayer/Ginko Bioworks (West Sacramento, CA, USA) and MBI (Davis, CA, USA). Many in the world of agricultural business would agree that managing a living organism as a product can be fraught with some problems. These are usually lack of reliability and consistency of outcomes when placing the live product on the market.

Some guidance for the development, and testing endophytes for eventual commercialization has recently been presented [71]. This work was done because of the problems mentioned above. Murphy et al. [71] began their work on barley by finding candidate source endophytes in a wild relative of *Hordeum mirnum* followed by testing in a carefully controlled environment and finally doing multi-year field testing. In their hands this approach was successful. It is to be strongly stressed in that what is generally missing in most studies of endophytes of crop plants is exactly what was avoided by Murphy et al. [71]. The organisms of interest to them were recovered by an extensive search and description of the endophytic flora of wild host plants growing in or near the center of origin of that species. It is entirely likely that that most plants serving mankind as food sources have long since lost their complement of normal flora as a result of movement from their centers of origin, intensive genetic selection techniques dealing only with the seeds or fruits of the plant and exposure of the crop to a myriad of pesticides. Evidence for this concept has been demonstrated in wild wheat related species vs. *Triticum aestivum* [72]. The wild types relatives of wheat contain a plethora of taxonomically diverse fungal endophytes that are not found in modern wheat. It turns out that most wild or native food source plants have yet to have their microflora studied. It behooves us to find and carefully preserve all natural sites that harbor wild relatives of mankind's major and minor food plants not only for the genetic diversity that they may hold but also for the unique microflora of these plants that may exist in such places.

9. Endophyte Biology as a Platform for Teaching Undergraduate Students

Finally, it should be mentioned that the concepts and techniques involved in isolating and characterizing endophytes and studying their biology represents an excellent opportunity for independent undergraduate participation. Several studies have demonstrated that an independent research experience is the best way to get undergraduates excited about a career in science [73]. In fact, the vast majority (around 80%) of the science faculties in US universities had an undergraduate research experience during their college careers. In this case a student can do her/his own collecting, plant identification, finding GPS coordinates and bringing the samples to the lab for further work. Endophyte isolation, microbial identification using classical methods and modern molecular methods offer a student a real opportunity to acquire independent project ownership which is a great way to encourage progress on a project. Ultimately, the student will have the possibility to develop novel bioassays for endophyte products or learn a myriad of other techniques involving chemistry, molecular biology, statistical methods and spectroscopic methods among others as applied to the microbes that were isolated. The student may have an opportunity to publish her/his work and possibly be involved in the patenting process. In fact, in this review I have cited many papers from my lab as the reference list to this paper. The names of Mitchell, Daisy, Tomsheck, Schaible, Mends and Nigg had all worked as undergraduate students in my laboratory and the majority of these students were women. Each of these students has had one or more senior authored papers from one or more individual research projects and their names appear in the Reference section of this paper [13,24–28,55]. The concept of giving undergraduate students an opportunity to work on endophyte projects has been as rewarding for them as it has for me.

Acknowledgments: This research received no specific grant.

Conflicts of Interest: The author declares no conflict of interest.

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ISBN 978-3-03897-247-1