BELOWGROUND FUNGAL COMMUNITY COMPOSITION AND DISTRIBUTION IN PEAT FORESTS AND A MANAGED PEAT ECOSYSTEM OF MALAYSIAN TROPICAL PEATLAND

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ABSTRACT

Fungi have a primary role in the decomposition of organic matter in tropical peatlands and contribute to maintaining ecosystem functions. Understanding the belowground fungal diversity and composition is a preliminary step to predicting the stability of ecosystem functions, especially in ecosystems that have been exposed to anthropogenic impacts. This study determined soil fungal diversity and community composition in two types of forest: Peat swamp forest (PSF) and logged-over secondary forest (LOF) and a managed ecosystem (oil palm plantation, OPP) in peatland in Sarawak, Malaysia. Fungi from peat samples were isolated and identified using 185 rDNA region sequencing. Fungal diversity was calculated, and Bray-Curtis dissimilarity was performed to compare fungal communities between the study sites. LOF has the highest diversity, followed by PSF and OPP. In addition, the fungal community composition is more similar between LOF and PSF than between the peat forests and OPP. Aspergillus spp. and Trichoderma spp. contributed to the great dissimilarities between the peat forests and the managed ecosystem due to their dominance. In conclusion, LOF is an important ecosystem that retains relatively high fungal diversity. Different fungal communities were observed, which contained ecologically important fungal groups incorporated into the peat forests and an oil palm plantation.

Keywords: fungal diversity, logged-over forest, primary forest, tropical peatland

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INTRODUCTION

A growing interest in belowground biodiversity in peatlands has been observed over the last two decades, owing to their functional importance in regulating nutrient cycling, decomposition, primary production, and climate change (Bach *et al.*, 2020; Song *et al.*, 2020). Belowground biodiversity, comprising primarily of microorganisms (bacteria, archaea, and fungi) and larger organisms (earthworms, ants, termites, and arachnids), play important ecological roles as symbionts, mutualists, decomposers, and pathogens (Bardgett

and Van Der Putten, 2014; Food and Agriculture Organisation, 2020; Tedersoo et al., 2014). Soil microorganisms, particularly bacteria, archaea, and fungi, are key indicators of various soil quality and health attributes, such as nutrient availability, carbon sequestration, incidence of plant diseases, and promotion of plant growth (Fierer et al., 2021). Their roles are crucial in terrestrial ecosystems, particularly in agricultural ecosystems, to address challenges and solutions for sustainable farming, food security, and climatic impacts. Understanding the community, structure, and diversity of soil microorganisms is critical for predicting ecosystem functioning in response to environmental changes so as to allow the implementation of conservation and management efforts (Egidi et al., 2019) and maintain ecosystem stability during climate change (Li et al., 2019).

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In peatlands, soil fungi constitute a large fraction of the belowground community (Juan-Ovejero et al., 2020). Most of the dominant soil fungi in peatlands are known as saprotrophs, which play a significant role in the decomposition of organic matter (Thormann, 2006; Thormann and Rice, 2007). Fungi produce a variety of extracellular enzymatic substances to assist in the degradation of simple and complex recalcitrant polymers, such as starch, cellulose, proteins, chitin, keratin, and lignin of wood (Eichlerová et al., 2015; Paredes et al., 2014). In addition, other fungi classified in the fungal guilds of pathogens, yeasts, ectomycorrhizal fungi, ericoid mycorrhizal, and dark septate endophytic fungi also contribute to the decomposition of organic matter (Andersen et al., 2013; Juan-Ovejero et al., 2020; Robinson et al., 2020). In a recent commentary, symbiotic fungi are considered ecosystem engineers due to their role in regulating the carbon sink in ombrotrophic peatlands via nutrient transfer (Barel and Robroek, 2023). Therefore, soil fungal communities and their diversity are recognised as the main components in regulating carbon dynamics and nutrient turnover in the peatland ecosystem (Juan-Ovejero et al.,

Soil fungal communities have been extensively identified and studied in many ecosystems across the globe (Egidi et al., 2019; Newsham et al., 2016; Shi et al., 2014) including in peatlands. However, studies are limited to bogs (Filippova and Thormann, 2014; Golovchenko et al., 2013; Morrison et al., 2020) and fens (Asemaninejad et al., 2017; Mpamah et al., 2017; Zhang et al., 2017), while tropical peatlands remain overlooked (Juan-Ovejero et al., 2020). As tropical peatlands are one of the largest carbon stock reservoirs (Page et al., 2022), it is important to protect and restore them by studying the diversity and functional role of their inhabiting soil fungi. This is because the carbon balance in peatland ecosystems is largely carried out by microbial activity, besides the interactions between plants and diverse soil communities (Barel and Robroek, 2023; Page et al., 2022). In Malaysia, tropical peatlands are exposed to anthropogenic disturbances, such as agricultural conversion, which led researchers to conduct studies on soil fungal communities. A few studies have emphasised the sensitivity of soil fungal communities towards the conversion of forest to oil palm cultivation (Kerfahi et al., 2014; Mcguire et al., 2015; Robinson et al., 2020). These studies showed that soil fungal communities can be an indicator or basis to monitor the impact of land use and response to environmental changes, especially in agricultural ecosystems. Thus, it is imperative to deepen the understanding of microorganisms in these ecosystems. Considering the importance of soil fungi and the limited studies in tropical

peatlands, this study was conducted to determine soil fungal communities and diversity in the tropical peatlands of Sarawak, Malaysia. The objective of this study is to determine soil fungal community, diversity, and structure under two types of forests, *i.e.*, peat swamp forest (PSF) and logged-over secondary forest (LOF), and a managed ecosystem in an oil palm plantation (OPP).

MATERIALS AND METHODS

Site Description and Sample Collection

Study sites were selected in the district of Sri Aman, Sarawak, Malaysia, representing the area of lowland tropical peatland. Peat samples were collected in August 2013 during the dry season (precipitation <200 mm/ sampling month) from peat swamp forest (PSF), loggedover secondary forest (LOF) and a managed peat ecosystem represented by an 8 year-old oil palm plantation (OPP). PSF is a gazetted mixed tropical peat swamp forest, owned by the Malaysian government, with an extensive area of 43 147 ha (Melling, 2016) and located along a riverbank of the Maludam River. PSF is a habitat for several important high-value timber species, such as Gonystylus bacanus, Dactylocladus stenostachys and Shorea spp. This forest contains diverse faunal species, with a few endemic fauna such as Presbytis chrysomelas crucige (Red Banded Langur) and Nasalis larvatus (Proboscis monkey) (Melling, 2016). LOF is a secondary forest that was recovered after timber extraction for land preparation years ago. It was covered with shrubs and small woody trees (Kusai et al., 2018). In OPP, routine management practices have been carried out throughout the year, such as the application of fertiliser and herbicides, tilling, compaction, and maintaining the water table for palm growth and production. The reduction of the water table within the oil palm plantation was maintained by a drainage system on site (Dhandapani et al., 2020). The total area of LOF was 2411.5 ha, while the planted OPP area was 4152.7 ha.

Peat samples were collected at ten sampling points at each study site. In PSF and OPP, the samples were collected in two transects with approximately a distance of 10 m to 100 m between each sampling point, while samples from LOF were taken in a line transect at a 100 m interval (Maidin *et al.*, 2016; Siti Ramlah *et al.*, 2016). The coordinates for each sampling point were recorded (Kusai *et al.*, 2018).

Peat soil cores were taken from peat surfaces up to 60 cm using a peat auger (Eijkelkamp, The Netherlands) with a half-cylindrical shape (diameter: 5.2 cm). A core sample collected in the peat auger was divided into four subsamples according to depths of 0-30 cm, 30-40 cm, 40-50 cm, and 50-60 cm and placed into designated 50 mL tubes before being transported to the laboratory for analysis. All peat samples were stored at 4°C for a short-term period of one month, and biological analyses such as the isolation of fungi were conducted within two weeks of sampling.

Isolation and Identification of Fungi

Peat samples were serially diluted up to a dilution of 10^{-5} for fungal isolation. A volume of 100 μL of dilutions 10^{-4} and 10^{-5} was spread evenly on potato dextrose agar (PDA) and malt extract agar (MEA) and incubated at $25 \pm 2^{\circ} \text{C}$ for three days. Colonies grown on both media were subcultured onto a new PDA to obtain pure cultures and used for identification.

The pure cultures were grown on PDA and incubated at $25 \pm 2^{\circ}\text{C}$ for three and seven days for fast-growing fungi and slow-growing fungi, respectively. The mycelium of the isolate was harvested, and genomic DNA was extracted using Fungi/ Yeast Genomic DNA Isolation Kit (NorgenBiotek Corp., Canada) according to the manufacturer's instructions.

A pair of primer; forward primer, EF4-F (5'-GG AAGGG[G/A]TGTATTTATTAG-3') and reverse primer fung5-R (5'-GTAAAAGTCCTGGTTCC CC-3') were employed to amplify the 18S rDNA region of fungi (Smit et al., 1999). A volume of 25 μL PCR master mix was used; 1× PCR buffer, 1.0 μM of each primer, 0.2 mM of each dNTP, 0.3 U of GoTaq Flexi DNA polymerase (Promega, Madison, WI, USA), 2.5 mM MgCl₂, 0.3% of bovine serum albumin and 1.0 µL DNA template. Amplification of 18S rDNA was carried out in a thermal cycler (Applied Biosystems Veriti™ Thermal Cycler, USA) with the following cycle: Initial denaturation at 94°C for 3 min, 30 cycles of denaturation, annealing and extension at 94°C for 1 min, 48°C for 1 min, 72°C for 1 min, respectively. The final extension was performed at 72°C for 5 min.

The PCR product was electrophoresed in 1% agarose gel, stained with FloroSafe DNA Stain (First BASE Laboratories, Malaysia) and run at 80 V for 30 min. The gel was viewed under UV to observe the amplified band of interest with a size of 550 bp. The amplified bands were excised and purified using a purification kit, EasyPure® Quick Gel Extraction Kit (Transgen Biotech, China). The purified products were sent to First BASE Laboratories (Malaysia) for sequencing according to the Sanger sequencing-based method. Sequences were compared with the nucleotide sequences in GenBank using Basic Local Alignment Search Tool (BLAST) analysis (http://blast.ncbi.nlm.nih.gov/blast/Blast.cgi) to verify the similarity of

the sequences and classify the fungal isolates into genus and species levels.

Diversity Indices and Data Analysis

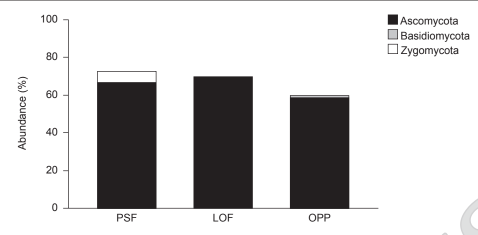
Species diversity in the study sites was calculated using the Shannon's diversity index (H') (Shannon and Weaver, 1963) and the Simpson diversity index (D) (Simpsons, 1949). Species richness (S) was determined using the total number of species in each study (Magurran, 2004; Pielou, 1969). Evenness (E) was also calculated to determine the distribution of abundance across the species in a community (Hill et al., 2003). Analysis of similarities (ANOSIM) was run to determine the significant variation in the fungal composition between PSF, LOF and OPP. Prior to analysis, the square root transformation was applied to the fungal composition data. The comparison of fungal composition between study sites was evaluated using pairwise Bray-Curtis distance and displayed using non-metric multidimensional scaling (NMDS). Similarity percentage (SIMPER) was also determined to identify the dominant species contributing to the fungal composition in the study sites. All the analyses were performed using PRIMER version 7 software (PRIMER-E Ltd., Plymouth, United Kingdom).

RESULTS

$Below ground \, Fungal \, Community \, and \, Composition$

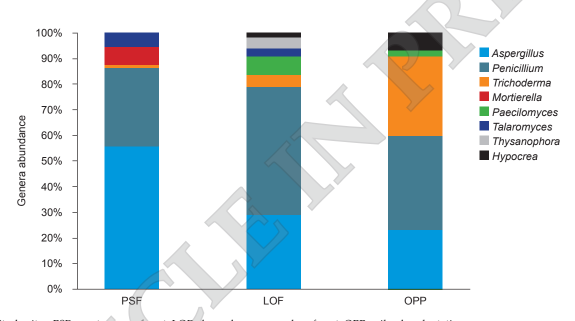
A total of 203 isolates were isolated and identified from all study sites, representing 3 phyla, 16 genera, and 30 species. The percentage of similarity between the isolated fungi and the sequences in the GenBank was high, between 81% and 100%. In total, 73 isolates were obtained from PSF, while 70 and 60 isolates were obtained from LOF and OPP, respectively. Three phyla comprising Ascomycota, Basidiomycota, and Zygomycota were identified using the 18S rDNA region, with Ascomycota being the most frequently isolated from the peat samples (12 genera) (Figure 1). Meanwhile, only Ascomycota was found in the LOF. A small proportion of Basidiomycota was obtained from the OPP, whereas Zygomycota was only found in PSF.

At the genus level, *Aspergillus*, *Penicillium* and *Trichoderma* were commonly found in all study sites (*Figure* 2). About 54.8% of the fungi found in the PSF were *Aspergillus*, and half of this number was from the LOF and OPP. *Penicillium* was highly isolated from the LOF at 47.1%, compared to PSF (30.2%) and OPP (34.9%). Meanwhile, *Trichoderma* was distinctly found in OPP (30%), compared to PSF (1.4%) and LOF (4.3%).



Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation.

Figure 1. Abundances of soil fungal taxa.



 $Note: Study\ sites: PSF-peat\ swamp\ forest; LOF-logged-over\ secondary\ forest; OPP-oil\ palm\ plantation.$

Figure 2. Abundance of fungal genera in all study sites. Genera less than 2% were excluded. Genera excluded were Cordyceps, Coniochaetales, Lecythophora, Meyerozyma, Myrothecium, Polyporus, Purpureocillium and Umbelopsis.

The common species present in all study sites were Aspergillus fumigatus, Penicillium corylophilum, P. roqueforti, Penicillium sp., and Trichoderma sp. (Table 1). The teleomorph of Trichoderma, i.e., Hypocrea, was found in the LOF (1.4%) and OPP (6.7%). A species of Basidiomycota, Polyporus ciliatus, was isolated at a low frequency (1.7%) in OPP. Meanwhile, Mortierella chlamydospora (6.8%) and Umbelopsis isabellina (1.4%), which are classified in the phylum Zygomycota, were only found in the PSF.

Five species unique to PSF, A. fischeri, A. flavipes, M. chlamydospora, P. purpurogenum and U. isabellina, were absent in the LOF and OPP. Likewise, five species from the OPP, i.e., Coniochaetales sp., H. rufa, Lecythophora sp., P. freii, and Polyporus ciliates,

were not found at other study sites. In contrast, nine species were found only in the LOF and not detected in the PSF and OPP. The species were Cordyceps cylindrica, Meyerozyma guilliermondii, Myrothecium sp., P. chrysogenum, P. commune, P. oxalicum, Purpureocillium lilacinum, Thysanophora sp., and Trichoderma harzianum.

Fungal Diversity

Based on Shannon's and Simpson's diversity indices, LOF displayed a higher fungal diversity, at 2.321 and 0.864 ($Table\ 2$), than the other sites. This was followed by PSF (H' = 2.166, D = 0.852) and OPP (H' = 2.049, D = 0.832). Species richness was also highest in the LOF, with 18 species. However,

PSF and OPP recorded the same richness, with 14 species. Species evenness ranged between 0.776 and 0.821. The least even distribution was observed in OPP (0.776), while the highest evenness was recorded in PSF (0.821).

The fungal community composition differed significantly between PSF, LOF, and OPP (ANOSIM, R global = 0.336; number of permutations = 999; P < 0.001; *Table 3*). The fungal community in PSF and OPP showed a significant dissimilarity (R = 0.452, P < 0.001). The ordination plot showed differences in species composition between PSF and LOF, where the composition resembled each other more than OPP (*Figure 3*).

The SIMPER analysis revealed that three species contributed to more than 80% of the total species identified in PSF (*Table 4*). Aspergillus

fumigatus contributed about 50.85% of the fungal assemblages in PSF, while *Penicillium* sp. (37.24%) and *Trichoderma* sp. (55.83%) contributed greatly to fungal assemblages within the LOF and OPP.

SIMPER analysis for dissimilarity indicated greater dissimilarity between PSF and OPP (average dissimilarity = 74.87%) than LOF and OPP (average dissimilarity = 73.24%) and between PSF and LOF (average dissimilarity = 60.87%, Table 5). The species that contributed to the largest differences were *Trichoderma* sp. and *A. fumigatus*, abundant in OPP and PSF. The taxa that contributed to the dissimilarity between LOF and OPP were *Trichoderma* sp. and *Penicillium* sp. The fungal community in PSF was more similar to LOF, with *Aspergillus* sp., *Penicillium* sp., and *P. roqueforti* as the main contributors.

TABLE 1. PERCENTAGE ABUNDANCE OF FUNGI ISOLATED IN DIFFERENT STUDY SITES

	Smarias	P	Percentage abundance (%)		
	Species	PSF	LOF	OPP	
	Aspergillus fischeri	1.4	0	0	
	Aspergillus flavipes	2.7	0	0	
	Aspergillus fumigatus	31.5	25.7	16.7	
	Aspergillus niger	1.4	1.4	0	
	Aspergillus nomius	5.5	0	1.7	
	Aspergillus sp.	12.3	0	3.3	
	Coniochaetales sp.	0	0	1.7	
	Cordyceps cylindrica	0	1.4	0	
	Hypocrea atroviridis	0	1.4	5.0	
	Hypocrea rufa	0	0	1.7	
	Lecythophora sp.	0	0	1.7	
	Meyerozyma guilliermondii	0	1.4	0	
	Mortierella chlamydospora	6.8	0	0	
	Myrothecium sp.	0	1.4	0	
	Paecilomyces sp.	0	7.1	1.7	
	Penicillium chrysogenum	0	1.4	0	
	Penicillium commune	0	5.7	0	
	Penicillium corylophilum	5.5	2.9	3.3	
	Penicillium freii	0	0	3.3	
	Penicillium oxalicum	0	1.4	0	
	Penicillium purpurogenum	1.4	0	0	
	Penicillium roqueforti	8.2	11.4	23.3	
	Penicillium sp.	15.1	24.3	5.0	
7	Polyporus ciliatus	0	0	1.7	
*	Purpureocillium lilacinum	0	1.4	0	
	Talaromyces purpureogenus	5.5	2.9	0	
	Thysanophora sp.	0	4.3	0	
	Trichoderma harzianum	0	1.4	0	
	Trichoderma sp.	1.4	2.9	30.0	
	Umbelopsis isabellina	1.4	0	0	

Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation.

TABLE 2. DIVERSITY INDICES OF SOIL FUNGAL COMMUNITIES

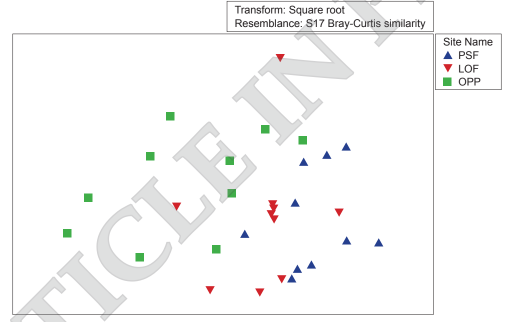
Sites	Total isolates	Shannon's diversity index	Simpson's diversity index	Species richness	Evenness
PSF	73	2.166	0.852	14	0.821
LOF	70	2.321	0.864	18	0.803
OPP	60	2.049	0.832	14	0.776

Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation.

TABLE 3. ANALYSIS OF SIMILARITIES (ANOSIM) IN SPECIES COMPOSITION OF FUNGI

Sites	R value ^a	P-value
PSF vs. LOF	0.2	0.003
PSF vs. OPP	0.452	0.001*
LOF vs. OPP	0.372	0.001*

Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation; vs. - versus; ^a ANOSIM; R global = 0.336; *Number of permutations = 999; P < 0.001.



Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation.

Figure 3. Non-metric multidimensional scaling (NMDS) ordination comparing the fungal community in different study sites.

TABLE 4. SIMPER ANALYSIS OF THE SIMILARITY OF SPECIES COMPOSITION

Sites	Average similarity	Species	Average abundance	Contribution (%)	Cumulative contribution (%)
		Aspergillus fumigatus	1.49	50.85	50.85
PSF	47.90	Aspergillus sp.	0.78	16.49	67.34
		Penicillium sp.	0.80	15.91	83.24
LOF	44.99	Penicillium sp.	1.21	37.42	37.42
		Aspergillus fumigatus	1.19	34.30	71.72
OPP	34.13	Trichoderma sp.	1.16	55.83	55.83
		Penicillium roqueforti	0.87	23.24	79.07

Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation.

TABLE 5. SIMPER ANALYSIS OF DISSIMILARITY OF FUNGAL COMMUNITIES

Study site	Average dissimilarity (%)	Species dissimilarity	Average abundance	Contribution (%)
DCF LOF	60.87	Aspergillus sp.	0.78	11.30
		Penicillium sp.	0.80	10.63
		Penicillium roqueforti	0.48	10.02
		Aspergillus fumigatus	1.49	7.97
		Mortierella chlamydospora	0.44	6.80
PSF vs. LOF		Penicillium corylophilum	0.34	5.98
		Talaromyces purpureogenus	0.34	5.63
		Paecilomyces sp.	0.00	5.47
		Penicillium commune	0.00	5.06
		Aspergillus nomius	0.34	4.72
	74.87	Trichoderma sp.	0.10	15.44
		Aspergillus fumigatus	1.49	13.37
		Penicillium roqueforti	0.48	11.17
PSF vs. OPP		Penicillium sp.	0.80	11.11
		Aspergillus sp.	0.78	9.51
		Mortierella chlamydospora	0.44	6.25
		Penicillium corylophilum	0.34	5.44
	73.24	Trichoderma sp.	0.20	15.54
		Penicillium sp.	1.21	15.16
		Aspergillus fumigatus	1.19	12.87
OF OPP		Penicillium roqueforti	0.74	10.94
LOF vs. OPP		Paecilomyces sp.	0.38	5.70
		Penicillium commune	0.34	4.81
		Penicillium corylophilum	0.20	4.46
		Hypocrea atroviridis	0.10	3.87

Note: Study sites: PSF - peat swamp forest; LOF - logged-over secondary forest; OPP - oil palm plantation. vs.- versus.

DISCUSSION

Fungi obtained in this study belonged to the Ascomycota, Basidiomycota, and Zygomycota taxa, commonly present across peatlands, such as in China (Zhang et al., 2017), Japan (Yabuki et al., 2014), Canada (Asemaninejad et al., 2017; Zhang et al., 2023), Europe (Elliott et al., 2015; Sun et al., 2016, Vinogradova et al., 2023), Malaysia (Ayob et al., 2018; Kusai et al., 2018), Brunei (Tripathi et al., 2016a), and Australia (Birnbaum et al., 2023). These taxa can degrade organic matter, including cellulose, hemicellulose, pectin, lignin, tannin, and polyphenolic compounds (Thormann, 2006). A large proportion of Ascomycota indicated that this taxon is predominant in the peatlands of Sarawak, either in natural or managed ecosystems, due to their physiological characteristics as heavy sporulators and fast-growing microorganisms (Thormann and Rice, 2007). Moreover, ascomycetes have high adaptability to environmental stresses by utilising resources, leading to more generalist strategies and consequently contributing to their dominance in soil (Egidi *et al.*, 2019). On the other hand, Basidiomycota is primarily a slow-growing fungus and a less prolific sporulator than Ascomycota (Thormann and Rice, 2007), which might explain the low abundance of Basidiomycota in Sarawak's peatland. Less distribution of Zygomycota in peatlands was commonly reported than the other two taxa (Juan-Ovejero *et al.*, 2020; Liu *et al.*, 2020).

The dominance of several genera was observed and identified in all study sites. Species belonging to the genus *Aspergillus* were greater in number and more frequently isolated than other genera in PSF. *Aspergillus fumigatus* dominated the fungal communities in PSF. It is a saprophytic fungus known as a recycler in nature by degrading plant polysaccharides (Benoit *et al.*, 2015; Kwon-Chung and Sugui, 2013). Diverse metabolites enable this species to grow in different environmental conditions (Houbraken *et al.*, 2014). Thus, it was also detected under managed ecosystem like oil palm plantations.

The genus *Penicillium* was more frequently isolated in LOF, with unknown *Penicillium* sp. making up the fungal communities. *Penicillium* is a predominant fungus found in a wide range of environments and substrates, including peat soil (Yabuki *et al.*, 2014). *Penicillium* was previously detected in the peatlands of Argentina (Paredes *et al.*, 2014), Finland (Sun *et al.*, 2016), and Russia (Dobrovol'skaya *et al.*, 2012). *Penicillium* produces extracellular enzymes to degrade polysaccharide such as cellulose in plant cells (Vaishnav *et al.*, 2018; Zhang *et al.*, 2014). Therefore, the presence of *Penicillium* is important for plant biomass degradation and carbon cycling in the logged-over secondary forest ecosystem.

Trichoderma spp. were the dominant fungi and contributed the most to the fungal community in OPP. They are root-colonising fungi and important plant-symbiotic microbes, with a high distribution in agricultural soils (Jiang et al., 2016). A sampling conducted in 2012 also found that this group of fungi was more highly distributed in oil palm plantations than in other ecosystems (Kusai et al., 2018). The existence of Trichoderma spp. in plantations enhances plant growth and nutrient uptake efficiency, increases photosynthesis, improves root structure, and induces systemic resistance to abiotic and biotic stresses (Harman, 2011; Shoresh et al., 2010; Tyśkiewicz et al., 2022). Metabolites produced by Trichoderma spp. (Mendoza-mendoza et al., 2018) provide a promising biological control for severe oil palm disease, such as basal stem rot caused by Ganoderma boninense (Alexander et al., 2017; Supramani et al., 2022). Thus, the presence of *Trichoderma* may serve as a bioindicator for monitoring palm health and productivity besides reducing the use of chemical fungicides to control soil-borne diseases. It is still unknown why Trichoderma is exclusively isolated from oil palm plantations. Trichoderma may adapt to obtain nutrients from other soil organisms (Chaverri and Samuels, 2013) or enriched nutrients from soil environments that favour their growth in oil palm plantations.

Fewer species of Basidiomycota and Zygomycota were isolated from the sites in comparison to ascomycetes. Only one species of Basidiomycota was found in OPP, *i.e.*, *Polyporus ciliatus*. The species was previously reported in the Russian bog (Grum-Grzhimaylo *et al.*, 2016) and Caucasus regions, such as Iran, Turkey, and Russia (Gjobad-Nehjad, 2011). Two species of zygomycetes isolated from PSF, *Mortierella chlamydospora* and *Umbelopsis isabellina*, were previously isolated by Kusai *et al.* (2018) in 2012 from the same site, PSF. Despite the low isolated numbers of *M. chlamydospora* and *U. isabellina*, these species still reproduce and were detected in PSF in this study. *Mortierella chlamydospora* has a

specialised structure with a thick wall and resistant spores that can survive for an extended period and remain dormant in peat soil (Lin *et al.*, 2014). *Umbelopsis isabellina* has been reported to be highly distributed in boreal forest in Canada (Summerbell, 2005), while *M. chlamydospora* was found in agricultural areas in the United Kingdom (Hunt *et al.*, 2004).

This study showed that primary and secondary logged-over forests and managed ecosystems have different fungal communities and dominant species within the study sites. The overall compositions of the isolate assemblage that make up the fungal communities in PSF and LOF were more similar to one another compared to those in OPP. Notable differences were observed in the fungal community of OPP and PSF compared with LOF, as shown in the NMDS ordination, which can be attributed to the impacts of anthropogenic disturbances in the OPP (Kerfahi et al., 2014). These findings are consistent with Nurulita et al. (2016), who reported a distinct microbial community in the peatland forest that was converted into an oil palm plantation and rubber plantation in Indonesia. A similar finding in mineral soil was also reported in Peninsular Malaysia (Mcguire et al., 2015). Both studies demonstrated that fungal communities differ between primary forest and oil palm plantations compared to regenerating forests. Based on a study of fungal mycelium production by Robinson et al. (2020), fungal communities of saprophytic, mycorrhizal, and pathogenic species were significantly different between oil palm plantations and forests (old-growth and selectively logged). After six months of installation of hyphalin growth bags, mycelium fungal communities were unaffected by the logging event since both peat forest types showed similar fungal communities (Robinson et al., 2020). Although oil palm plantations showed different fungal communities, they can still support several ecologically important fungal groups like plant-symbionts (Trichoderma spp.) and saprotrophs (Aspergillus spp. and Penicillium spp.) (Sun et al., 2016), which are beneficial to plant development and decomposition processes, respectively.

In this study, LOF showed a higher diversity for both diversity indices when compared with PSF and OPP. This finding is consistent with a study by Mcguire *et al.* (2015), who revealed that the diversity of soil fungal communities is higher in regenerating forests (selectively logged 50 years ago) than in primary forest and oil palm plantations. Several aspects need to be considered when explaining this result, such as soil physical and chemical properties (Juan-Ovejero *et al.*, 2020) at each study site. Besides, the vegetation composition and chemical composition of litter could also contribute to the present results (Artz, 2009). LOF, a regenerating

forest logged years ago, would accumulate wood debris on the upper layer of peat soil, which might attract a more diverse species of fungi for the decaying processes. Logged-over secondary forests can retain the same taxonomy and structure of microbes as in a forest (Tripathi *et al.*, 2016b).

In this study, PSF was expected to have a higher fungal diversity than LOF and OPP due to the absence of anthropogenic activities at the site. Interestingly, lower fungal diversity and richness were recorded in PSF than in LOF. The result might be due to the dominance of Aspergillus spp., which may exclude less competitive or slowgrowing species during isolation (Thormann, 2006), and subsequently reduce diversity. A similar observation was recorded in OPP for *Trichoderma* spp. A study by Repečkienė *et al.* (2012) reported how the abundance of fast-growing and sporulating species like Trichoderma and Penicillium suppressed the growth of other fungal species in peatland. However, studies by Mcguire et al. (2015); Nurulita et al. (2016), and Brinkmann et al. (2019) showed that the fungal diversity-based Shannon diversity index in OPP was similar to or even slightly higher than the primary forest ecosystem. No significant effect on the fungal diversity due to the conversion of forests to agricultural lands, such as rubber and oil palm plantations, was reported in their studies that were conducted in Malaysia and Indonesia. Nurulita et al. (2016) who focused on peatland, showed that fungal diversity in OPP was not significantly affected by oil palm practices compared to rubber practices. The contrasting finding may be due to the method of analysis by Nurulita et al. (2016), who used the cultureindependent method PCR-DGGE, which identified a higher rate of microbial diversity compared to the culture-dependent method (Ayob and Kusai, 2021).

This study showed that the slightly reduced fungal diversity in OPP compared to the natural ecosystems is likely to be related to the addition of fertiliser (Berkelmann et al., 2020), organic matter stock (Allen et al., 2016), and vegetation richness (Drescher et al., 2016) which can all exert some effects on the fungal diversity in OPP. However, a managed ecosystem like OPP can support considerable fungal diversity and richness. Numerous factors can also influence the structure of the fungal diversity and community in ecosystems, such as soil properties (Birkhofer et al., 2012; Lauber et al., 2008; Zhang et al., 2016), vegetation diversity (Allingham et al., 2023; Dassen et al., 2017), hydrology (Lamit et al., 2021) and tree species (Prescott and Grayston, 2013; Sun et al., 2016; Urbanov et al., 2015) and further investigation is needed to provide more concrete evidence.

CONCLUSION

In summary, the composition of the fungal community and diversity were different between forests (PSF and LOF) and a managed peat ecosystem (OPP). However, similar species composition was recorded between PSF and LOF in comparison to OPP. Remarkable results were found in LOF, which recorded the highest fungal diversity and richness compared to PSF and OPP. OPP does not always negatively impact diversity as they still support several fungal species that are important to carry out ecosystem functions, such as the decomposition of organic matter. However, proper and sustainable management within the OPP needs to be applied to preserve its fungal diversity. Natural and managed peat ecosystems have their unique species composition, which is ecologically important in regulating the decomposition of peat and, consequently, for the carbon cycle. The presence and absence status of particular fungal species could be an indicator for monitoring ecosystem disturbances. Nevertheless, further studies using advanced technology, such as next-generation sequencing (NGS), especially for spatio-temporal analysis, are necessary for a more comprehensive understanding of soil fungal diversity in mediating biogeochemical cycles in tropical peatlands.

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