



Morpho-phylogenetic evidence for the first freshwater record of *Trichoderma protrudens* Samuels & P. Chaverri (Hypocreales, Sordariomycetes)

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Abstract. *Trichoderma protrudens* Samuels & P. Chaverri was originally described from the trunk of *Theobroma cacao* L. in Kerala, India, in 2008. Morphologically, it is defined by trichoderma-like conidiophores bearing subulate or lageniform phialides, green subglobose to obovoidal conidia, and the presence of chlamydospores in culture. Multilocus phylogenetic analyses based on ITS, *rpb2*, and *tefl-a* sequence data confirmed the identity of the isolates from the Philippines as *T. protrudens*, with robust support values (100% ML, 1.00 BPP) clustering them with the holotype CBS 121320. This study represents the first global record of *T. protrudens* in a freshwater ecosystem, expanding this species' known ecological range into aquatic environments. These findings emphasize the ecological versatility of *T. protrudens* and underscore the importance of further investigations into the fungal diversity of freshwater habitats.

Key words. Biodiversity, fungal systematics, microbial taxonomy, mycology, Philippines, tropical mycology

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INTRODUCTION

Lignicolous freshwater fungi, an important ecological group, thrive in freshwater habitats by utilizing submerged decaying plant matter (Jones et al. 2014). These fungi are vitally important in freshwater ecosystems in that they decompose lignocellulose and contribute to the carbon and nutrient cycles (Wong et al. 1998; Bucher et al. 2004; Hyde et al. 2016). Since 1996, extensive research has led to the description of over 3,870 freshwater fungal species, primarily ascomycetes (Calabon et al. 2022; Goh and Hyde 1996). The discovery of new taxa continues to accelerate, with no signs of slowing down (Calabon et al. 2023).

Trichoderma Pers. (Hypocreales) was first described by Pearson with *T. viride* Pers. as the type species (Pearson 1794). It is the most diverse member of the Hypocreales, with a global distribution and an estimated 400+ species (Sadfi-Zouaoui et al. 2009; Jaklitsch and Voglmayr 2015; Ma et al. 2020; Wijayawardene et al. 2020; Cai and Druzhinina 2021; Cai et al. 2022; Tang et al. 2022). The diversity and interest in *Trichoderma* stem from its production of numerous secondary metabolites with ecological and economic significance (Khan et al. 2020a). In agriculture, *Trichoderma* inoculants have been used for pest and disease control, and as growth promoters (Khan et al. 2020b; Zin and Badaluddin 2020; Subramaniam et al. 2022; Tyškiewicz et al. 2022; Yao et al. 2023; Kredics et al. 2024). Additionally, some *Trichoderma* species exhibit stress tolerance and can absorb heavy metals and organic pollutants, making them potential agents for environmental remediation (Bishnoi et al. 2007; Zafra and Cortés-Espinosa 2015; Cuevas and Banaay 2022).

In freshwater ecosystems, *Trichoderma* species play a crucial ecological role through their lignocellulolytic activity, which allows them to colonize and degrade lignocellulosic substrates, even in submerged aquatic habitats (Savory 1954; Calabon et al. 2023; Huang et al. 2024). The presence of saprobic *Trichoderma* in freshwater environments has been well-documented across several regions. For instance, studies by Au et al. (1992), Pinruan et al. (2014), Goh et al. (2018), and Concha et al. (2024) reported their occurrence on various submerged decaying plant substrates and sediments in China, Korea, Thailand, and the Philippines. Zheng et al. (2021) expanded the understanding of *Trichoderma* diversity in aquatic habitats by describing eight endophytic species associated with freshwater plants in China. These findings underscore the adaptability and functional importance of *Trichoderma* in freshwater ecosystems, particularly in processes involving organic matter decomposition and nutrient cycling.



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Traditionally, *Trichoderma* species have been identified and classified based on morphological analysis and growth rates (Bissett 1984, 1991a, 1991b; Harman and Kubicek 2002; Siddiquee 2017). However, morphological traits can be homoplastic, leading to challenges in identification (Druzhinina et al. 2005, 2006). DNA barcoding, using internal transcribed spacer (ITS), translation elongation factor 1-alpha (*tefl- α*), and RNA polymerase II gene (*rpb2*), has become a widely used method for *Trichoderma* and other hypocrealean fungi (Kuhls et al. 1996; Kullnig-Gradinger et al. 2002; Chaverri et al. 2015; Inglis et al. 2020; Cai and Druzhinina 2021; Cai et al. 2022). The simultaneous use of *tefl- α* and *rpb2* for phylogenetic analyses is particularly effective in identifying novel *Trichoderma* species, as these protein-coding genes offer greater resolution than ribosomal genes. (Hanada et al. 2008; Stielow et al. 2015; Zhu and Zhuang 2015; Qin and Zhuang 2016; Ye et al. 2023). The concatenation of multiple loci has been widely adopted to enhance resolution and accuracy in phylogenetic analyses, particularly in the reconstruction of fungal phylogenies (Delsuc et al. 2005; Robbertse et al. 2006; Marcet-Houben and Gabaldón 2009; Cai and Druzhinina 2021). A standardized protocol has been developed, requiring a *tefl- α* similarity of $\geq 97\%$ and an *rpb2* similarity of $\geq 99\%$ for species identification. Failure to meet these criteria suggests the potential for a novel species (Cai and Druzhinina 2021).

Aquatic *Trichoderma* species have been previously isolated from submerged substrates like sand sponges and decaying wood in marine environments (Paz et al. 2010; Gal-Hemed et al. 2011; Goh et al. 2018; Zheng et al. 2021). While freshwater *Trichoderma* has not been extensively surveyed in the Philippines, it has found application in agriculture as a sustainable alternative to synthetic fertilizers and fungicides (Cuevas et al. 2005; Launio et al. 2020). In this study, *Trichoderma* strains were isolated from submerged wood and bamboo substrates collected from a freshwater habitat in Iloilo, Philippines. This discovery was based on a comprehensive morphological and multilocus phylogenetic analysis, highlighting the diversity and potential of *Trichoderma* in freshwater ecosystems.

METHODS

Wood samples were collected from a waterfall in Miagao, Iloilo, Philippines (Figures 1, 2). These specimens were examined using the methods outlined by Senanayake et al. (2020). The incubated wood samples were initially observed under a stereomicroscope. The morphology of the observed fruiting bodies on the wood substrate was photographed using a Quaiqu Simul-Focal Trinocular dissecting microscope. Microscopic characters were captured using an Olympus BX53 microscope. The microphotographs were then processed using Adobe Photoshop (v. 21.1) and important characters were measured with ImageJ v. 154j. Single-spore isolation was conducted using potato dextrose agar (PDA) plates, following the procedures described by Senanayake et al. (2020). Mycelial growth patterns were observed on PDA, malt extract agar (MEA), and cornmeal agar (CMA) after seven and fourteen days. Fungal herbarium specimens and pure living cultures were stored and deposited in the University of the Philippines Visayas–Division of Biological Sciences Herbarium (UPVMI) and Culture Collection (UPVMICC) (Accessed on 2025-9-15), respectively.

Fresh fungal mycelia were scraped from colonies on PDA plates and placed in sterile 1.5 mL microcentrifuge tubes. Genomic DNA was extracted using QIAGEN DNeasy extraction kits following the manufacturer's protocol. Polymerase chain reaction (PCR) was used to amplify the ITS, *tefl- α* , and *rpb2* markers. The ITS markers were amplified using ITS5 and ITS4 primers (White et al. 1990). *tefl- α* and *rpb2* were amplified using EF1-983f and EF1-2218R (Rehner and Buckley 2005) and fRPB2-5f and fRPB2-7cr (Liu et al. 1999), respectively. Each 25 μ L PCR reaction consisted of 18.37 μ L nuclease-free water, 2.5 μ L genomic DNA, 2.5 μ L PCR buffer solution, 0.5 μ L dNTPs, 0.5 μ L forward and reverse primers, and 0.125 μ L HotStartaq DNA polymerase. The PCR program began with an initial denaturation at 95 °C for 15 min, followed by 35 cycles of denaturation

Figure 1. Location map showing the collection site of *Trichoderma protrudens*. The blue and red circles denote the new record in the Philippines (this study) and India (Degenkolb et al. 2008), respectively. The map was created using ArcGIS.

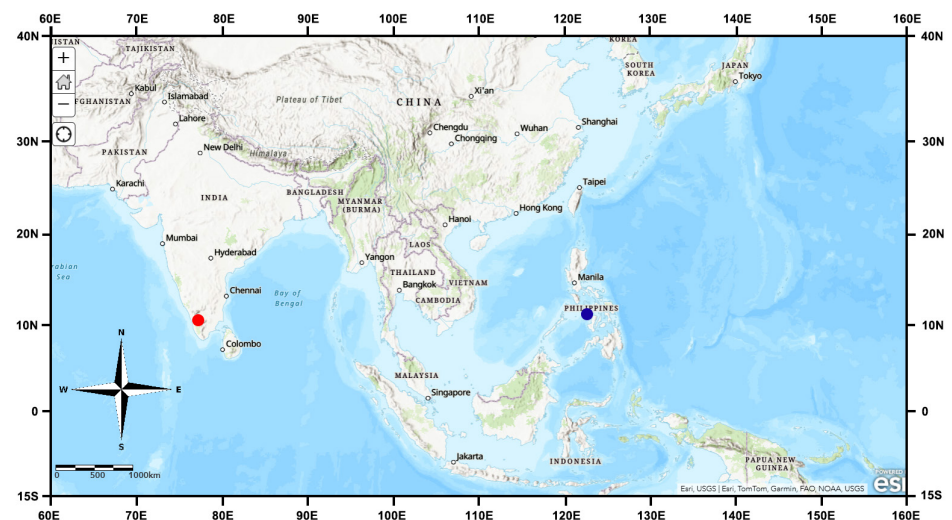




Figure 2. Habitat and collection of *Trichoderma protrudens* in the Philippines. **A.** Dilaan Falls, Miagao, Iloilo. **B, C.** Collection and sorting of samples. **D, E.** Unidentified submerged decaying woods: **(D)** UPVMI 24-0008 and **(E)** UPVMI 24.0009 (arrows point to colonies of *T. protrudens* on the natural substrate).

(1 min at 95 °C), annealing (30 s at 56 °C), and elongation (1 min at 72 °C). A final extension step was performed at 72 °C for 10 min. The presence of amplicons was confirmed using agarose gel electrophoresis on a Mupid One Electrophoresis System. The PCR products were sequenced on the Illumina iSeq100 platform at the Philippine Genome Center Visayas, University of the Philippines Visayas, Miagao, Philippines.

The taxa included in our phylogenetic analysis (Table 1) were assembled based on the closest matches from the BLAST search results and from recently published data of Brito et al. (2023). Sequences generated for the three target genes were combined with additional sequences retrieved from GenBank (Table 1). Three separate datasets, one for each gene, were aligned using MAFFT v. 7 (Katoh et al. 2019). Alignments were then manually refined where necessary with BioEdit v. 7.0.9.0 (Hall 1999). TrimAl v. 1.3 was employed for automatic trimming of aligned sequences. The online tool ALTER (Glez-Peña et al. 2010) facilitated conversion of the alignment file to Phylip format. Both individual gene data and combined data were subjected to phylogenetic analysis using the maximum-likelihood (ML) method. RAxML v. 8 software on the CIPRES web portal (Stamatakis 2014) (<http://www.phylo.org/portal2/>; Miller et al. 2010) was utilized for this purpose. The GTR+GAMMA model of nucleotide evolution was adopted, and RAxML rapid bootstrapping was performed with 1,000 replicates. Additionally, Bayesian-inference analysis was conducted with MrBayes v. 3.2.6 on XSEDE at the CIPRES web portal (Ronquist and Huelsenbeck 2003). Newly generated sequences were deposited in GenBank (Table 1).

Table 1. Taxa used in this study for the analysis of combined ITS rDNA, *rpb2*, and *tefl-a* sequence data and their GenBank accession numbers. The newly generated sequences are indicated with asterisk (*).

Taxon	Strain / voucher number	ITS	<i>rpb2</i>	<i>tefl-a</i>
<i>Protocrea farinosa</i>	CBS 121551	MH863119	EU703935	EU703889
<i>P. pallida</i>	CBS 121552	EU703922	EU703944	EU703897
<i>Trichoderma albolutescens</i>	CBS 119286	FJ860721	FJ860517	FJ860609
<i>T. applanatum</i>	7792	KJ783291	KJ634726	KJ634759
<i>T. arundinaceum</i>	37MTS	OR880618	–	PP263665
	GJS 05-184	EU330933	EU338308	EU338280
	NRRL 3199	EU330932	EU338307	EU338279
<i>T. atlanticum</i>	CBS 120632	FJ860781	FJ860546	FJ860649
<i>T. atroviride</i>	CBS:142.95	AF456917	EU341801	AF456891
<i>T. aurantioeffusum</i>	CBS 119284	FJ860728	FJ860520	FJ860613
	CPK 3119	FJ860730	FJ860521	FJ860614
	S565	–	–	KJ665430
<i>T. austriacum</i>	CBS 122494	FJ860735	FJ860525	FJ860619
<i>T. brasiliensis</i>	COAD 2324	MK713497	MT300488	MT300486

Taxon	Strain / voucher number	ITS	rpb2	teff- α
	COAD 2642	MK713498	MT300490	MT300487
	COAD 2643	MK713499	MT300489	MT300492
	COAD 2644	MK713514	MT300491	MT300493
<i>T. brevicompactum</i>	CBS 109720	AY857249	DQ857347	AY324176
	CBS 112443	EU330943	EU338319	EU338281
	CBS 112444	EU330938	EU338314	EU338296
	CBS 112447	EU330942	EU338318	EU338300
<i>T. citrinum</i>	CBS 894.85	MH861923	AF545561	DQ835441
<i>T. grande</i>	HMAS:248749	–	KX066266	KX066254
	HMAS:273788	–	–	KX066255
<i>T. limonium</i>	HMAS:248751	–	KX066259	KX066247
	HMAS:248754	–	KX066260	KX066248
<i>T. luteoeffusum</i>	CBS 120537	FJ860773	FJ860543	FJ860645
<i>T. margaretense</i>	C.P.K. 3127	FJ860741	FJ860529	FJ860625
<i>T. oligosporum</i>	HMAS:252870	KJ783296	KJ634731	KJ634764
<i>T. peltatum</i>	GJS 08-207	HM466659	HQ260610	KR135819
<i>T. phellincola</i>	CBS 119283	FJ860808	FJ860569	FJ860672
<i>T. polysporum</i>	CBS:820.68	MH859230	DQ087238	AF400989
<i>T. protopulvinatum</i>	CBS 739.83	DQ835405	DQ835463	DQ835487
<i>T. protrudens</i>	CBS 121320	EU330946	EU338322	EU338289
	UPVMICC 24-0008*	PQ451471	PQ790167	PQ790164
	UPVMICC 24-0009*	PQ451472	–	PQ790165
<i>T. pseudostramineum</i>	TUFC60104	JQ797392	JQ797408	JQ797400
<i>T. psychrophilum</i>	C.P.K. 2435	FJ860819	FJ860576	FJ860682
<i>T. pulvinatum</i>	CBS 121279	FJ860820	FJ860577	FJ860683
<i>T. rhododendri</i>	CBS 119288	FJ860822	FJ860578	FJ860685
<i>T. rodmanii</i>	GJS 91-88	EU330948	EU338324	EU338286
	CBS:121553	FJ860824	FJ860580	FJ860687
	CPK 2852	FJ860825	FJ860581	FJ860688
<i>T. seppoi</i>	CBS:122498	FJ860833	FJ179617	FJ179581
<i>T. stercorarium</i>	CBS:148.85	MH861866	EF469103	AF543782
<i>T. subsulphureum</i>	M 141	DQ835509	DQ835522	DQ835492
<i>T. tiantangzhaiense</i>	HMAS 252872	KJ783295	KJ634730	KJ634763
<i>T. turrialbense</i>	CBS 112445	EU330945	EU338321	EU338284
	BBA 72294	EU330944	EU338320	EU338282
	SFC20160125-C300	–	–	MZ634448
<i>T. undulatum</i>	PPRI:19365	KX267810	KX351319	KX267789
<i>T. victoriense</i>	CBS:140064	DQ835505	DQ835517	DQ835500
<i>T. viride</i>	CBS:119325	DQ323428	EU711362	DQ672615

RESULTS

Trichoderma protrudens Samuels & P. Chaverri, Mycol. Progr. 7(3): 212 (2008)

Figure 3

New record. PHILIPPINES – ILOILO • Miagao, Dilaan Falls; 10°44'37.98"N, 122°11'22.0"E; 23.IX.2023; B.S. Dangallo leg.; isolated from submerged decaying unidentified wood; tropical monsoon climate; GenBank ITS rDNA PQ451471; *teff- α* PQ790164; *rpb2* PQ790167; UPVMI 24-0008; living culture UPVMICC 24-0008; GenBank ITS rDNA PQ451472; *teff- α* PQ790165; UPVMI 24-0009; living culture UPVMICC 24-0009.

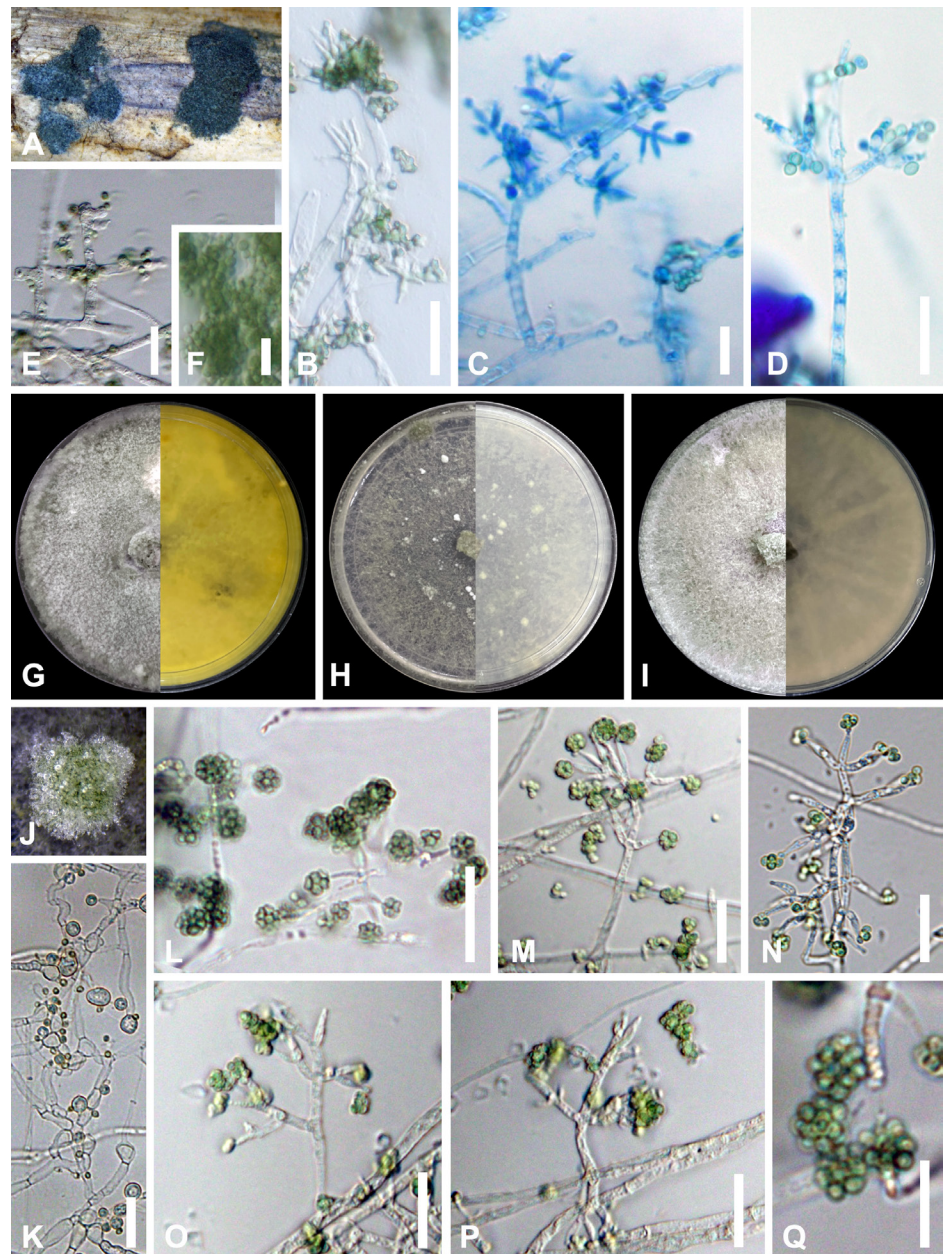


Figure 3. *Trichoderma protrudens* (24-0008) **A.** Appearance of fungi on bamboo. **B-E.** Conidiophore and phialides: (**B-D**) with and (**E**) without lactophenol cotton blue stain. **F.** Conidia. **G-I.** Colonies after 14 days: (**G**) on potato dextrose agar; (**H**) on corn meal agar; (**I**) on malt extract agar. **J.** Active sporulation on CMA. **K.** Chlamydozoospores. **L-Q.** Conidiophores and phialides with conidia clusters. Scale bars: B-E, K, L-P = 20 μm ; F, Q = 10 μm .

Identification. *Saprobic* on decaying wood submerged in freshwater. *Description on natural substrate:* sexual morph undetermined. Asexual morph: hyphomycetous. Conidial pustules, dark bluish green, irregular shape, outline not well defined. Conidiophores $35\text{--}155 \times 1.9\text{--}4.1 \mu\text{m}$ ($\bar{x} = 67.1 \times 2.9 \mu\text{m}$, $n = 30$), trichoderma-like, hyaline, often symmetry, repeatedly branched toward the base forming a pyramidal structure with phialides in whorl, smooth-walled, often with a main axis, side branches in straight angles with main axis. Phialides $10\text{--}18 \mu\text{m}$ length ($\bar{x} = 14.2 \mu\text{m}$, $n = 50$), $1.9\text{--}3.8 \mu\text{m}$ maximum width ($\bar{x} = 2.8 \mu\text{m}$), $1.7\text{--}2.7 \mu\text{m}$ base width ($\bar{x} = 2.3 \mu\text{m}$), typically formed in whorls of 3 or rarely solitary, subulate or lageniform. Conidia $3.5\text{--}5.0 \times 1.9\text{--}2.9 \mu\text{m}$ ($\bar{x} = 4.1 \times 2.5 \mu\text{m}$, $n = 50$), green, smooth, subglobose to obovoidal.

Culture characteristics: on MEA: after 72 h, colony radius reached 33–38 mm at 25 °C, colonies on MEA reached 90 mm diameter within 5 days at 25 °C; Colony radial, mycelium common, aerial hyphae more abundant in colony margin. Conidiation starting after 6 days, light-green conidia formed scarcely on aerial hyphae: few pustules. On PDA: after 72 h, colony radius reached 32–36 mm at 25 °C. Conidiation starting after eight days, light-green conidia formed scarcely on aerial hyphae, more abundant around the margin of the colony; few pustules. No pigment diffusing through the agar; no distinctive odor. On CMA: after 72 h, colony radius reached 28–31 mm at 25 °C, colonies reached 90 mm diameter at four days and 25 °C; cottony white aerial mycelium, light-green sporulation starting from the center of the plate, pustules abundant. Conidiation starting after 5 days, light-green conidia formed on aerial hyphae around margin of colony. No pigment diffusing through the agar; no distinctive odor. Conidiophores trichoderma-like, micronematous, pyramidal with phia-

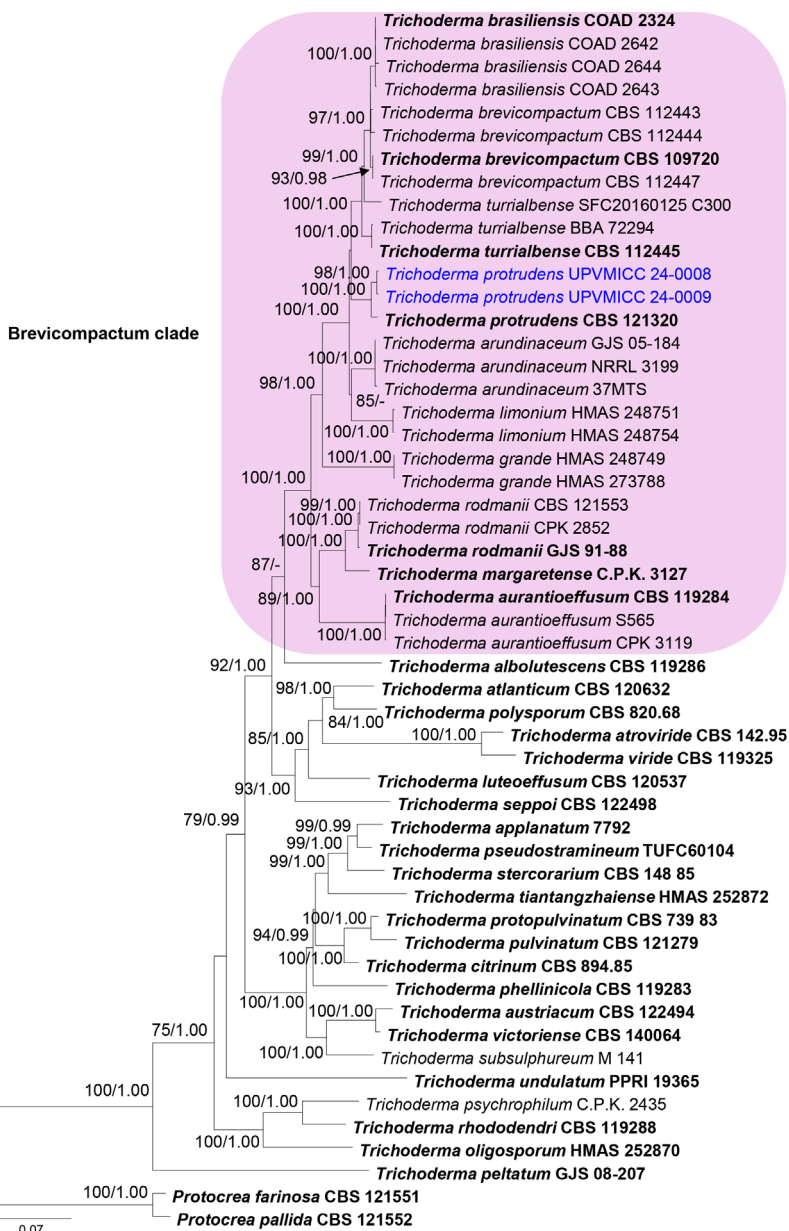
lides in whorl, smooth-walled. Phialides 9–27 µm length (\bar{x} = 16, n = 50), 2–5 µm maximum width (\bar{x} = 3 µm), 1.8–4.4 µm base width (\bar{x} = 2.7 µm), hyaline, typically formed in whorls of 3 or rarely solitary, subulate or lageniform. Conidia 1.6–3.3 × 1.9–2.9 µm (\bar{x} = 2.2 × 2.1 µm, n = 50), pale greenish, smooth, subglobose to obovoidal. Chlamydospores 2.4–9.2 × 2.6–9.7 (\bar{x} = 6.6 × 5.9 µm, n = 30), hyaline, abundant, globose.

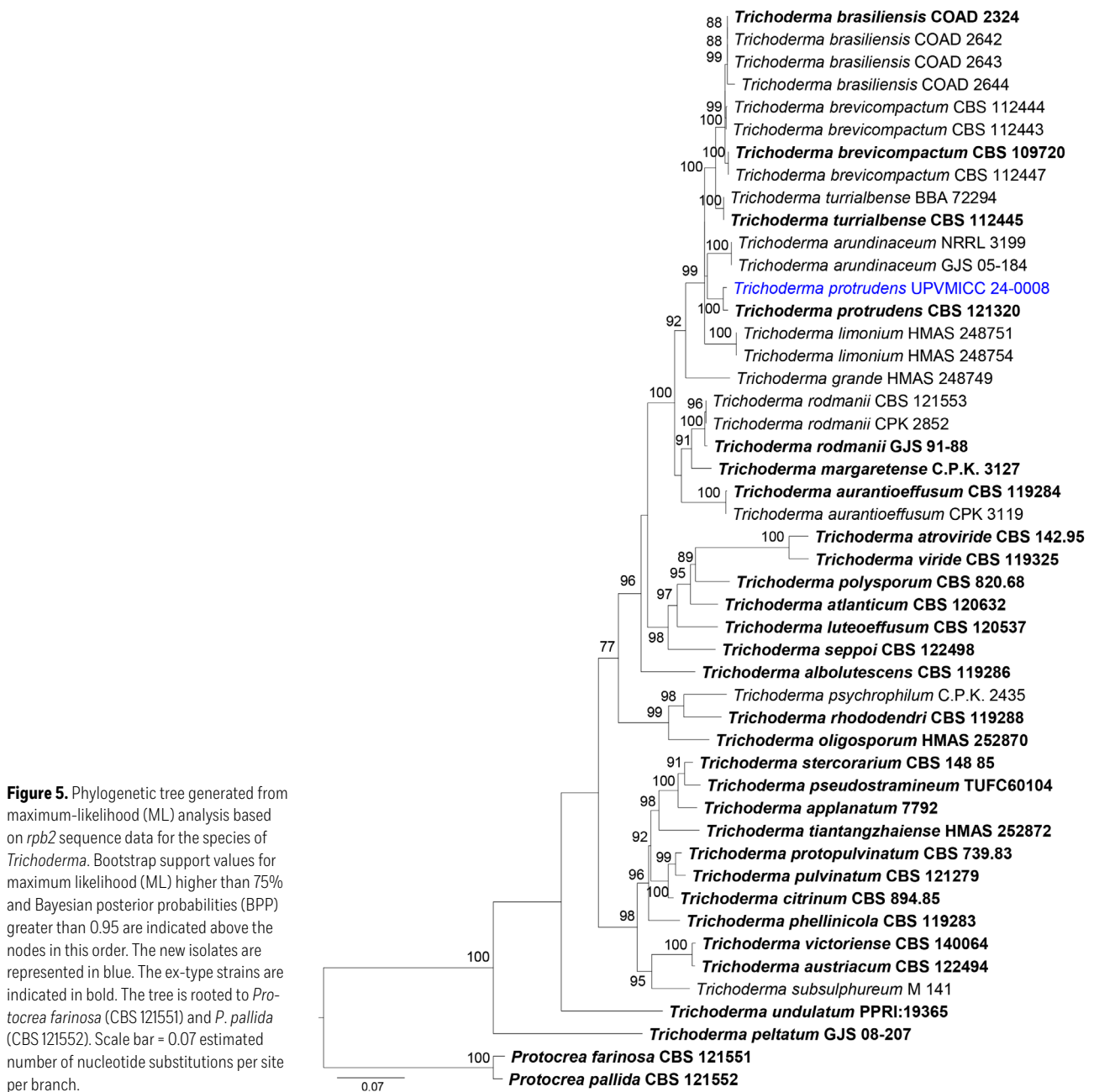
In a multilocus phylogenetic analysis, strains UPVMICC 24-0008 and UPVMICC 24-0009 clustered with the holotype CBS 121320 with strong statistical support (100% ML, 1.00 BPP) (Figure 4). The single-locus trees revealed that these strains fall within the brevicompactum clade (Figures 5, 6). In the *rpb2* phylogenetic tree, strain UPVMICC 24-0008 clustered with the holotype CBS 121320 with strong statistical support (100% ML) (Figure 5). However, in the *tefl-a* locus, both UPVMICC 24-0008 and UPVMICC 24-0009 exhibited a distinct lineage basal to *T. protrudens* (Figure 6). This unique lineage in the *tefl-a* tree could be attributed to the use of primers EF1-983f and EF1-2218R, as opposed to the standard *tefl-a* barcodes (EF1 and EF2) recommended by Cai and Druzhinina (2021). Morphologically, *T. protrudens* UPVMICC 24-0008 and UPVMICC 24-0009 have trichoderma-like conidiophore with smaller conidia [1.6–3.3 × 1.9–2.9 µm vs. (2.5–)2.7–3.2(–3.5) × (2.2–)2.5–2.7(–3.0)], and develops chlamydospores in the culture medium, features that distinguish the isolate from the holotype CBS 121320 (Degenkolb et al. 2008). Based on morphological characteristics and phylogenetic analyses of ITS, *rpb2*, and *tefl-a* regions, these two isolates have been identified as *T. protrudens*, marking the first report of this species in a freshwater ecosystem globally.

Habitat. Trunk of *Theobroma cacao* (Degenkolb et al. 2008); on decaying wood submerged in a freshwater stream (this study).

Known distribution. India (Degenkolb et al. 2008), Philippines (this study).

Figure 4. Phylogenetic tree generated from maximum-likelihood (ML) analysis based on ITS, *rpb2*, and *tefl-a* sequence data for the species of *Trichoderma*. Bootstrap support values for maximum likelihood (ML) higher than 75% and Bayesian posterior probabilities (BPP) greater than 0.95 are indicated above the nodes in this order. The new isolates are represented in blue. The ex-type strains are indicated in bold. The tree is rooted to *Protocrea farinosa* (CBS 121551) and *P. pallida* (CBS 121552). Scale bar = 0.07 estimated number of nucleotide substitutions per site per branch.



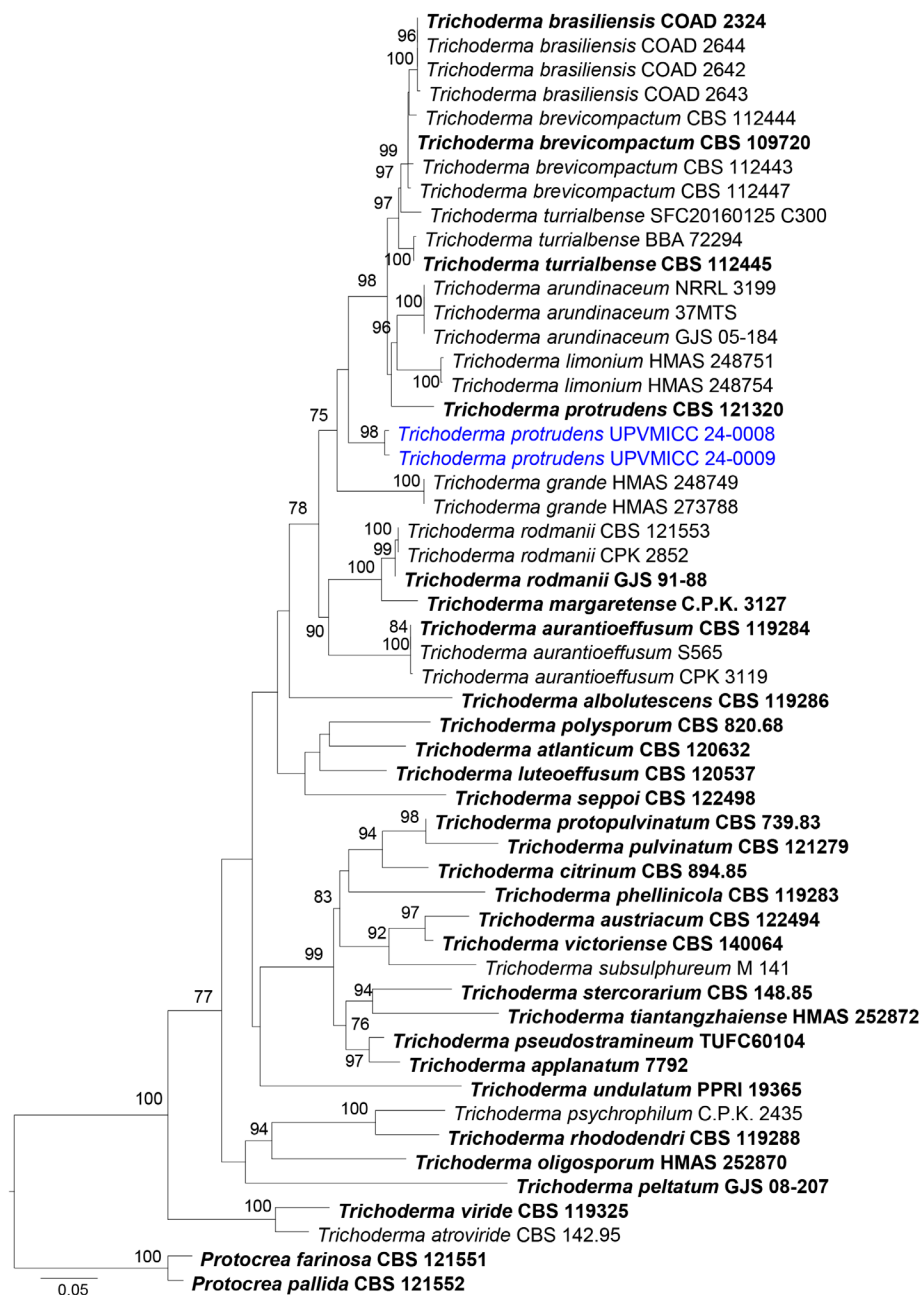


DISCUSSION

Based on a multilocus phylogenetic analysis of combined datasets of ITS, *rpb2*, and *tef1- α* , the newly isolated strains were placed within the brevicompactum clade. The brevicompactum clade, one of the infrageneric clades of *Trichoderma*, is distinguished by its relatively small number of species (only 14) and its characteristic morphological features, including globose to subglobose conidia and ampulliform to lageniform phialides. While some species within this clade have been observed to produce sexual morphs (e.g., *T. rodmanii* (Samuels & P. Chaverri) Jaklitsch & Voglmayr, *T. grande* W.T. Qin & W.Y. Zhuang, *T. limonium* W.T. Qin & W.Y. Zhuang), the majority are known only as asexual morphs (Degenkolb et al. 2008).

Aquatic Hypocreales encompass 80 genera across six families. Most of these genera (57) belong to the family Nectriaceae (Calabon et al. 2022). Over the past decade, there has been a surge in the discovery and description of novel freshwater Hypocreales species and genera. For instance, *Penicillifer sinicus* Z.Q. Zeng & W.Y. Zhuang, *Fusicolla aquaeductum* (Radlk. & Rabenh.) Gräfenhan, Seifert & Schroers, and *Aquanectria jacinthicolor* S.K. Huang, Jeewon & K.D. Hyde were identified in China, while the genera *Varicosporella* Lechat & J. Fourn. and *Cosmosporella* S.K. Huang, Jeewon & K.D. Hyde were discovered in France and China, respectively (Lechat and Fournier, 2015; Huang et al. 2018; Zeng and Zhuang 2022). The adaptations of Hypocreales to freshwater habitats remain a subject of ongoing research. While there is no clear evidence of specialized

Figure 6. Phylogenetic tree generated from maximum-likelihood (ML) analysis based on *tefl-a* sequence data for the species of *Trichoderma*. Bootstrap support values for maximum likelihood (ML) higher than 75% and Bayesian posterior probabilities (BPP) greater than 0.95 are indicated above the nodes in this order. The new isolates are represented in blue. The ex-type strains are indicated in bold. The tree is rooted to *Protocrea farinosa* (CBS 121551) and *P. pallida* (CBS 121552). Scale bar = 0.05 estimated number of nucleotide substitutions per site per branch.



appendages, some species of Nectriaceae exhibit apical rings that may contribute to their aquatic lifestyle (Bao et al. 2023). The total number of *Trichoderma* species described is estimated to be between 400 and 600 (Sadfi-Zouaoui et al. 2009; Jaklitsch and Voglmayr 2015; Ma et al. 2020; Wijayawardene et al. 2020). According to *Index Fungorum* (as of 2024-10-10), there are 476 individual species recorded, excluding former species and duplicates. Recent studies have identified sixteen *Trichoderma* species in freshwater environments, where they function as saprobes and endophytes on various substrates across China, Korea, Philippines, and Thailand (Au et al. 1992; Pinruan et al. 2014; Goh et al. 2018; Zheng et al. 2021; Concha et al. 2024). A synopsis of freshwater *Trichoderma* species is provided in Table 2. These species are classified within six distinct *Trichoderma* species complexes: atroviride (*T. uncinatum*), hamatum (*T. pseudoasperelloides*), harzianum (*T. afroharzianum*, *T. guizhouense*, *T. harzianum*, *T. pinicola*), koningii (*T. tibetica*), longibrachiatum (*T. aquaticum*, *T. capillare*, *T. reesei*, *T. saturnisporum*), and viridescens (*T. asymmetricum*, *T. inconspicuum*, *T. paraviridescens*, *T. scorpioideum*). Despite this relatively small number in freshwater environments, the past decade has witnessed a significant increase in the discovery of *Trichoderma* species in terrestrial habitats, particularly in China (Qin and Zhuang 2016; Zheng et al. 2021; Ye et al. 2023; Zhao et al. 2023, 2024).

Taxonomic research on freshwater fungi in the Philippines remains underexplored, with most species' identification based primarily on morphological characteristics. Over the past two decades, only four species of freshwater fungi have been described: *Paoayensis lignicola* Cabanela, Jeewon & K.D. Hyde (Cabanela et al.

Table 2. Synopsis of morphological characters of freshwater *Trichoderma* species.

Taxon	Conidiophore		Phialides			Conidia			Chlamydospores		Reference	
	Size (µm)	Shape	No.	Size (µm)	Shape	Size (µm)	Shape	Septa	Wall	Size (µm)		Shape
<i>T. afroharzianum</i>	–	Pyramidal with opposing, widely spaced branches	1–5	3.5–17.5 × 2.0–4.2	Lageniform to ampulliform	2.0–4.5 × 2.0–4.0	Subglobose to ovoid	–	–	Rare	–	Chaverri et al. 2015
<i>T. aquaticum</i>	5.5–19.1	Pyramidal; alternately or irregularly branched	3–5	5.2–11.5 × 2.5–4.2	Lageniform, rare ampulliform, slightly curved	3.3–4.8 × 2.4–3.3	Mostly ovoid, ellipsoidal, rarely subglobose	–	Thin	4.3–7.6 × 3.91–7.2	Globose	Zheng et al. 2021
<i>T. asymmetricum</i>	–	Tree-like, asymmetrically branched	1–3	4.4–12.6 × 2.3–4.4	Occasionally lageniform, with obtuse apex	3.5–5.0 × 3.2–4.1	Subglobose to globose, sometimes broadly ellipsoidal, rarely ovoid	–	–	–	–	Zheng et al. 2021
<i>T. capillare</i>	–	Lateral branching with branches terminating in whorls of few phialides; sometimes, lacking branches lack obvious patterns	–	4.0–14.0 × 2.2–4.5	Lageniform, nearly cylindrical or conspicuously swollen below middle, straight or less frequently asymmetric or hooked	2.2–4.5 × 1.7–4.0	Subglobose to broadly ellipsoidal	–	–	Not observed	–	Samuels et al. 2012
<i>T. guizhouense</i>	–	Verticillate, forms a more or less pyramidal structure	2–3	4.5–10 × 2–3	Ampulliform to lageniform	2–3 (diam.)	Globose	–	Smooth	Not observed	–	Li et al. 2012
<i>T. harzianum</i>	–	Pyramidal with opposing branches	2–5	4.7–16 × 2.7–4.7	Ampulliform to lageniform	2.2–4.2 × 2.0–3.7	Subglobose to ovoid	–	–	Rare	–	Chaverri et al. 2015; Jang et al. 2018
<i>T. inconspicuum</i>	–	Tree-like to irregular branched	3–4	6.4–16.2 × 1.9–4.0	Narrow lageniform, some swollen in the middle, ampulliform	2.8–4.5 × 2.6–4.0	Subglobose to globose; few are ovoid	–	Thin	6.6–10.1 × 5.8–8.7	Globose, ellipsoidal	Zheng et al. 2021
<i>T. paraviride</i>	–	Curved to sinuous, branched irregularly or paired in tree fashion	3–5	9.2–16.4 × 2.1–4.1	Spindly to ellipsoid, rarely cylindrical, sometimes curved	3.4–5.0 × 3.2–4.8	Globose to subglobose	–	Thin	–	–	Zheng et al. 2021
<i>T. paraviridescens</i>	–	Tree/branch-like from, wide side branches	2–3	5.6–12.3 × 2.0–3.4	Lageniform	2.3–4.9 × 1.82–3.5	Globose to subglobose	–	–	6.08–11.17 (diam.)	Globose	Goh et al. 2018
<i>T. pinicola</i>	–	Pyramidal, branched repeatedly toward base	–	9–20 × 2–6	Subulate or lageniform	3.5–5.0 × 1.9–2.9	Oblong to ellipsoid	–	–	6.2–10 × 4–7	Globose	Concha et al. 2024
<i>T. protrudens</i>	–	broad central axis with lateral and often terminal fertile branches	–	–	Lageniform to ampulliform	(2.5–)2.7–3.2(–3.5) × (2.2–)2.5–2.7(–3.0)	Subglobose to ovoidal	–	–	–	–	Degenkolb et al. 2008
<i>T. pseudoasperelloides</i>	–	Dendrifor, with little rebranching	3–5	5.0–9.0 × 2.0–5.0	Spindle-shaped and lageniform	3.7–4.1 × 2.7–3.8	Subglobose to ellipsoidal, rarely globose	–	Thin	–	–	Zheng et al. 2021
<i>T. reesei</i>	–	Long central axis with solitary phialides before secondary branching	–	3.0–11.5 × 1.8–4.0	Lageniform, asymmetric, swollen or not at middle, straight, less frequently sinuous	3.0–6.0 × 1.5–3.5	Ellipsoidal to oblong	–	–	–	Subglobose to pyriform	Goh et al. 2018
<i>T. saturni-sporum</i>	–	Highly variable in form	–	5.5–14.0 × 2.0–3.5	Lageniform to ampulliform, often curved	3.0–6.0 × 2.0–4.0	Subglobose to ellipsoidal or sub-cylindrical	–	–	–	–	Goh et al. 2018
<i>T. scorpioidium</i>	–	Slightly curved main axis, generally verticillate branches	2–3	6.5–13.3 × 2.0–3.9	Narrow lageniform, some are ellipsoidal	3.3–4.4 × 2.4–3.8	Globose to subglobose, a few ovoidal to ellipsoidal	–	Thin	–	–	Zheng et al. 2021
<i>T. tibetica</i>	–	Slightly curved main axis, slightly upward and sometimes perpendicular branches	2–4	5.2–12.3 × 2.1–3.5	Lageniform, sometimes globose, ellipsoidal, or pyramidal	3.7–4.8 × 2.9–4.0	Ellipsoidal to ovoid, subglobose rarely noted	–	–	–	–	Zheng et al. 2021
<i>T. uncinatum</i>	–	Hard-discernable, slightly curved main axis, irregular alternate branches	2–5	4.3–10.3 × 2.3–3.9	Lageniform, sometimes ampulliform to subglobose	3.1–4.4 × 2.7–4.0	Globose to subglobose, rarely ovoidal or ellipsoidal	–	Thin	–	–	Zheng et al. 2021

2007), *Phaeoacremonium philippinense* M.S. Calabon, Bagacay, Nim, Sadaba & E.B.G. Jones (Calabon et al. 2024), *Sporoschisma chiangraiense* N.G. Liu & K.D. Hyde, and *Trichoderma pinicola* Seung Y. Oh, M.S. Park & Y.W. Lim (Concha et al. 2024), that employed a phylogenetic analysis of combined nuclear ribosomal DNA loci (LSU, SSU, ITS) and protein-coding genes (*tub2*, *act*, *tefl-a*). The Philippines remains underexplored in terms of freshwater fungal systematics research, especially when compared to nearby Asian countries such as China and Thailand (Zhang et al. 2011; Hu et al. 2013; Calabon et al. 2022). This underscores the vast yet largely untapped fungal diversity present in Philippine freshwater ecosystems. Comprehensive taxonomic surveys are essential to bridge this knowledge gap, integrating both morphological and molecular phylogenetic approaches to better characterize the region's fungal biodiversity. Globally, the occurrence and ecological roles of

Trichoderma in freshwater ecosystems are also underrepresented in scientific literature. Despite its ecological significance, few studies have been published, leaving its geographical distribution and functional roles largely unexplored. To address these gaps, extensive surveys are necessary to elucidate the ecological functions and distribution patterns of *Trichoderma* in freshwater environments.

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ADDITIONAL INFORMATION

Conflict of interest

The authors declare that no competing interests exist.

Ethical statement

No ethical statement is reported.

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Author contributions


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Data availability

All data that support the findings of this study are available in the main text.

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