Identification and Pathogenicity of *Paramyrothecium* Species Associated with Leaf Spot Disease in Northern Thailand

Patchareeya Withee 1, Sukanya Haituk 1, Chanokned Senwanna 1, Anuruddha Karunarathna 1, Nisachon Tamakaew 1, Parichad Pakdeeniti 1, Nakarin Suwannarach 2,3, Nakarin Suwannarach 2,3, Jaturong Kumla 2,3, Piyawan Suttiprapan 1,4, Paul W. J. Taylor 5, Milan C. Samarakoon 1,* and Ratchadawan Cheewangkoon 1,2,4,*

1 Department of Entomology and Plant Pathology, Faculty of Agriculture, Chiang Mai University, Chiang Mai 50200, Thailand; patchareeya_withee@cmu.ac.th (P.W.); sukanay_hai@cmu.ac.th (S.H.); chanokned.swnn@gmail.com (C.S.); anumandrack@yahoo.com (A.K.); nisachon_t@cmu.ac.th (N.T.); parichad_pakdee@cmu.ac.th (P.P.); piyawan.s@cmu.ac.th (P.S.)

2 Research Center of Microbial Diversity and Sustainable Utilization, Chiang Mai University, Chiang Mai 50200, Thailand

3 Department of Biology, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand

4 Innovative Agriculture Research Centre, Faculty of Agriculture, Chiang Mai University, Chiang Mai 50200, Thailand

5 Faculty of Veterinary and Agricultural Sciences, The University of Melbourne, Parkville, VIC 3010, Australia; paulwjt@unimelb.edu.au

* Correspondence: milanchameerasam.s@cmu.ac.th (M.C.S.); ratchadawan.e@cmu.ac.th (R.C.)

Abstract: Species of *Paramyrothecium* that are reported as plant pathogens and cause leaf spot or leaf blight have been reported on many commercial crops worldwide. In 2019, during a survey of fungi causing leaf spots on plants in Chiang Mai and Mae Hong Son provinces, northern Thailand, 16 isolates from 14 host species across nine plant families were collected. A new species *Paramyrothecium vignicola* sp. nov. was identified based on morphology and concatenated (ITS, cmdA, rpb2, and tub2) phylogeny. Further, *P. breviseta* and *P. folicola* represented novel geographic records to Thailand, while *P. eichhorniae* represented a novel host record (*Psophocarpus* sp., *Centrosema* sp., *Aristolochia* sp.). These species were confirmed to be the causal agents of the leaf spot disease through pathogenicity assay. Furthermore, cross-pathogenicity tests on *Coffea arabica* L., *Commelina benghalensis* L., *Glycine max* (L.) Merr., and *Diefenbachia seguine* (Jacq.) Schott revealed multiple host ranges for these pathogens. Further research is required into the host–pathogen relationship of *Paramyrothecium* species that cause leaf spot and their management. Biotic and abiotic stresses caused by climate change may affect plant health and disease susceptibility. Hence, proper identification and monitoring of fungal communities in the environment are important to understand emerging diseases and for implementation of disease management strategies.

Keywords: climate change; diversity; food security; multi-gene phylogeny; new species; plant pathology; taxonomy

1. Introduction

Plant diseases have a high impact on food security [1] and fungi play a major role in plant diseases [2]. Foliar fungal pathogens severely affect the yield and health of commercial crops [3]. Leaf spots are an early indicator of foliar diseases and may initially occur on the adaxial leaf surfaces and then appear on the abaxial leaf surface.

*Paramyrothecium* species have been frequently identified to cause leaf spot and blight disease on a wide range of vegetables, ornamental plants, and economic crops [4–7]. Disease symptoms caused by *Paramyrothecium* may also include stem and crown canker and fruit rot [8–10]. Lombard et al. [4] designated an epitype for the generic type *Paramyrothecium roridum* (≡*M. roridum*). *Paramyrothecium* species are distinguished from related *Myrothecium* species...
sensu stricto and other myrothecium-like genera by the presence of 1–3 septate, thin-walled setae surrounding the sporodochia. Currently, there are 19 species listed in Index Fungorum (http://www.indexfungorum.org/; accessed on 14 April 2022).

*Paramyrothecium roridum* and *P. foliicola* are well-known pathogens that cause leaf spot or leaf blight and have been reported on many commercial crops and a wide range of hosts, such as soybean, strawberry, and muskmelon [5,8,9]. Rennberger and Keinath [11] isolated *P. foliicola* and *P. humicola* from watermelon and two other cucurbits and confirmed their pathogenicity on watermelons, tomatoes, and southern peas. Aumentado and Balandres [12] reported *P. foliicola* causing crater rot in eggplant and 45 plant species from 21 plant families and were tested for the pathogenicity on detached fruit or leaf assays. Furthermore, *P. foliicola* is pathogenic to cucumber seedlings and watermelon, causing stem canker [13]. Due to the lignicolous nature of the *Paramyrothecium*, they are being used as bio-pesticides for the control of weeds and insects [14,15]. Interestingly, several important secondary metabolites or toxins found in *Paramyrothecium* include trichothecenes macrolides such as roridin, verrucarin, and mytoxin B, which are important for some medicinal and biotechnological applications [16–18].

In Thailand, only *P. eichhorniae* has been reported and this was identified as the cause of the leaf blight disease of water hyacinth [19]. The diversity of *Paramyrothecium* species in Thailand is unknown. As a result, surveys and additional research on the distribution of *Paramyrothecium* in Thailand is required. The objective of this study was to identify and describe *Paramyrothecium* spp. from northern Thailand and assess their pathogenicity across a broad range of potential host plant species.

2. Results

2.1. Symptoms

Leaf spots varying in size and shape, depending on the host, were most visible on the upper surface. The leaf spots consisted of small brown spots or necrotic lesions with a dark border, while in older lesions, small sporodochia were visible (Figure 1f,n,o). Necrotic lesions appeared dark gray or black on *Centrosema* sp., *Coccinia grandis*, *Oroxylum indicum*, *Solanum virginianum*, *Tectona grandis*, *Vigna mungo*, *Vigna* sp., and *V. unguiculata* (Figure 1a,d,e,g,h–k,m), and surrounded by a prominent yellow halo on *Lablab purpureus*, *Psophocarpus* sp., and *Spilanthes* sp. (Figure 1b,c,l). Lesions on *Aristolochia* sp., *Coffea arabica*, and *Commelina benghalensis* consisted of light to dark brown concentric rings with a target-like appearance, and small sporodochia that appeared on lower and upper surfaces (Figure 1f,n,o).
Figure 1. Symptoms on different hosts caused by *Paramyrothecium* (left) and sporodochia on the host surface (right); (a) *Solanum virginianum*; (b) *Lablab purpureus*; (c) *Psophocarpus* sp.; (d, i) *Vigna* sp.; (e) *Coccinia grandis*; (f) *Commelina benghalensis*; (g) *Tectona grandis*; (h) *Vigna mungo*; (j) *Vigna unguiculata*; (k) *Oroxylum indicum*; (l) *Spilanthes* sp.; (m) *Centrosema* sp.; (n) *Aristolochia* sp.; (o) *Coffea arabica*. Scale bars: (c, g, j) = 1 mm; (b, d–f, l) = 2 mm; (a, i, k) = 4 mm; (h, m) = 5 mm; (o) = 6 mm; (n) = 1 cm.

2.2. Culture Morphology

Diverse culture characters were observed on PDA at room temperature (25–30 °C) (Figure 2). Eleven isolates of *Paramyrothecium* sp. (SDBR-CMU374, SDBR-CMU375, SDBR-CMU376, SDBR-CMU377, SDBR-CMU378, SDBR-CMU379, SDBR-CMU380, SDBR-CMU382, SDBR-CMU387, SDBR-CMU388, and SDBR-CMU389) (Figure 2a–j, l) formed whitish colonies with entire to slightly undulated margins, radial or in concentric rings with sporodochia, covered with slimy olivaceous green to black conidial masses, while the other four isolates (SDBR-CMU383, SDBR-CMU384, SDBR-CMU385, and SDBR-CMU386) (Figure 2m–p) formed abundant white aerial mycelium with sporodochia forming on the stroma and surface of the medium, covered by slimy olivaceous green to black conidial masses. Isolate SDBR-CMU381 (Figure 2k) produced exudates with brown pigment into the medium.
2.3. Phylogenetic Analysis

The phylogenetic tree topologies of the ML and BI analyses for concatenated ITS, cmdA, rpb2, and tub2 were similar. Hence, a phylogenetic tree from ML analyses is used to represent the results of both ML and BI analyses. The dataset comprised 53 taxa with 1760 characters (ITS: 1–542; cmdA: 543–824; rpb2: 825–1548; tub2: 1549–1760), including gaps. The GTR+G+I model was the best-fit model for all loci. The best scoring likelihood tree was selected on the basis of the ML analysis, with a final ML optimization likelihood value of $-8176.4871$, as shown in Figure 3. Sixteen new isolates were clustered into four distinct clades in Paramyrothecium (see the notes).

2.4. Taxonomy

Isolates from symptomatic living leaves of different hosts were recognized under Paramyrothecium based on taxonomy (Table 1) and multi-gene phylogeny (Figure 3). The morphologies of the Paramyrothecium species are described herein.
Figure 3. Phylogram generated from maximum likelihood analysis based on combined ITS, cmdA, rpb2, and tub2 sequenced data. Fifty-three strains are included in the combined sequence analyses, which comprise 1760 characters with gaps. Single gene analyses were also performed, and topology and clade stability were compared from combined gene analyses. Striaticonidium cinctum (CBS 932.69), S. humicola (CBS 388.97), and S. synnematum (CBS 479.85) are used as the outgroup taxa. The best scoring RAxML tree with a final likelihood value of $-8176.4871$ is presented. The matrix had 524 distinct alignment patterns. Estimated base frequencies were as follows; A = 0.2266, C = 0.2915, G = 0.2681, T = 0.2138; substitution rates AC = 1.1215, AG = 5.1556, AT = 1.0792, CG = 1.2292, CT = 11.1203, GT = 1.0000; gamma distribution shape parameter $\alpha$ = 0.3855. The bootstrap support ($\geq 50\%$) of ML and the posterior probability values ($\geq 0.9$) of BI analyses are indicated above or below the respective branches. The fungal isolates from this study are indicated in red. The type species are indicated in bold.
Table 1. Synopsis of *Paramyrothecium* type species.

| Species                  | Host                  | Location | Conidiophores (µm) | Conidiogenous Cells (µm) | Conidia (µm) | Setae (µm) | References |
|--------------------------|-----------------------|----------|--------------------|--------------------------|--------------|------------|------------|
| *Paramyrothecium acadiense* | Tussilago farfara     | Canada   | 9–14 × 2–2.5       | –                        | 0–1-septate, 5.5–16.5 × 1.5–2.5 | –          | [20]       |
| *P. breviseta*           | unknown               | India    | 6–9 × 2–4          | 6–11 × 1–2               | aseptate, 4–5 × 1–2 | 1–3-septate, 25–40 × 2–3 | [4]        |
| *P. cupuliforme*         | Soil                  | Namibia  | 15–25 × 2–4        | 4–11 × 1–3               | aseptate, 6–8 × 1–2 | 1–3-septate, 45–90 × 2–3 | [4]        |
| *P. eichhorniae*         | Eichhornia crassipes  | Thailand | 15–40 × 2–3        | (8--)11–17(–20) × 2–3    | aseptate, 5–6.5 × 1.5–2.5 | 1–3-septate, 40–120 × 2–3 | [19]       |
| *P. foeniculicola*       | Foeniculum vulgare    | Netherlands | 7–17 × 2–3      | 6–16 × 1–2               | aseptate, 5–7 × 1–2 | –          |           |
| *P. foliicola*           | Decaying leaf         | Brazil   | 15–25 × 2–3        | 8–14 × 1–2               | aseptate, 5–6 × 1–2 | 1–3-septate, 60–100 × 2–3 | [4]        |
| *P. guiyangense*         | Soil                  | China    | 10–60 × 1–3        | 8–18 × 1.6–2.7           | aseptate, 6.6–9.0 × 2–3 | 1–3-septate, 60–120 × 1–3 | [21]       |
| *P. humicola*            | Soil                  | USA      | 12–22 × 2–3        | 8–13 × 1–3               | aseptate, 6–7 × 1–2 | 1–2-septate, 55–65 × 2–3 | [4]        |
| *P. lathyri*             | Lathyrus tuberosus    | Russia   | 5–10 × 2–3.5       | 5–10 × 2–3               | aseptate, (8–)9 × 2(–2.5) | 3–10-septate, up to 300 × 3–4 | [22]       |
| *P. nigrum*              | Soil                  | Spain    | 25–45 × 2–4        | 8–13 × 1–2               | aseptate, 5–6 × 1–2 | 1–3-septate, 60–100 × 2–3 | [4]        |
| *P. parvum*              | Viola sp.             | UK       | 12–26 × 2–4        | 7–23 × 1–2               | aseptate, 4–5 × 1–2 | –          |           |
| *P. pituitipietianum*     | Grielum humifusum     | South Africa | 20–35 × 3–4      | 20–35 × 3–4 | aseptate, (7–)9–10(–12) × 2(–2.5) | 7–10-septate, 100–300 × 4–5 | [23]       |
| *P. roditum*             | Gardenia sp.          | Italy    | 15–40 × 2–4        | 7–33 × 2–3               | aseptate, (5–)6.5–7.5×(–8) × 2 | 1–3(–4)-septate, 60–100 × 2–6 | [4]        |
| *P. salvadorae*          | Salvadora persica     | Namibia  | 20–40 × 3–4        | 8–15 × 2–2.5             | aseptate, (8–)10–12(–13) × 2–2.5 | 5–10- septate, 100–200 × 2.5–3 | [24]       |
| *P. sinense*             | Rhizosphere soils of Poa sp. | China | 20–30 × 2–3       | 7–16 × 1–3               | aseptate, 6–7 × 2–3 | 1–3-septate, 45–90 × 1–3 | [25]       |
| *P. tellicola*            | Soil                  | Turkey   | 15–30 × 2–4        | 7–17 × 1–3               | aseptate, (7–)7.5–8.5 × 9 × 1–3 | 1–3-septate, 45–80 × 2–3 µm | [4]        |
| *P. terrestris*           | Soil                  | Turkey   | 15–30 × 2–3        | 7–12 × 2–3               | aseptate, (7–)7.5–8.5 × 1–3 | 1–3-septate, 35–70 × 2–3 | [4]        |
| *P. verruridum*          | Soil                  | China    | 20–40 × 1.5–2.5    | 12–20 × 1.7–2.7          | aseptate, 6.8–7.8 × 2–2.7 | 1–3-septate, 40–120 × 2–3 | [21]       |
| *P. vignicola*           | Vigna sp.             | Thailand | 40–60 × 2–3        | 11–16 × 1–3               | aseptate, 5–7 × 1–3 µm | 3–8-septate, 80–155 × 2–3 | This study |
| *P. viridisporum*        | Soil                  | Turkey   | 15–35 × 2–3        | 6–12 × 3–5               | aseptate, 3–5 × 2 µm | 1–3-septate, 60–140 × 2–3 | [4]        |
2.4. Taxonomy

Isolates from symptomatic living leaves of different hosts were recognized under the genus *Paramyrothecium* (Table 1).

**Table 1.** Conidiomata of *Paramyrothecium* spp. arising from different hosts.

| Host        | Conidiomata Characteristics |
|-------------|----------------------------|
| Foeniculum  | Covered by slimy olivaceous green to black conidial masses. |
| Tussilago   | Covered by a green mucoid layer. |
| Soil        | Septate, smooth, 40–60 × 2–3 µm. |

**Figure 4.** *Paramyrothecium vignicola* (CRC4-H, holotype); (a) leaf spot of *Vigna* sp.; (b) sporodochia on leaf; (c) sporodochial conidiomata on PDA; (d,e) conidiophores and conidiogenous cells; (f,g) conidiogenous cells; (h) setae; (i) conidia. Scale bars: (b,e) = 1 mm; (d–h) = 10 µm; (i) = 5 µm.

Mycobank: MB 843763.

Etymology: Name reflects the host genus *Vigna*, from which the species was collected.

Holotype: SDBR–CMU376.

Description: Sexual morph: unknown. Asexual morph: *Conidiomata* sporodochial, stromatic, superficial, cupulate, scattered or gregarious, oval or irregular in outline, (60–90–300(–385) µm diam, (70–)140–180(–200) µm deep, with a white to creamy setose fringe surrounding an olivaceous green agglutinated slimy mass of conidia. *Stroma* poorly developed, hyaline. *Setae* arising from the stroma thin-walled, hyaline, 3–8-septate, straight becoming sinuous above the apical septum, 80–155 µm long, 2–3 µm wide, tapering to an acutely rounded apex. *Conidiophores* arising from the basal stroma, consisting of a stipe and a penicillately branched conidiogenous apparatus; stipes unbranched, hyaline sometimes covered by a green mucoid layer, septate, smooth, 40–60 × 2–3 µm; primary branches aseptate, unbranched, smooth, 10–26 × 2–3 µm ($\overline{x} = 18 \pm 3$ µm, n = 20); secondary branches aseptate, unbranched, smooth, 10–17 × 2–3 µm ($\overline{x} = 13 \pm 3$ µm, n = 20); terminating in a whorl of 3–6 conidiogenous cells; conidiogenous cells phialidic, cylindrical to subcylindrical, hyaline, smooth, straight to slightly curved, 11–16 × 1–3 µm ($\overline{x} = 13 \pm 2$ µm, n = 20), with conspicuous collarettes and periclinal thickenings. *Conidia* aseptate, hyaline, smooth, cylindrical to ellipsoidal, 5–7 × 1–3 µm ($\overline{x} = 6 \pm 2$ µm, n = 20), rounded at both ends.

Culture characteristics: Colonies on PDA, dense, circular, flattened, slightly raised, floccose, white aerial mycelium, radiating with concentric ring of sporodochia forming, covered by slimy olivaceous green to black conidial masses.

Material examined: Thailand, Mae Hong Son Province, on living leaf of *Vigna* sp. (*Fabaceae*), 11 September 2019, N. Tamakaew, CRC4-H (holotype); ex-type living culture SDBR-CMU376; *ibid.*, on living leaf of *Solanum virginianum* (*Solanaceae*), 11 September 2019,
N. Tamakaew, CRC1-H, living culture SDBR-CMU389; ibid., on living leaf of Lablab purpureus (Fabaceae), 11 September 2019, N. Tamakaew, CRC2-H, living culture SDBR-CMU374; ibid., on living leaf of Coccinia grandis (Cucurbitaceae), CRC6-H, living culture SDBR-CMU377; Chiang Mai province, on living leaf of Commelina benghalensis (Commelinaceae), 20 November 2019, P. Withee, CRC14-H, living culture SDBR-CMU381; ibid., on living leaf of Vigna mungo (Fabaceae), 5 December 2019, N. Tamakaew, CRC144-H, living culture SDBR-CMU384; ibid., on living leaf of Vigna sp. (Fabaceae), CRC145-H, living culture SDBR-CMU385; ibid., on living leaf of Vigna unguiculata (Fabaceae), 10 February 2020, P. Withee, CRC146-H, living culture SDBR-CMU386.

Notes: Based on ITS, cmdA, rpb2 and tub2 phylogeny (Figure 3) and cmdA and tub2 (data not shown), Paramyrothecium folicola formed two distinct clades. The clade with Paramyrothecium folicola type (CBS 113121) was treated as the Paramyrothecium sensu stricto. Eight of the new strains clustered with eight previously described Paramyrothecium strains (as P. folicola) and formed a well-supported clade (100% BS/1.00 PP) (Paramyrothecium sensu lato) closely related to P. eichhorniae and P. folicola (Figure 3). Based on morphology and phylogeny, we introduce a new species to accommodate taxa in P. folicola sensu lato. Paramyrothecium vignicola differs from P. eichhorniae and P. folicola with longer setae (up to 155 μm vs. up to 120 μm and up to 100 μm). The conidia of P. vignicola (5–7 × 1–3 μm) are slightly larger than those of P. eichhorniae (5–6.5 × 1.5–2.5 μm) [20] and P. folicola (5–6 × 1–2 μm) [4]. Paramyrothecium vignicola differs from other Paramyrothecium species by its 3–8-septate, thin-walled setae surrounding the sporodochia. In BLAST searches of NCBI GenBank, the closest matches of the sequences are Paramyrothecium: P. folicola (CBS 11321) with 98.98% similarity in ITS sequence, 93.89% similarity in cmdA. P. vignicola, 96.32% in tub2 with P. folicola (CBS 11321). Based on phylogenetic evidence and morphological differences, P. vignicola is a new species.

Paramyrothecium breviseta L. Lombard & Crous, in Lombard et al., Persoonia 36: 207 (2016) (Figure 5).

Figure 5. Paramyrothecium breviseta (CRC13-H); (a,b) leaf spot of Coffea arabica; (c) sporodochia on leaf; (d) conidiophores and conidiogenous cells; (e–g) conidiogenous cells; (h) setae; (i) conidia. Scale bars: (d) = 20 μm; (e–h) = 10 μm; (i) = 5 μm.
Description: Sexual morph: unknown. Asexual morph: Conidiomata sporodochial, stromatic, cupulate, superficial, scattered or rarely gregarious, oval or irregular in outline, 135–790 µm diam, 9–15 µm deep, with a white setose fringe surrounding an olivaceous green to black agglutinated slimy mass of conidia. Setae arising from the stroma thin-walled, hyaline, 1–5-septate, straight to flexuous, 25–120 µm long, 2–3 µm wide, tapering to an acutely rounded apex. Conidiophores arising from the basal stroma, consisting of a stipe and a penicillately branched conidiogenous apparatus; stipes unbranched, hyaline, septate, smooth, 6–9 × 2–4 µm; primary branches aseptate, unbranched, smooth, 12–24 × 3–4 µm (\(\overline{x} = 18 \times 3 \, \mu m, n = 20\)); secondary branches aseptate, unbranched, smooth, 10–17 × 2–4 µm (\(\overline{x} = 12 \times 3 \, \mu m, n = 20\)); terminating in a whorl of 3–6 conidiogenous cells; conidiogenous cells phialidic, cylindrical to subcylindrical, hyaline, smooth, straight to slightly curved, 5–7 × 1–2 µm (\(\overline{x} = 6 \times 2 \, \mu m, n = 20\)), rounded at both ends. Conidia aseptate, hyaline, smooth, cylindrical to ellipsoidal, 5–7 × 1–2 µm (\(\overline{x} = 6 \times 2 \, \mu m, n = 20\)), rounded at both ends.

Culture characteristics: Colonies on PDA, dense, circular, flattened, slightly raised, floccose, white aerial mycelium, radiating with concentric ring of sporodochia forming, covered by slimy olivaceous green to black conidial masses.

Material examined: Thailand, Chiang Mai, on living leaf of Coffea arabica (Rubiaceae), 20 November 2019, R. Cheewangkoon and P. Withee, CRC13-H, living culture SDBR-CMU387; ibid., CRC12-H, living culture SDBR-CMU388.

Notes: Phylogenetically, SDBR-CMU387 and SDBR-CMU388 formed a well-supported clade closely related to Paramyrothecium breviseta L. Lombard & Crous (Figure 2). Paramyrothecium breviseta was collected on an unknown substrate in India [4] and in this study, we collected P. breviseta from Coffea arabica (Rubiaceae) in Chiang Mai Province. The morphology of the fresh specimen is similar to that described by Lombard et al. [4], but the conidia (5–7 × 1–2 µm vs. 4–5 × 1–2 µm) and setae (25–120 × 2–3 vs. 25–40 × 2–3 µm) are longer. However, this is the first host report of leaf spot causing P. breviseta on C. arabica in Thailand.

Paramyrothecium eichhorniae J. Unartngam, A. Unartngam & U. Pinruan, in Pinruan et al., Mycobiology 50: 17 (2022) (Figure 6).

Description: Sexual morph: unknown. Asexual morph: Conidiomata sporodochial, stromatic, superficial, cupulate, scattered or gregarious, oval or irregular in outline, (60–)70–250(–500) µm diam, (60–)70–270(–370) µm deep, with a white setose fringe surrounding an olivaceous green to dark green slimy mass of conidia. Setae arising from the stroma thin-walled, hyaline, 1–5-septate, straight to flexuous, 60–120 µm long, 2–3 µm wide, tapering to an acutely rounded apex. Conidiophores arising from the basal stroma, consisting of a stipe and a penicillately branched conidiogenous apparatus; stipes unbranched, hyaline, septate, smooth, 15–40 × 2–3 µm; primary branches aseptate, unbranched, smooth, 10–17 × 2–3 µm (\(\overline{x} = 12 \times 3 \, \mu m, n = 20\)); secondary branches aseptate, unbranched, smooth, 7–14 × 2–3 µm (\(\overline{x} = 10 \times 3 \, \mu m, n = 20\)); terminating in a whorl of 3–6 conidiogenous cells; conidiogenous cells phialidic, cylindrical to subcylindrical, hyaline, smooth, straight to slightly curved, 11–17 × 2–3 µm (\(\overline{x} = 14 \times 2 \, \mu m, n = 20\)), with conspicuous collarettes and periclinal thickenings. Conidia aseptate, hyaline, smooth, cylindrical to ellipsoidal, 5–7 × 1–2 µm (\(\overline{x} = 6 \times 2 \, \mu m, n = 20\)), rounded at both ends.

Culture characteristics: Colonies on PDA, entire to slightly undulated margins, with sporodochia forming on the surface of the medium, covered by slimy olivaceous green to black conidial masses.
**Figure 6.** *Paramyrothecium eichhorniae* (CRC143); (a) leaf spot of *Aristolochia* sp.; (b) sporodochia on leaf; (c) sporodochial conidiomata on PDA; (d) sporodochia; (e,f) conidiogenous cells; (g) setae; (h) conidia. Scale bars: (b,c) = 1 mm; (d,g) = 20 µm; (e,f) = 10 µm; (h) = 5 µm.

Material examined: Thailand, Mae Hong Son Province, on living leaf of *Psophocarpus* sp. (*Fabaceae*), 11 September 2019, N. Tamakaew, CRC3-H, living culture SDBR-CMU375; *ibid.*, on living leaf of *Oroxylum indicum* (*Bignoniaceae*), 11 September 2019, N. Tamakaew, CRC8-H, living culture SDBR-CMU378; *ibid.*, on living leaf of *Spilanthes* sp. (*Asteraceae*), 11 September 2019, N. Tamakaew, CRC148-H, living culture SDBR-CMU379; *ibid.*, on living leaf of *Centrosema* sp. (*Fabaceae*), 11 September 2019, N. Tamakaew, CRC11-H, living culture SDBR-CMU380; Chiang Mai province, on living leaf of *Aristolochia* sp. (*Aristolochiaceae*), January 2020, P. Suttiprapan, CRC143-H, living culture SDBR-CMU383.

Note: Based on multigene phylogeny, five isolates in this study clustered with *Paramyrothecium eichhorniae*, which was associated with water hyacinth (*Eichhornia crassipes*) and recently described from Thailand [10]. Morphologically, the conidiogenous cells of our collections are similar to those of the holotype of *P. eichhorniae*. However, the conidia of *P. eichhorniae* in this study are thinner than reported by Pinruan et al. [19] (5–7 × 1–2 µm vs. 5–6.5 × 1.5–2.5 µm) and have more septa in setae than the holotype (1–5 vs. 1–3 septate). This is the first report of *P. eichhorniae* on *Psophocarpus* sp., *Centrosema* sp., and *Aristolochia* sp. from Thailand.

*Paramyrothecium folicola* L. Lombard & Crous, in Lombard et al., Persoonia 36: 209 (2016) (Figure 7).

Description: Sexual morph: unknown. Asexual morph: *Conidiomata* sporodochial, stromatic, superficial, cupulate, scattered or gregarious, oval or irregular in outline, (60–)100–170(–245) µm diam, (70–)140–165(–200) µm deep, with a white to creamy setose fringe surrounding an olivaceous green agglutinated slimy mass of conidia. *Stroma* poorly developed, hyaline. *Setae* arising from the stroma thin-walled, hyaline, 1–4(–8)-septate, straight becoming sinuous above the apical septum, 35–175 µm long, 2–3 µm wide, tapering to an acutely rounded apex. *Conidiophores* arising from the basal stroma, consisting of a stipe.
and a penicillately branched conidiogenous apparatus; stipes unbranched, hyaline sometimes covered by a green mucoid layer, septate, smooth, 20–75 × 2–4 μm; primary branches aseptate, unbranched, smooth, (10–)17–21(–26) × 2–3(–4) μm (X = 15 × 3 μm, n = 20); secondary branches aseptate, unbranched, smooth, (7–)9–17(–19) × 2–3(–4) μm (X = 14 × 3 μm, n = 20); terminating in a whorl of 3–6 conidiogenous cells; conidiogenous cells phialidic, cylindrical to subcylindrical, hyaline, smooth, straight to slightly curved, 10–17 × 1–3 μm (X = 13 × 2 μm, n = 20), with conspicuous collarettes and periclinal thickenings. Conidia aseptate, hyaline, smooth, cylindrical to ellipsoidal, 5–8 × 1–3 μm (X = 7 × 2 μm, n = 20), rounded at both ends.

Figure 7. Paramyrothecium foliicola (CRC15); (a) sporodochia on leaves of Tectona grandis (b) sporodochial conidiomata on PDA; (c,d) sporodochia (e,f) conidiogenous cells; (g) setae; (h) conidia. Scale bars: (a) = 500 μm; (b) = 1 mm; (c) = 30 μm; (d,g) = 20 μm; (h) = 5 μm.

Culture characteristics: Colonies on PDA, abundant white aerial mycelium with sporodochia forming on the aerial mycelium and surface of the medium, covered by slimy olivaceous green to black conidial masses.

Materials examined: Thailand, Chiang Mai, on living leaf of Tectona grandis (Lamiaceae), 20 November 2019, P. Withee, CRC15-H, living culture SDBR-CMU382.

Notes: Based on our phylogenetic analysis (Figure 3), SDBR-CMU382 isolates were clustered with Paramyrothecium foliicola. The morphology of our collection (CRC15-H) is similar to that of P. foliicola described by Lombard et al. [4]. However, our collection has longer conidiophores (20–75 × 2–4 vs. 15–25 × 2–3 μm) and more septa in setae (1–4(–8) vs. 1–3 septate), conidiogenous cells (10–17 × 1–3 vs. 8–14 × 1–2 μm) and conidia (5–8 × 1–3 vs. 5–6 × 1–2 μm). This may be due to distribution, environment, and morphological variability within the species. This is the first report of P. foliicola from Tectona grandis in Thailand.
2.5. Pathogenicity Test and Cross Pathogenicity

Koch’s postulates confirmed that all the fungal isolates were able to cause disease in unwounded leaves of Commelina benghalensis and Glycine max (Figure 8b,c). The SDBR-CMU383 isolate infected all inoculated plants and was highly aggressive on most, except for C. benghalensis. No infection was observed in the unwounded inoculation of Coffea arabica and Dieffenbachia seguine (Figure 8a,d). Leaves receiving sterilized distilled water remained healthy. The fungi were re-isolated from the diseased leaf tissues in each experiment, and each isolated fungus was identical to the inoculated fungus. Further, Koch’s postulates confirmed that all isolates of Paramyrothecium vignicola, P. breviseta, P. eichhorniae, and P. foliicola were pathogenic to their original host plants. Cross pathogenicity tests showed that all isolates infected inoculated (wounded) C. arabica, C. benghalensis, G. max, and D. seguine leaves (Table 2). The symptoms showed light to dark brown and irregular to round lesions, which had scattered olive-coloured sporodochia and dark exudates of spore masses (Figure 8).

Figure 8. Pathogenicity test (a,b) and cross pathogenicity (c,d); Control (left): (a) Paramyrothecium breviseta on Coffea arabica; (b) P. vignicola on Commelina benghalensis; (c) P. vignicola on Glycine max; (d) P. vignicola on Dieffenbachia seguine; (w) wound and (uw) unwounded. Scale bars: (a–c) = 1 cm; (d) = 6 cm.
Table 2. Pathogenicity test and cross pathogenicity of *Paramyrothecium* species on original hosts and other plant species.

| Species | Original Host | Isolates | **Plant Hosts** |
|---------|---------------|----------|----------------|
|         |               |          | Coffea arabica | Commelina benghalensis | Glycine max | Dieffenbachia seguine |
|         |               |          | w | uw | w | uw | w | uw | w | uw |
| *P. vignicola* | Vigna sp. | SDBR-CMU376 | + | - | + | - | + | + | + | + | + | + | + | + | + | - |
|           | Lablab purpureus | SDBR-CMU374 | + | - | + | + | + | + | + | + | - |
|           | Commelina benghalensis | SDBR-CMU377 | + | - | + | - | + | + | + | + | - |
|           | Vigna mungo | SDBR-CMU384 | + | - | + | + | + | - |
|           | Vigna sp. | SDBR-CMU385 | + | - | + | + | + | + | + | - |
|           | Vigna unguiculata | SDBR-CMU386 | + | - | + | + | + | - |
|           | Solanum virginianum | SDBR-CMU389 | + | - | + | - |
| *P. brevista* | Coffea arabica | SDBR-CMU387 | + | - | + | - |
|          | Coffea arabica | SDBR-CMU388 | + | - | + | + | + | + | + | + | - |
| *P. eichhorniae* | Psophocarpus sp. | SDBR-CMU375 | + | - | + | + | + | - |
|           | Oroxylum indicum | SDBR-CMU378 | + | - | + | + | + | - |
|           | Spilanthes sp. | SDBR-CMU379 | + | - | + | + | + | - |
|           | Centrosema sp. | SDBR-CMU380 | + | - | + | + | + | - |
|           | Aristolochia sp. | SDBR-CMU383 | + | - | - | + | + | - |
| *P. foliicola* | Tectona grandis | SDBR-CMU382 | + | - | + | + | + | - |

Note: (-) No symptoms (+) Symptoms (w) wound and (uw) unwound.; Superscript “T” indicates type species.

3. Discussion

The new species *Paramyrothecium vignicola* was described using morphology and multi-gene phylogeny and the host range included *Solanum virginianum* (*Solanaceae*), *Lablab purpureus* (*Fabaceae*), *Coccinia grandis* (*Cucurbitaceae*), *Commelina benghalensis* (*Commelinaceae*), *Vigna* sp., *V. mungo*, and *V. unguiculata* (*Fabaceae*). Multi-gene phylogeny using ITS, *cmdA*, *rpb2*, and *tub2* sequence data clearly identified *P. eichhorniae*, *P. vignicola*, *P. breviseta*, and *P. foliicola* as distinct species within *Paramyrothecium*. Further, multi-gene phylogeny precisely demonstrated the species delineation of *Paramyrothecium*.

The pathogenicity assays showed that *P. vignicola*, *P. breviseta*, *P. eichhorniae*, and *P. foliicola* isolated from different hosts from different locations in northern Thailand can all cause leaf spot disease on different host families, including *Rubiaceae*, *Fabaceae*, *Commelinaceae*, and *Araceae*. However, the disease severity was related to the plant species and inoculation methods, where *Paramyrothecium* spp. could not cause disease in *Coffea arabica* and *Dieffenbachia seguine* without wounding. Wounding involves the breakage of the plant’s first barrier of defense; cuticle and epidermal cells. The tissue then becomes more susceptible to the pathogens. Some species cannot infect non-wounded leaves, hence they are weakly aggressive on these hosts [15]. On the other hand, *Commelina benghalensis* and *Glycine max* were susceptible to all isolates. These results are similar to those of Rennberger and Keinath [11] and Aumentado and Balendres [12], in which *Paramyrothecium* species were able to infect original and non-original hosts within the same family (host shift ability) and different families (host jump ability).

For species diversity and distribution, more gene studies and more reference sequences are needed to resolve the species boundaries of *Paramyrothecium*. Field inspections are needed to confirm the importance of this pathogen and prove that diseases associated with *Paramyrothecium* species are threats to economic crops in Thailand. The information on the spread of related species to new areas is necessary as climate change may enable saprotrophic fungi to switch their nutritional mode across a wider host range, even if an area is predicted to be at risk from an introduced pathogen. It may be the case that few
of the susceptible host species are present in this predicted area [26], so for the risk to be realized, climate change should also favor the migration of susceptible species or increase the susceptibility of the resident hosts.

*Paramyrothecium* leaf spot occurs in commercially important plants (*Coffea arabica, Tectona grandis, Vigna mungo,* and *V. unguiculata*) as well as on non-commercial plants (*Aristolochia* sp., *Centrosema* sp., *Coccinia grandis, Commelina benghalensis, Lablab purpureus, Oroxylum indicum, Psophocarpus* sp., *Solanum virginianum, Spilanthes* sp., and *Vigna* sp.). In cross pathogenicity assays, all the isolates from host plants could induce the disease on non-original hosts. *Paramyrothecium* species can stay in non-commercial plants, and they can infect commercially important crops. Hence, *Paramyrothecium* leaf spot disease has the potential to be an emerging fungal disease in Thailand. Thus, more research on *Paramyrothecium* is required for epidemiology studies and management strategies in agriculture, horticulture, and plantation forestry.

4. Materials and Methods

4.1. Sample Collection

Symptomatic plant leaves were collected from fields or forests in different locations in northern Thailand. The name of the host, location, and collection dates were recorded. Specimens were taken to the lab, and infected leaves were examined directly using the stereo microscope (Zeiss Stemi 305) to observe the fungal structures (sporodochia). Symptomatic leaves without fungal structures were also incubated in moist chambers (Petri dishes containing moist filter paper). Leaves were inspected daily for *Paramyrothecium*-like fungi.

4.2. Fungal Isolation and Taxonomic Description

Fungal structures on leaf samples were mounted in lactic acid and photographed under a light microscope (Axiowision Zeiss Scope-A1). Measurements were made with the Tarosoft (R) Image Frame Work program (Tarosoft, Bangkok, Thailand). The fungi were isolated using the single spore isolation technique [27]. Cultures were plated onto fresh PDA and incubated at 25–30 °C in daylight to promote sporulation. Cultural characteristics were observed after 14 days. The specimens were deposited in the fungal collection library at the Department of Entomology and Plant Pathology (CRC), Faculty of Agriculture, Chiang Mai University, Chiang Mai, Thailand. Pure fungal isolates were deposited in the Culture Collection of the Sustainable Development of Biological Resources Laboratory (SDBR), Faculty of Science, Chiang Mai University, Chiang Mai, Thailand.

4.3. DNA Extraction, Amplification, and Analyses

Fungal mycelia were grown on PDA at 25–30°C for 7 days and DNA was extracted by using the DNA Extraction Mini Kit (FAVORGEN, Ping-Tung, Taiwan) following the manufacturer’s instructions. DNA amplifications were performed by polymerase chain reaction (PCR). The relevant primer pairs used in this study are listed in Table 3.

| Gene Regions | Primers | Sequence (5′→3′) | Length (bp) | References |
|--------------|---------|-----------------|-------------|------------|
| ITS          | ITS5    | GGA AGT AAA AGT CGT AAC AAG G | ca. 600 | [28] |
|              | ITS4    | TCC GCT TAT TGA TAT GC | | |
|              | CAL–228F | GAG TTC AAG GAG GCC TTC TCC C | | |
|              | CAL–737R | CAT CTT TCT GCC CAT GG | CAL–228F–CAL–737R: 470–570 | |
|              | CAL2Rd  | TGR TCN GCC TCD CGG ATC ATC TC | CAL–228F–CAL2Rd: 680–745 | [29,30] |
| cmdA         | CAL–228F | GAG TTC AAG GAG GCC TTC TCC C | | |
|              | CAL–737R | CAT CTT TCT GCC CAT GG | CAL–228F–CAL–737R: 470–570 | |
|              | CAL2Rd  | TGR TCN GCC TCD CGG ATC ATC TC | CAL–228F–CAL2Rd: 680–745 | [29,30] |
| rpb2         | RPB2–5F | GAY GAY MGW GAT CAY TTY GG | ca. 1000 | [31] |
|              | RPB2–7cR | GAY GAY MGW GAT CAY TTY GG | | |
| tub2         | Bt2a    | GCC AAT CAA ATC GGT TCT GTC | ca. 320 | [32] |
|              | Bt2b    | ACC CTC AGT GTA GTG ACC CTT GGC | | |

Table 3. Gene regions and primer sequences used in this study.
The quality of PCR amplification was confirmed on 1% agarose gel electrophoresis and viewed under ultraviolet light, and the sizes of amplicons were determined against a HyperLadder™ I molecular marker (BIOLINE). Further purification of PCR products was performed using the PCR Clean-up Gel Extraction NucleoSpin® Gel and PCR Clean-Up Kit (Macherey-Nagel, Düren, Germany). The purified PCR fragments were sent to the 1st Base Company (Kembangan, Selangor, Malaysia). The obtained nucleotide sequences were deposited in GenBank.

Sequences were assembled using SeqMan 5.00 and the closely related taxa for newly generated sequences were selected from GenBank® based on BLAST searches of the NCBI nucleotide database (http://blast.ncbi.nlm.nih.gov/; accessed on 4 March 2022). The reference nucleotide sequences of representative genera in Stachybotriaceae are in Table 4. The individual gene sequences were initially aligned by MAFFT version 7 [33] (http://mafft.cbrc.jp/alignment/server/; accessed on 4 March 2022) and improved manually where necessary in BioEdit v.7.0.9.1 [34]. The final alignment of the combined multigene dataset was analyzed and inferred the phylogenetic trees based on maximum likelihood (ML) and Bayesian inference (BI) analyses. The ML analyses were carried out on RAxML-HPC2 on XSEDE (v. 8.2.8) [35,36] via the CIPRES Science Gateway platform [37]. Maximum likelihood bootstrap values (BS) equal or greater than 50% are defined above each node. The BI analyses were performed by MrBayes on XSEDE, MrBayes 3.2.6 [38] via the CIPRES Science Gateway. Bayesian posterior probabilities (PP) [39,40] were determined by Markov Chain Monte Carlo Sampling (BMCMC). Six simultaneous Markov chains were run from random trees for 2,000,000 generations, and trees were sampled every 100th generation. The run was stopped when the standard deviation of split frequencies was reached at less than 0.01. The first 20% of generated trees representing the burn-in phase of the analysis were discarded, and the remaining trees were used for calculating PP in the majority rule consensus tree. The Bayesian posterior probabilities (BYPP) equal to or greater than 0.9 are defined above the nodes. The phylogenetic tree was visualized in FigTree v.1.4.3 [41] and edited in Adobe Illustrator CC 2021 version 23.0.3.585 and Adobe Photoshop CS6 version 13.0. (Adobe Systems, New York, USA).

4.4. Pathogenicity Tests and Cross Pathogenicity

Koch’s postulates were used to confirm the pathogenicity of all the isolates on their original hosts. Cross pathogenicity of all the isolates was performed in healthy leaves of selected economically important plants in northern Thailand, including Coffea arabica (Rubiaceae) and Glycine max (Fabaceae) and widespread herbaceous plants including Commelina benghalensis (Commelinaceae) and Diefenbachia seguine (Araceae). Healthy leaves were surface disinfected with 70% ethanol, washed two times with sterile distilled water, and air-dried under laminar flow. Conidial suspensions (10⁶ conidia/mL) were prepared for all fungal isolates in sterile distilled water. The conidia (10 µL of spore suspension) were placed on the upper surface of the leaves. In addition, the leaves were also wounded before inoculation. The upper epidermis was wounded approximately 2 cm from the mid-vein by pricking with a sterile needle to about 1 mm depth. Three wounds were made for each leaf, vertically on each side of the mid-vein. Control leaves received drops of sterile distilled water. All inoculated leaves were placed in a moist chamber at 25–30 °C under daylight condition. After 7 days, symptoms were recorded, compared, and confirmed with the original morphology and molecular relationships.
Table 4. Taxa used in the phylogenetic analyses and their corresponding GenBank numbers.

| Species                        | Isolate No. | Substrate                                      | Location      | GenBank Accession Numbers |
|--------------------------------|-------------|------------------------------------------------|---------------|---------------------------|
|                                |             |                                                |               |                           |
| **Myrothecium inundatum**      | CBS 275.48  | On decaying pileus of *Russula nigricans*      | England       | KU846452 KU846435 – KU846533 |
| M. simplex                     | CBS 582.93  | On decaying agaric                             | Japan         | KU846456 KU846439 – KU846537 |
| **Paramyrothecium acadiense**  | CBS 123.96 ^T= DAOMC 221473 = UAMH 7653 | On leaves of *Tussilago farfara*                | Canada        | KU846288 – KU846350 KU846405 |
|                                | CBS 544.75  | unknown                                        | India         | KU846289 KU846262 KU846351 KU846406 |
| P. breviseta                   | DRL3        | On leaves of *Coffea canephora*                | China         | MT853067 MT897897 – MT897999|
|                                | DRL4        | On leaves of *C. canephora*                    | China         | MT853068 MT897898 – MT897900 |
|                                | SDBR-CMU387 | On living leaf of *C. arabica*                 | Thailand      | MZ373251 OM810407 ON033773 OM982450 |
| P. cupuliforme                 | CBS 127789  | On surface soil in desert                      | Namibia       | KU846291 KU846264 KU846353 KU846408 |
|                                | CBS 126167  | On surface soil in desert                      | Namibia       | KU846290 KU846263 KU846352 KU846407 |
| P. eichhorniae                 | TBRC 10637  | On leaf of *Eichhornia crassipes*              | Thailand      | MT973996 MT975319 MT977540 MT977540 |
|                                | KKFC 474    | On leaf of *E. crassipes*                      | Thailand      | MT973995 MT975318 MT977541 MT9775316 |
|                                | SDBR-CMU375 | On living leaf of *Psophocarpus sp.*           | Thailand      | MZ373241 OM810411 ON033781 ON033770 |
|                                | SDBR-CMU378 | On living leaf of unidentified plant           | Thailand      | MZ373246 OM810414 ON033782 ON033772 |
|                                | SDBR-CMU379 | On living leaf of unidentified plant           | Thailand      | MZ373247 OM810415 ON033783 ON033768 |
|                                | SDBR-CMU380 | On living leaf of *Centrosemia* sp.            | Thailand      | MZ373250 OM810416 ON033784 ON033771 |
|                                | SDBR-CMU383 | On living leaf of *Aristolochia* sp.           | Thailand      | MZ373255 OM810418 ON033785 ON033769 |
| P. foeniculica                 | CBS 331.51 ^T= IMI 140051 | On leaf sheath *Foeniculum vulgare*   | The Netherlands | KU846292 – KU846354 KU846409 |
| P. foliicola                   | CBS 113121 ^T= INIFAT CO2/104 ^T  | On rotten leaf of unknown host                | Brazil        | KU846294 KU846266 – KU846411 |
|                                | SDBR-CMU382 | On decaying leaf of *Tectona grandis*          | Thailand      | MZ373254 – ON033775 OM982452 |
Table 4. Cont.

| Species               | Isolate No.       | Substrate                  | Location          | GenBank Accession Numbers        |
|-----------------------|-------------------|----------------------------|-------------------|----------------------------------|
|                       |                   |                            |                   | ITS   | cmdA     | rpb2     | tub2   |
| **P. guiyangense**    | GUCC 201608501    | From soil                  | China             | KY126418 | KY196193 | –        | KY196201 |
|                       | HGUP 2016–8001    | From soil                  | China             | KY126417 | KY196192 | –        | KY196200 |
| **P. humicola**       | CBS 127295        | From tallgrass prairie soil | USA               | KU846295 | –        | KU846356 | KU846412 |
|                       | MU4               | On leaf of *Citrullus lanatus* | USA               | MN227389 | MN593629 | MN397959 | MN398054 |
| **P. nigrum**         | CBS 116537        | From soil                  | Spain             | KU846296 | KU846267 | KU846357 | KU846413 |
|                       | LC12188           | Rhizosphere soils of *Poa* sp. | China             | MK478871 | MK500252 | MK500261 | MK500269 |
| **P. parvum**         | CBS 257.35        | On *Viola* sp.             | UK                | KU846298 | –        | KU846359 | KU846415 |
|                       | CBS 142.42        | From dune sand             | France            | KU846297 | KU846268 | KU846358 | KU846414 |
| **P. pituitipetianum**| CPC38688          | On stems of *Grielum humifusum* | South Africa      | MW175358 | MW173100 | –        | MW173139 |
|                       | CBS 357.89        | On *Gardenia* sp.          | Italy             | KU846300 | KU846270 | KU846361 | KU846417 |
|                       | CBS 372.50        | On twig of *Coffee* sp.    | Colombia          | KU846301 | KU846269 | KU846360 | KU846416 |
| **P. roridum**        | CGMCC3.19212      | Rhizosphere soils of *Poa* sp. | China             | MH793296 | MH885437 | MH818824 | MH793313 |
|                       | LC12137           | Rhizosphere soils of *Poa* sp. | China             | MH793295 | MH885436 | MH818822 | MH793312 |
| **P. tellicola**      | CBS 478.91        | From soil                  | Turkey            | KU846302 | KU846272 | KU846363 | KU846419 |
| **P. terrestris**     | CBS 564.86        | From soil under *Lycopersicon esculentum* | Turkey            | KU846303 | KU846273 | KU846364 | KU846420 |
|                       | CBS 566.86        | From soil beneath *Helianthus annuus* | Turkey            | KU846305 | KU846275 | KU846366 | KU846422 |
| **P. verruridum**     | HGUP 2016–8006    | From soil                  | China             | KY126422 | KY196197 | –        | KY196205 |
### Table 4. Cont.

| Species                  | Isolate No. | Substrate                | Location       | ITS          | cmdB         | rpb2         | tub2         |
|--------------------------|-------------|--------------------------|----------------|--------------|--------------|--------------|--------------|
| *P. vignicola*           | SDBR-CMU389 | On living leaf of *Solanum virginianum* | Thailand       | MZ373239     | OM810409     | ON033776     | ON009013     |
|                          | SDBR-CMU374 | On living leaf of *Lablab purpureus* | Thailand       | MZ373240     | OM810410     | ON033777     | ON009014     |
|                          | SDBR-CMU376 | On living leaf of *Vigna* sp. | Thailand       | MZ373242     | OM810412     | ON033778     | ON009015     |
|                          | SDBR-CMU377 | On living leaf of *Coccinia grandis* | Thailand       | MZ373244     | OM810413     | ON033779     | ON009016     |
|                          | SDBR-CMU381 | On living leaf of *Commelina benghalensis* | Thailand       | MZ373253     | OM810417     | ON033780     | ON009017     |
|                          | SDBR-CMU384 | On living leaf of *Vigna mungo* | Thailand       | MZ373256     | OM810419     | ON033786     | –             |
|                          | SDBR-CMU385 | On living leaf of *Vigna* sp. | Thailand       | MZ373257     | OM810420     | ON033787     | ON009018     |
|                          | SDBR-CMU386 | On living leaf of *V. unguiculata* | Thailand       | MZ373258     | OM810421     | ON033788     | ON009019     |
| *P. viridisporum*        | CBS 873.85  | From soil                | Turkey         | KU846308     | KU846278     | KU846369     | KU846425     |
|                          | CBS 125835  | Rhizosphere soils of bunchgrass | USA           | KU846310     | KU846280     | KU846371     | KU846427     |
| *Striaticonidium cinctum*| CBS 932.69  | From agricultural soil    | The Netherlands | KU847239     | KU847216     | KU847290     | KU847329     |
| *S. humicola*            | CBS 388.97  | From soil in tropical forest | Papua New Guinea | KU847241     | KU847217     | KU847291     | KU847331     |
| *S. synnematum*          | CBS 479.85  | From leaf of unknown palm | Japan          | KU847242     | KU847218     | KU847292     | KU847332     |
| *Tangerinosporium thalictricola* | CBS 317.61 = IMI 034815 | On *Thalictrum flavum* | UK             | KU847243     | KU847219     | –             | KU847333     |
| *Xenomyrothecium tongaense* | CBS 598.80  | On dead thallus of *Halimeda* sp. | Tonga         | KU847246     | KU847221     | KU847295     | KU847336     |

Note: CBS: Culture collection of the Centraalbureau voor Schimmelcultures, Fungal Biodiversity Centre, Utrecht, The Netherlands; CGMCC: China General Microbiological Culture Collection Center; CPC: Collection of P.W. Crous; DAOMC: The Canadian Collection of Fungal Cultures; GUCC: Guizhou University Culture Collection, Guiyang, China; HGUP: Herbarium of Guizhou University, Plant Pathology, China; IMI: International Mycological Institute, CABI-Bioscience, Egham, Bakeham Lane; INIFAT: INIFAT Fungus Collection, Ministerio de Agricultura Habana; KKFC: Kasetsart.Kamphaengsaen Fungal Collection, Thailand; LC: Collection of Lei Cai, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China; MUCL: Mycothèque de l’Université Catholique de Louvain, Belgium; SDBR-CMU: the Culture Collection of the Sustainable Development of Biological Resources Laboratory, Faculty of Science, Chiang Mai University, Chiang Mai, Thailand; TBRC: Thailand Bioresource Research Center, Thailand. Species obtained in this study are in bold. Superscript "T" indicates type species and "–" represents the absence of sequence data in GenBank.
5. Conclusions

Leaf spots caused by *Paramyrothecium* spp. were isolated from commercially important plants (*Coffee arabica*, *Tectona grandis*, *Vigna mungo*, and *V. unguiculata*), and non-commercial plants (*Aristolochia* sp., *Centrosema* sp., *Coccinia grandis*, *Commelina benghalensis*, *Lablab purpureus*, *Oroxylum indicum*, *Psophocarpus* sp., *Solanum virginianum*, *Spilanthes* sp., and *Vigna* sp.) in northern Thailand. Based on morphology and concatenated (ITS, *cmdA*, *rpb2*, and *tub2*) phylogeny, *P. vignicola*, *P. breviseta*, *P. eichhorniae*, and *P. folicola* were identified. The pathogenicity of each isolate was proven using Koch’s postulates. The pathogenicity assay revealed that all the isolates can cause the leaf spot disease. Interestingly, cross pathogenicity assay proved the ability of all 16 isolates to cause the disease on a wide range of hosts.

Author Contributions: Conceptualization, R.C. and M.C.S.; methodology, P.W., S.H., N.T., C.S., A.K., PS. and PP.; validation, R.C., M.C.S., N.S., J.K. and P.W.J.T.; investigation, P.W., S.H., N.T., C.S., A.K. and P.P.; writing—original draft preparation, P.W., S.H. and C.S.; writing—review and editing, all authors; supervision, R.C.; funding acquisition, R.C. All authors have read and agreed to the published version of the manuscript.

Funding: This research project was supported by Fundamental Fund 2022 (FF65/002), Chiang Mai University.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Publicly available datasets were analyzed in this study. These data can be found here: https://www.ncbi.nlm.nih.gov/ (access on 30 June 2022).

Acknowledgments: This research work was partially supported by Graduate School Chiang Mai University, TA/RA Scholarship, and Chiang Mai University. We thank Hiran A. Ariyawansa for his valuable suggestions and help.

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

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