

New species in *Dictyosporium*, new combinations in *Dictyocheirospora* and an updated backbone tree for Dictyosporiaceae

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Abstract

A survey of freshwater fungi on submerged wood in China and Thailand resulted in the collection of three species in *Dictyocheirospora* and four species in *Dictyosporium* including two new species in the latter genus. Morphological characters and phylogenetic analyses based on ITS, LSU and TEF1 α sequence data support their placement in *Dictyocheirospora* and *Dictyosporium* (Dictyosporiaceae). An updated backbone tree is provided for the family Dictyosporiaceae. Descriptions and illustrations of the new taxa and re-collections are provided. Four new combinations are proposed for *Dictyocheirospora*.

Keywords

2 new taxa, asexual morph, Dothideomycetes, phylogeny, taxonomy

Introduction

The family Dictyosporiaceae was introduced by Boonmee et al. (2016) to accommodate mostly aquatic lignicolous species with cheiroid, digitate, palmate and/or dictyosporous conidia and their sexual morphs that form a monophyletic clade in the class Dothideomycetes.

Dictyosporium, the type genus of the family, has been reported worldwide from dead wood and plant litter in terrestrial and aquatic habitats (Hyde and Goh 1998, Ho et al. 2002, Pinnoi et al. 2006, Pinruan et al. 2007). Corda (1836) established the genus with *D. elegans* Corda as the type species. The holomorph genus is characterised by dark brown, subglobose superficial ascomata, bitunicate cylindrical asci and hyaline, fusiform uniseptate ascospores with or without a sheath; sporodochial colonies, micronematous to macronematous conidiophores and cheiroid, digitate complanate conidia with several parallel rows of cells. Goh et al. (1999) reviewed the genus accepting 22 species and the remaining 16 species were doubtful or excluded. Tsui et al. (2006) first considered that the genus is closely related to Massarinaceae (Pleosporales) based on phylogenetic analysis using SSU and LSU sequence data. Tanaka et al. (2015) and Boonmee et al. (2016) confirmed the phylogenetic placement of *Dictyosporium* in Dictyosporiaceae (Massarineae, Pleosporales). Recent comparisons of *Dictyosporium* species were provided by Whitton et al. (2012), Prasher and Verma (2015) and Silva et al. (2015) with up to 48 accepted species. Since Silva et al. (2015), *D. araucariae* S.S. Silva, R.F. Castañeda & Gusmão, *D. hydei* I.B. Prasher & R.K. Verma, *D. indicum* I.B. Prasher & R.K. Verma, *D. olivaceosporum* Kaz. Tanaka, K. Hiray., Boonmee & K.D. Hyde, *D. palmae* Abdel-Aziz, *D. pseudomusae* Kaz. Tanaka, G. Sato & K. Hiray., *D. sexualis* Boonmee & K.D. Hyde, *D. splendidum* Alves-Barb., Malosso & R.F. Castañeda and *D. wuyiense* Y. Zhang & G.Z. Zhao were newly introduced to the genus (Prasher and Verma 2015, Tanaka et al. 2015, Abdel-Aziz 2016, Boonmee et al. 2016, da Silva et al. 2016, Alves-Barbosa et al. 2017, Zhang et al. 2017) and nine species were re-assigned to *Dictyocheirospora*, *Jalapriya* and *Vikalpa* (Boonmee et al. 2016). Wijayawardene et al. (2017a) provided information on the availability of cultures and references to accessible sequence data.

Dictyocheirospora was introduced by Boonmee et al. (2016) with *Di. rotunda* D'souza, Bhat & K.D. Hyde as the type species. *Dictyocheirospora* is morphologically similar to *Dictyosporium* except in having cheiroid, non-complanate or cylindrical conidia, mostly with conidial arms closely gathered together at the apex. Ten species are accepted in the genus including four species transferred from *Dictyosporium* (Boonmee et al. 2016, Wang et al. 2016, Hyde et al. 2017, Li et al. 2017).

During a survey of freshwater fungi on submerged wood along a north/south gradient in the Asian/Australasian region (Hyde et al. 2016), two new freshwater species and five previously described species were collected and identified based on phylogenetic analyses and morphological characters. We therefore introduce *Dictyosporium tubulatum* and *Dictyosporium tratense* as new species, with an illustrated account and

phylogenetic evidence for the new taxa. An updated backbone tree based on the combined ITS, LSU and TEF1 α sequence data is provided for Dictyosporiaceae. Four new combinations are proposed in *Dictyocheirospora*.

Materials and methods

Collection and examination of specimens

Specimens of submerged, decaying wood were collected from streams in Chiang Rai, Prachuap Khiri Khan, Phang Nga and Trat Provinces, Thailand, in December 2014, 2015, April 2016 and Guizhou Province, China, in October 2016. Specimens were brought to the laboratory in plastic bags and incubated in plastic boxes lined with moistened tissue paper at room temperature for one week. Morphological observations were made using a Motic SMZ 168 Series dissecting microscope for fungal structures on natural substrate. The fungal structures were collected using a syringe needle and transferred to a small drop of distilled water on a clean slide and covered with a cover glass. The fungi were examined using a Nikon ECLIPSE 80i compound microscope and photographed with a Canon 550D, 600D or 70D digital camera fitted to the microscope. Measurements were made with the TAROSOFT (R) IMAGE FRAME WORK programme and images used for figures were processed with ADOBE PHOTOSHOP CS6 software. Single spore isolations were made on to potato dextrose agar (PDA) or water agar (WA) and later transferred on to malt extract agar (MEA) or PDA following the method of Chomnunti et al. (2014). Specimens (dry wood with fungal material) are deposited in the herbarium of Mae Fah Luang University (MFLU), Chiang Rai, Thailand and Kunming Institute of Botany, Academia Sinica (HKAS), China. Axenic cultures are deposited in Mae Fah Luang University Culture Collection (MFLUCC). Facesoffungi and Index Fungorum numbers are registered as outlined in Jayasiri et al. (2015) and Index Fungorum (2018).

DNA extraction, PCR amplification and sequencing

Isolates were grown on PDA and/or MEA medium at 25 °C for one month. Fungal mycelium was scraped off and transferred to a 1.5-ml microcentrifuge tube using a sterilised lancet for genomic DNA extraction. Ezup Column Fungi Genomic DNA Purification Kit (Sangon Biotech, China) was used to extract DNA following the manufacturer's instructions. ITS, LSU and TEF1 α gene regions were amplified using the primer pairs ITS5 or ITS1 with ITS4 (Vilgalys and Hester 1990), LROR with LR5 or LR7 (White et al. 1990) and EF1-983F with EF1-2218R (Rehner 2001). The amplifications were performed in a 25 μ l reaction volume containing 9.5 μ l ddH₂O, 12.5 μ l 2 \times Taq PCR Master Mix with blue dye (Sangon Biotech, China), 1 μ l of DNA template and 1 μ l of each primer (10 μ M). The amplification condition for ITS, LSU and TEF1 α consisted of initial denaturation at 94 °C for 3 min; followed by 40 cycles of 45 s at 94 °C, 50 s at

56 °C and 1 min at 72 °C and a final extension period of 10 min at 72 °C. Purification and sequencing of PCR products were carried out using the above-mentioned PCR primers at Sangon Biotech (Shanghai) Co. Ltd. in China.

Phylogenetic analyses

The taxa included in the phylogenetic analyses were selected and obtained from previous studies and GenBank (Boonmee et al. 2016, Wang et al. 2016, Li et al. 2017). Three gene regions (ITS, LSU and TEF1 α) were used for the combined sequence data analyses. SEQMAN v. 7.0.0 (DNASTAR, Madison, WI) was used to assemble consensus sequences. The sequences were aligned using the online multiple alignment programme MAFFT v.7 (<http://mafft.cbrc.jp/alignment/server/>) (Katoh and Standley 2013). The alignments were checked visually and improved manually where necessary.

Phylogenetic analysis of the sequence data consisted of maximum likelihood (ML) using RAxML-HPC v.8 (Stamatakis 2006, Stamatakis et al. 2008) on the XSEDE Teragrid of the CIPRES science Gateway (<https://www.phylo.org>) (Miller et al. 2010) with rapid bootstrap analysis, followed by 1000 bootstrap replicates. The final tree was selected amongst suboptimal trees from each run by comparing likelihood scores under the GTRGAMMA substitution model.

Maximum parsimony (MP) analyses were performed with PAUP v. 4.0b10 (Swofford 2003) using the heuristic search option with 1000 random taxa addition and tree bisection and reconnection (TBR) as the branch swapping algorithm. All characters were unordered and of equal weight and gaps were treated as missing data. Maxtrees were unlimited, branches of zero length were collapsed and all multiple, equally parsimonious trees were saved. Clade stability was assessed using a bootstrap (BT) analysis with 1000 replicates, each with 10 replicates of random stepwise addition of taxa (Hillis and Bull 1993).

The programme MRMODELTEST2 v. 2.3 (Nylander 2008) was used to infer the appropriate substitution model that would best fit the model of DNA evolution for the combined datasets for Bayesian inference analysis with GTR+G+I substitution model selected. Posterior probabilities (PP) (Rannala and Yang 1996, Zhaxybayeva and Gogarten 2002) were determined by Markov Chain Monte Carlo sampling (MCMC) in MRBAYES v. 3.0b4 (Huelsenbeck and Ronquist 2001). Six simultaneous Markov chains were run for 1 million generations, with trees sampled every 100 generations (resulting in 10000 trees). The first 2000 trees, representing the burn-in phase of the analyses were discarded and the remaining 8000 trees were used for calculating posterior probabilities (PP) in the majority rule consensus tree (Larget and Simon 1999).

The resulting trees were printed with FIGTREE v. 1.4.0 (<http://tree.bio.ed.ac.uk/software/figtree/>) and the layout was created in MICROSOFT POWERPOINT for Mac v. 15.19.1. The alignment of phylogenetic analyses and resultant tree were deposited in TreeBASE (www.treebase.org, submission number 22802). Sequences generated in this study were submitted to GenBank (Table 1).

Table 1. Isolates and sequences used in this study (newly generated sequences are indicated in bold, ex-type strains are indicated with ^T after strain number).

| Species | Source | GenBank accession number | | |
|--|-----------------------------------|--------------------------|-----------------|-----------------|
| | | ITS | LSU | TEF1 α |
| <i>Aquadictyospora lignicola</i> | MFLUCC 17-1318 ^T | MF948621 | MF948629 | MF953164 |
| <i>Aquaticheirosora lignicola</i> | HKUCC 10304 ^T | AY864770 | AY736378 | – |
| <i>Cheirosporium triseriale</i> | HMAS 180703 ^T | EU413953 | EU413954 | – |
| <i>Dendryphiella eucalyptorum</i> | CBS 137987 ^T | KJ869139 | KJ869196 | – |
| <i>Dendryphiella fasciculata</i> | MFLUCC 17-1074 ^T | MF399213 | MF399214 | – |
| <i>Dendryphiella paravinoso</i> | CBS 141286 ^T | KX228257 | KX228309 | – |
| <i>Dictyocheirosora aquatica</i> | KUMCC 15-0305 ^T | KY320508 | KY320513 | – |
| <i>Dictyocheirosora bannica</i> | KH 332 ^T | LC014543 | AB807513 | AB808489 |
| <i>Dictyocheirosora bannica</i> | MFLUCC 16-0874 | MH381765 | MH381774 | – |
| <i>Dictyocheirosora garethjonesii</i> | MFLUCC 16-0909 ^T | KY320509 | KY320514 | – |
| <i>Dictyocheirosora garethjonesii</i> | DLUCC 0848 | MF948623 | MF948631 | MF953166 |
| <i>Dictyocheirosora gigantea</i> | BCC 11346 | DQ018095 | – | – |
| <i>Dictyocheirosora heptaspora</i> | CBS 396.59 | DQ018090 | – | – |
| <i>Dictyocheirosora indica</i> | MFLUCC 15-0056 | MH381763 | MH381772 | MH388817 |
| <i>Dictyocheirosora pseudomusae</i> | yone 234 ^T | LC014550 | AB807520 | AB808496 |
| <i>Dictyocheirosora rotunda</i> | MFLUCC 14-02 ⁹³ T | KU179099 | KU179100 | – |
| <i>Dictyocheirosora rotunda</i> | MFLUCC 17-0222 | MH381764 | MH381773 | MH388818 |
| <i>Dictyocheirosora rotunda</i> | MFLUCC 17-1313 | MF948625 | MF948633 | MF953168 |
| <i>Dictyocheirosora subramanianii</i> | BCC 3503 | DQ018094 | – | – |
| <i>Dictyocheirosora vinaya</i> | MFLUCC 14-0294 ^T | KU179102 | KU179103 | – |
| <i>Dictyosporium alatum</i> | ATCC 34953 ^T | NR_077171 | DQ018101 | – |
| <i>Dictyosporium aquaticum</i> | MF 1318 ^T | KM610236 | – | – |
| <i>Dictyosporium bulbosum</i> | yone 221 | LC014544 | AB807511 | AB808487 |
| <i>Dictyosporium digitatum</i> | KH 401 | LC014545 | AB807515 | AB808491 |
| <i>Dictyosporium digitatum</i> | yone 280 | LC014547 | AB807512 | AB808488 |
| <i>Dictyosporium elegans</i> | NBRC 32502 ^T | DQ018087 | DQ018100 | – |
| <i>Dictyosporium hughesii</i> | KT 1847 | LC014548 | AB807517 | AB808493 |
| <i>Dictyosporium metosporum</i> | MFLUCC 10-0131 ^T | KP710944 | KP710945 | – |
| <i>Dictyosporium nigroapice</i> | BCC 3555 | DQ018085 | – | – |
| <i>Dictyosporium nigroapice</i> | MFLUCC 17-2053 | MH381768 | MH381777 | MH388821 |
| <i>Dictyosporium olivaceosporum</i> | KH 375 ^T | LC014542 | AB807514 | AB808490 |
| <i>Dictyosporium sexualis</i> | MFLUCC 10-0127 ^T | KU179105 | KU179106 | – |
| <i>Dictyosporium</i> sp. | MFLUCC 15-0629 | MH381766 | MH381775 | MH388819 |
| <i>Dictyosporium stellatum</i> | CCFC 241241 ^T | NR_154608 | JF951177 | – |
| <i>Dictyosporium strelitziae</i> | CBS 123359 ^T | NR_156216 | FJ839653 | – |
| <i>Dictyosporium tetrasporum</i> | KT 2865 | LC014551 | AB807519 | AB808495 |
| <i>Dictyosporium thailandicum</i> | MFLUCC 13-0773 ^T | KP716706 | KP716707 | – |
| <i>Dictyosporium tratense</i> | MFLUCC 17-2052^T | MH381767 | MH381776 | MH388820 |
| <i>Dictyosporium tubulatum</i> | MFLUCC 15-0631^T | MH381769 | MH381778 | MH388822 |
| <i>Dictyosporium tubulatum</i> | MFLUCC 17-2056 | MH381770 | MH381779 | – |
| <i>Dictyosporium wuyiense</i> | CGMCC 3.18703 ^T | KY072977 | – | – |
| <i>Dictyosporium zhejiangense</i> | MW-2009a ^T | FJ456893 | – | – |
| <i>Digitodesmium bambusicola</i> | CBS 110279 ^T | DQ018091 | DQ018103 | – |

| Species | Source | GenBank accession number | | |
|---|-----------------------------|--------------------------|----------|---------------|
| | | ITS | LSU | TEF1 α |
| <i>Gregarithecium curvisporum</i> | KT 922 ^T | AB809644 | AB807547 | – |
| <i>Jalapriya inflata</i> | NTOU 3855 | JQ267362 | JQ267363 | – |
| <i>Jalapriya pulchra</i> | MFLUCC 15-0348 ^T | KU179108 | KU179109 | – |
| <i>Jalapriya pulchra</i> | MFLUCC 17-1683 | MF948628 | MF948636 | MF953171 |
| <i>Jalapriya toruloides</i> | CBS 209.65 | DQ018093 | DQ018104 | – |
| <i>Periconia igniaria</i> | CBS 379.86 | LC014585 | AB807566 | AB808542 |
| <i>Periconia igniaria</i> | CBS 845.96 | LC014586 | AB807567 | AB808543 |
| <i>Pseudocoleophoma calamagrostidis</i> | KT 3284 ^T | LC014592 | LC014609 | LC014614 |
| <i>Pseudocoleophoma polygonicola</i> | KT 731 ^T | AB809634 | AB807546 | AB808522 |
| <i>Pseudocoleophoma typhicola</i> | MFLUCC 16-0123 ^T | KX576655 | KX576656 | – |
| <i>Pseudodictyosporium elegans</i> | CBS 688.93 ^T | DQ018099 | DQ018106 | – |
| <i>Pseudodictyosporium indicum</i> | CBS 471.95 | DQ018097 | – | – |
| <i>Pseudodictyosporium thailandica</i> | MFLUCC 16-0029 ^T | KX259520 | KX259522 | KX259526 |
| <i>Pseudodictyosporium wauense</i> | NBRC 30078 | DQ018098 | DQ018105 | – |
| <i>Pseudodictyosporium wauense</i> | DLUCC 0801 | MF948622 | MF948630 | MF953165 |
| <i>Vikalpa australiensis</i> | HKUCC 8797 ^T | DQ018092 | – | – |

Phylogenetic results

The analysed dataset consisted of combined ITS (557 bp), LSU (803 bp) and TEF1 α (918 bp) sequence data (a total of 2278 characters including gaps) for 59 taxa in Dictyosporiaceae with *Periconia igniaria* E.W. Mason & M.B. Ellis (CBS 379.86, CBS 845.96) as the outgroup taxon. The best scoring RAxML tree is shown in Figure 1.

Phylogenetic analyses indicated the placement of three isolates (MFLUCC 15-0056, MFLUCC 16-0874 and MFLUCC 17-0222) within the genus *Dictyocheirosora*. Five isolates (MFLUCC 15-0629, MFLUCC 17-2052, MFLUCC 17-2056, MFLUCC 15-0631 and MFLUCC 17-2053) nested in *Dictyosporium*. Phylogenetic results showed that *Dictyocheirosora indica* (MFLUCC 15-0056) clustered with *Dictyocheirosora subramaniani* (B. Sutton) D'souza, Boonmee & K.D. Hyde (BCC 3503) with good support. *Dictyocheirosora bannica* (MFLUCC 16-0874) was placed as sister taxon to the ex-type strain *Dictyocheirosora bannica* (KH 332). *Dictyocheirosora rotunda* (MFLUCC 17-0222) grouped together with *Dictyocheirosora rotunda* (MFLUCC 17-1313) and the ex-type strain *Dictyocheirosora rotunda* (MFLUCC 14-0293) with strong support. The strain *Dictyosporium* sp. (MFLUCC 15-0629) clustered as sister taxon to *Dictyosporium digitatum* J.L. Chen, C.H. Hwang & Tzean (KH 401), *Dictyosporium aquaticum* Abdel-Aziz (MF 1318) and *Dictyosporium stellatum* G.P. White & Seifert (CCFC 241241). The new taxon *Dictyosporium tratense* (MFLUCC 17-2052) formed a single clade within *Dictyosporium* which is distinct from other species in the genus. The new collection *Dictyosporium nigroapice* (MFLUCC 17-2053) was placed as sister taxon to a previous isolate *Dictyosporium nigroapice* (BCC 3555). Two isolates of the new taxon *Dictyosporium tubulatum* (MFLUCC 15-0631 and MFLUCC 17-2056) nested in *Dictyosporium* as sister clade to *Dictyosporium nigroapice* (MFLUCC 17-2053 and BCC 3555).

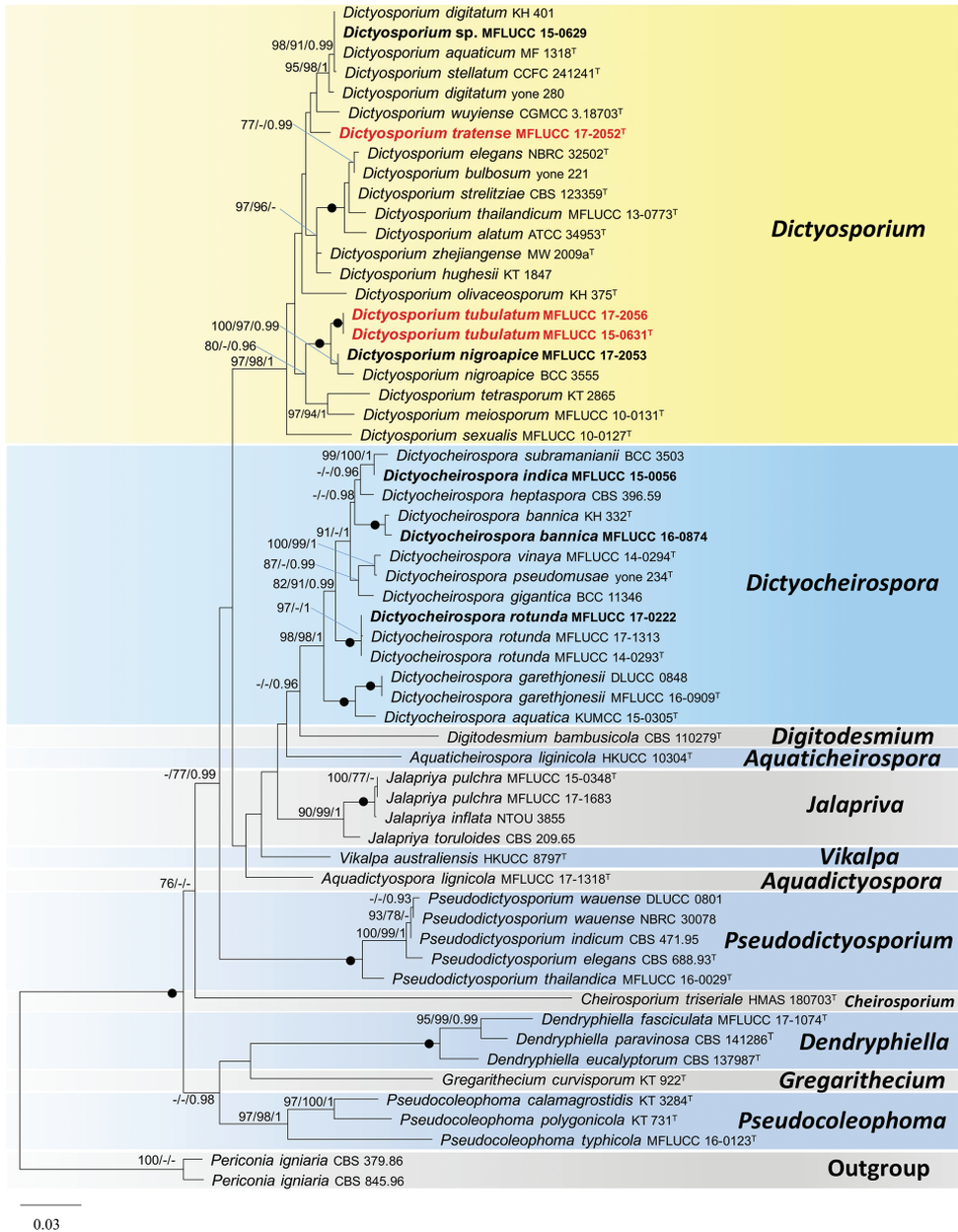


Figure 1. Maximum likelihood majority rule consensus tree for the analysed Dictyosporiaceae isolates based on a dataset of combined ITS, LSU and TEF1 α sequence data. Bootstrap support values for maximum likelihood (ML) and maximum parsimony (MP) greater than 75% and Bayesian posterior probabilities greater than 0.95 are indicated above the nodes as MLBS/MPBS/PP. The scale bar represents the expected number of changes per site. The tree is rooted with *Periconia igniaria* (CBS 379.86, CBS 845.96). The strain numbers are noted after the species names with ex-type strains indicated with ^T. The new collections are in bold with new taxa in red. Branches with 100% ML BS, 100% MP BS and 1.0 PP are shown as black nodes. Genera are indicated as coloured blocks.

Taxonomy

Dictyocheiросpora species

***Dictyocheiросpora bannica* Kaz. Tanaka, K. Hiray., Boonmee & K.D. Hyde, Fungal Diversity 80: 467 (2016)**

Index Fungorum number: IF551997

Facesoffungi number: FoF02014

Figure 2

Material examined. THAILAND. Phang Nga Province, Bann Tom Thong Khang, on decaying wood submerged in a freshwater stream, 17 Dec 2015, J. Yang, Site 7-70-1 (MFLU 18-1040, HKAS 102131), living culture MFLUCC 16-0874 (Additional SSU sequence GenBank MH381759).

Notes. The phylogenetic result showed the strain MFLUCC 16-0874 clustered with the ex-type (KH 332) of *Dictyocheiросpora bannica*. The morphological examination of this collection matched well with the holotype of *Dictyocheiросpora bannica* (Boonmee et al. 2016). *Dictyocheiросpora bannica* was previously collected in Japan, while this is a new record for Thailand.

***Dictyocheiросpora hydei* (I.B. Prasher & R.K. Verma) J. Yang & K.D. Hyde, comb. nov.**

Index Fungorum number: IF554773

Facesoffungi number: FoF04679

Basionym. *Dictyosporium hydei* I.B. Prasher & R.K. Verma, Phytotaxa 204 (3): 196 (2015).

Holotype. INDIA. Himachal Pradesh, Bilaspur, on bark of *Tecoma stans*, 17 September 2013, I.B. Prasher and R.K. Verma (PAN 30364).

Notes. Considering the latest generic concept of *Dictyocheiросpora* and *Dictyosporium*, we suggest that *Dictyosporium hydei* should be referred to *Dictyocheiросpora* with the key character of non-complanate or cylindrical conidia with conidial arms closely gathered together at the apex. We have not examined the holotype of *Dictyocheiросpora hydei*. The details provided by Prasher and Verma (2015) are adequate being illustrative and descriptive.

***Dictyocheiросpora indica* (I.B. Prasher & R.K. Verma) J. Yang & K.D. Hyde, comb. nov.**

Index Fungorum number: IF554774

Facesoffungi number: FoF04680

Figure 3

Basionym. *Dictyosporium indicum* I.B. Prasher & R.K. Verma, Phytotaxa 204 (3): 194 (2015).



Figure 2. *Dictyocheiropora bannica* (MFLU 18-1040) **a** Colonies on submerged wood **b** Conidia and conidiophores **c–f** Conidia **g** Germinated conidium **h, i** Culture, h from above, i from reverse. Scale bars: **a** = 200 μm , **b, g** = 50 μm , **c–f** = 30 μm .

Holotype. INDIA. Himachal Pradesh, Mandi, on petiole of *Phoenix rupicola*, 19 November 2012, I.B. Prasher and R.K. Verma (PAN 30313).

Material examined. THAILAND. Chiang Rai, stream flowing in Tham Luang Nang Non Cave, on decaying submerged wood, 25 November 2014, J. Yang, YJ-3 (MFLU 15-1169 **reference specimen designated here**, HKAS 102135), living culture MFLUCC 15-0056 (Additional SSU sequence GenBank MH381757).

Notes. Collection MFLU 15-1169 was identified as *Dictyocheiropora indica* (*Dictyosporium indicum*) based on morphological examination. Phylogenetic analyses indicated the placement of this taxon within *Dictyocheiropora* and sister to *Di. subramanianii* (BCC 3503). *Dictyocheiropora subramanianii* differs from *Di. indica* in lacking appendages. *Dictyocheiropora indica* resembles *Di. musae* in having non-complanate, cylindrical conidia with globose to subglobose appendages. However, conidial append-

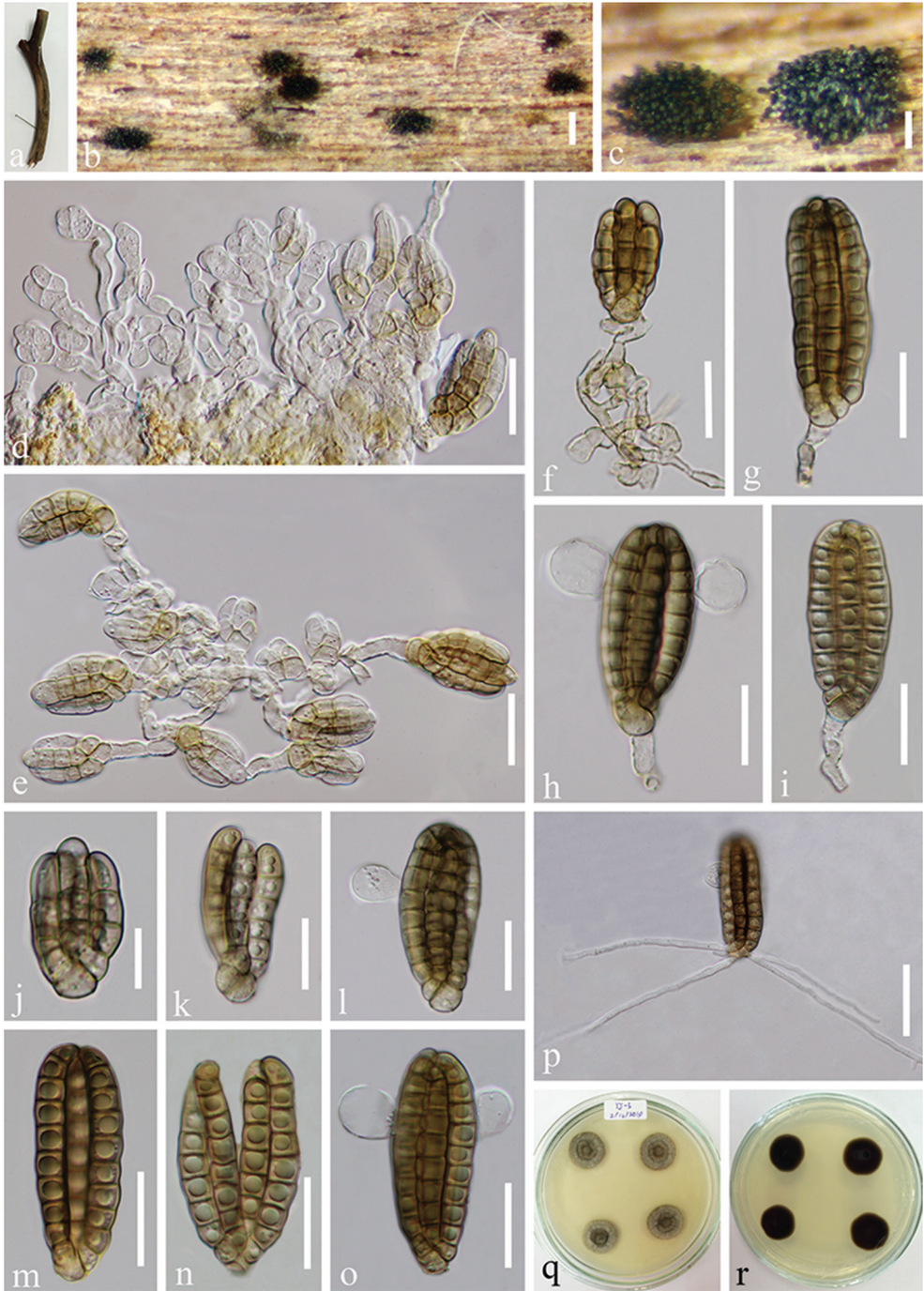


Figure 3. *Dictyocheirospora indica* (MFLU 15-1169, reference specimen). **a** Substrate **b, c** Colonies on woody substrate **d, e** Conidial formation **f–i** Conidia with partial conidiophores **j–o** Conidia **p** Germinated conidium **q–r** Culture, q from above, r from reverse. Scale bars: **b** = 200 μ m, **c** = 100 μ m, **d–i, l–o** = 20 μ m, **j** = 10 μ m, **k** = 15 μ m, **p** = 30 μ m.

ages of *Di. indica* are attached at the subapical cells, while appendages of *Di. musae* are attached at the central cells of the outer cell-row. The conidial size of *Di. indica* (33–48 × 13–18 µm) is smaller than that of *Di. musae* (45–65 × 20–27 µm) (Photita et al. 2002, Prasher and Verma 2015). In this study, sequence data of our collection *Dictyocheiropora indica* (MFLUCC 15-0056) was generated and, as there is no sequence data available for the previous collection (*Dictyosporium indicum*), we therefore designated our collection as the reference specimen (*sensu* Ariyawansa et al. 2014) for *Dictyocheiropora indica*.

***Dictyocheiropora musae* (Photita) J. Yang, K.D. Hyde & Z.Y. Liu, comb. nov.**

Index Fungorum number: IF554775

Facesoffungi number: FoF04681

Basionym. *Dictyosporium musae* Photita, Mycotaxon 82: 416 (2002)

Holotype. THAILAND. Mae Hong Son Province, Sob Mei, Huay Thicha Village, on decaying petioles of *Musa acuminata*, 23 November 2000, W. Photita (PDD 74135).

Notes. *Dictyocheiropora musae* is morphologically similar to *Di. hydei* in having non-complanate, cylindrical conidia with globose to subglobose appendages. However, *Dictyocheiropora musae* differs in having appendages in the middle cells while *Di. hydei* has appendages on the basal cells (Photita et al. 2002, Prasher and Verma 2015).

***Dictyocheiropora rotunda* D'souza, Bhat & K.D. Hyde, Fungal Diversity 80: 465 (2016)**

Index Fungorum number: IF551581

Facesoffungi number: FoF01262

Figure 4

Material examined. CHINA. Guizhou Province, Anshun city, Gaodang village, 26°4.267'N, 105°41.883'E, on decaying wood submerged in Suoluo river, 19 October 2016, J. Yang, GD 2-3 (MFLU 18-1041, HKAS 102132), living culture MFLUCC 17-0222 (Additional SSU sequence GenBank MH381758).

Notes. This species is known in China and Thailand from freshwater habitats (Boonmee et al. 2016, Wang et al. 2016).

***Dictyocheiropora tetraploides* (L. Cai & K.D. Hyde) J. Yang & K.D. Hyde, comb. nov.**

Index Fungorum number: IF554776

Facesoffungi number: FoF04682

Basionym. *Dictyosporium tetraploides* L. Cai & K.D. Hyde, Sydowia 55 (2): 132 (2003)

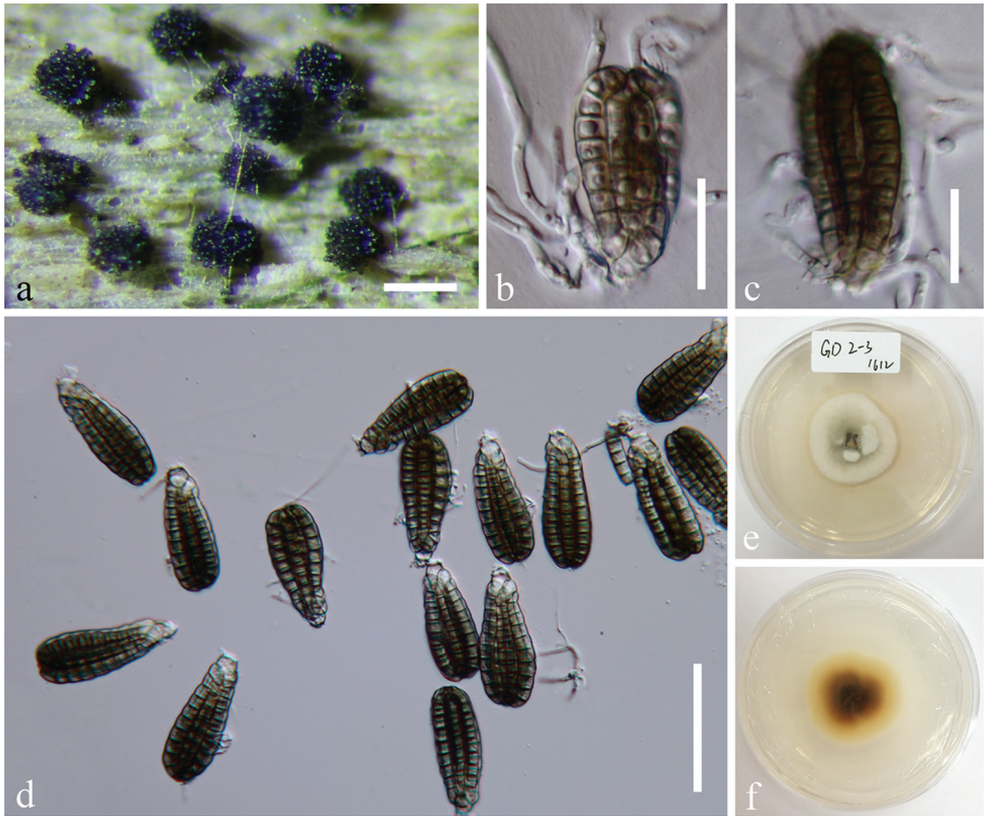


Figure 4. *Dictyocheirospora rotunda* (MFLU 18-1041). **a** Colonies on submerged wood **b, c** Germinated conidia **d** Conidia **e, f** Culture, e from above, f from reverse. Scale bars: **a** = 200 μ m, **b, c** = 20 μ m, **d** = 50 μ m.

Holotype. CHINA. Yunnan, Xishuangbanna, Menglun, a small stream, on submerged wood, 21 June 2002, L. Cai (HKUM 17146).

Notes. *Dictyocheirospora tetraploides* is morphologically similar to *Di. musae* in conidial shape, size, colour and appendages. However, conidia of *Di. tetraploides* have 5-rowed cells, while those of *Di. musae* are 7-rowed cells (Photita et al. 2002, Cai et al. 2003).

Dictyosporium species

Dictyosporium tubulatum J. Yang, K.D. Hyde & Z.Y. Liu, sp. nov.

Index Fungorum number: IF554771

Facesoffungi number: FoF04677

Figure 5

Etymology. Referring to the tubular conidial appendages.



Figure 5. *Dictyosporium tubulatum* (MFLU 15-1166, holotype). **a, b** Colonies on woody substrate **c** Squash mount of a sporodochium **d-g** Conidia **h-i** Conidia with conidiophores **j-l** Conidia with appendages **m** lateral view of a conidium **n** Germinated conidium **o, p** Culture, o from above **p** from reverse. Scale bars: **a** = 1000 μm , **b** = 200 μm , **c, n** = 30 μm , **d, e** = 10 μm , **f-m** = 15 μm .

Description. *Saprobic* on decaying plant substrates. **Asexual morph:** Colonies punctiform, sporodochial, scattered, dark brown to black, glistening. *Mycelium* mostly immersed, composed of smooth, septate, branched, hyaline to pale brown hyphae. *Conidiophores* micronematous, mononematous, septate, cylindrical, hyaline to pale brown, smooth-walled, $6.5\text{--}15 \times 3.5\text{--}6 \mu\text{m}$, sometimes reduced to conidiogenous cells. *Conidiogenous cells* monoblastic, integrated, terminal, determinate, hyaline to pale brown. *Conidia* acrogenous, solitary, cheiroid, smooth-walled, complanate, yellowish-brown to medium brown, mostly consisting of four arms closely compact with side arms lower than middle arms, rarely with five arms, 5–7-euseptate in each arm, guttulate, $(22\text{--})29\text{--}35(\text{--}38) \times (14\text{--})17\text{--}19(\text{--}22) \mu\text{m}$ ($\bar{x} = 32.5 \times 18 \mu\text{m}$, $n = 40$), with hyaline, tubular, elongated appendages which are $19\text{--}24 \times 3.5\text{--}7 \mu\text{m}$ and mostly attached at the apical part of two outer arms. **Sexual morph:** Undetermined.

Cultural characteristics. Conidia germinating on PDA within 24 h and germ tubes produced from the basal cell. Colonies on MEA reaching 5–10 mm diam. in a week at 25 °C, in natural light, circular, with fluffy, dense, white mycelium on the surface with entire margin; in reverse yellow in the middle and white at the margin.

Material examined. THAILAND. Prachuap Khiri Khan Province, near $12^{\circ}30.19'N$, $99^{\circ}31.35'E$, on decaying wood submerged in a freshwater stream, 25 December 2014, J. van Strien, Site 5-11-1 (MFLU 15-1166 **holotype**, HKAS 102136 **isotype**), ex-type living culture MFLUCC 15-0631; *ibid.* Trat Province, Amphoe Ko Chang, $12^{\circ}08'N$, $102^{\circ}38'E$, on decaying wood submerged in a freshwater stream, 27 April 2017, Y.Z. Lu, YJT 22-2 (MFLU 18-1044, HKAS 102137 **paratype**), living culture MFLUCC 17-2056.

Notes. Phylogenetic analyses showed that *Dictyosporium tubulatum* nested in *Dictyosporium* and sister to *D. nigroapice*. *Dictyosporium tubulatum* morphologically resembles *D. alatum* Emden, *D. canisporum* L. Cai & K.D. Hyde and *D. thailandicum* D' souza, D.J. Bhat & K.D. Hyde in conidial ontogeny and conidial shape, colour and appendages. *Dictyosporium tubulatum* differs from the three species in the number of conidial cell rows. There are mostly four conidial columns in *D. tubulatum* while mostly five columns in the others. *Dictyosporium tubulatum* has smaller conidia ($25\text{--}38 \times 14\text{--}22 \mu\text{m}$) than those in *D. canisporum* ($32.5\text{--}47.5 \times 20\text{--}25 \mu\text{m}$) but has similar conidial size with *D. alatum* ($26\text{--}32 \times 15\text{--}24 \mu\text{m}$) and *D. thailandicum* ($15.4\text{--}34.5 \times 14.5\text{--}20.6 \mu\text{m}$) (Cai et al. 2003, Liu et al. 2015). Based on the molecular phylogeny, *D. tubulatum* is distinct from *D. thailandicum* and *D. alatum*. Unfortunately, molecular data are unavailable for *D. canisporum*.

***Dictyosporium tratense* J. Yang & K.D. Hyde, sp. nov.**

Index Fungorum number: IF554772

Facesoffungi number: FoF04678

Figure 6

Etymology. Referring to the collecting site in Trat province, Thailand.

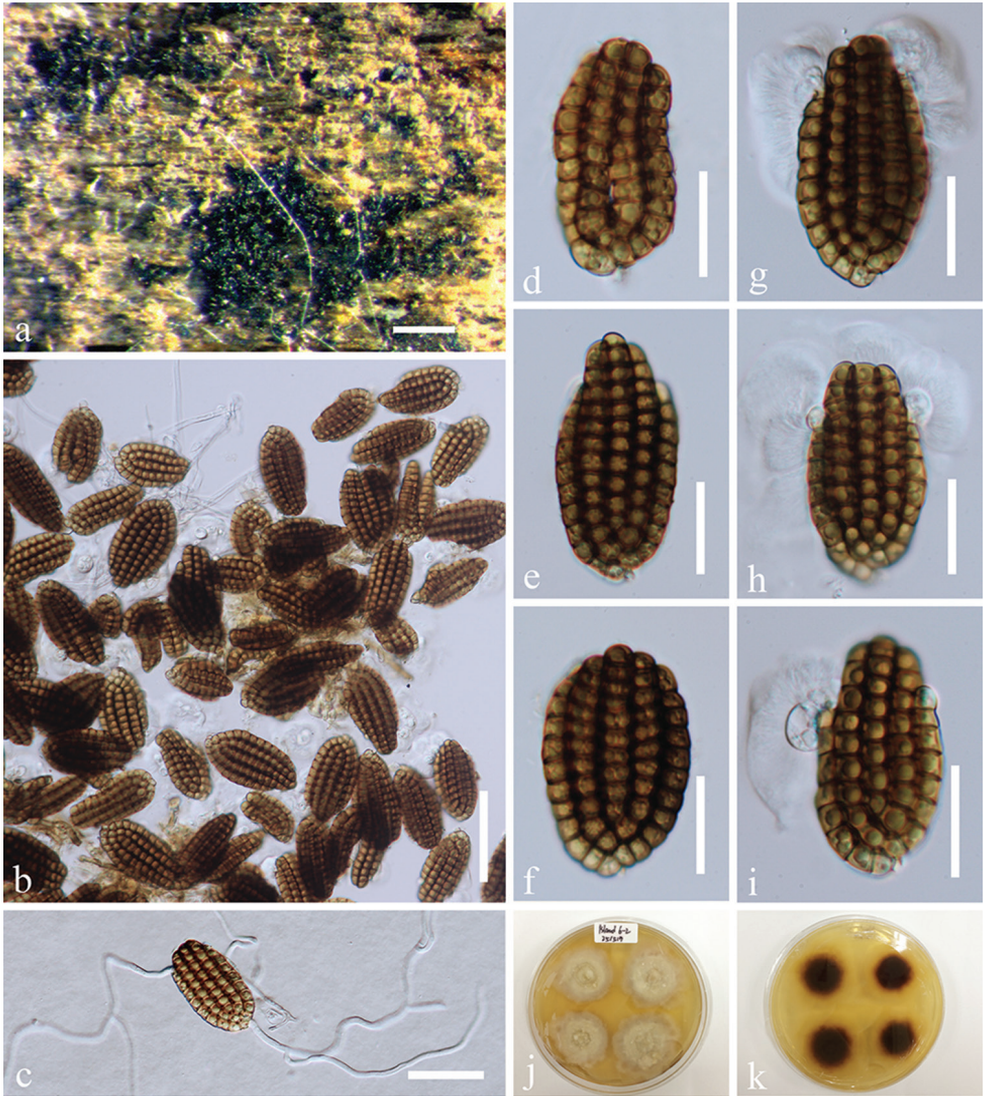


Figure 6. *Dictyosporium tratense* (MFLU 18-1042, holotype). **a** Colonies on submerged wood **b** Squash mount of a sporodochium **c** Germinated conidium **d–i** Conidia **j, k** Culture **j** from above **k** from reverse. Scale bars: **a** = 200 μm , **b** = 50 μm , **c** = 30 μm , **d–i** = 20 μm .

Description. *Saprobic* on decaying plant substrates. **Asexual morph:** *Colonies* punctiform, sporodochial, scattered, black, glistening. *Mycelium* mostly immersed, composed of smooth, septate, branched, hyaline to pale brown hyphae. *Conidiophores* micronematous, mononematous, septate, cylindrical, hyaline to pale brown, smooth-walled, sometimes reduced to conidiogenous cells. *Conidiogenous cells* monoblastic, integrated, terminal, determinate, hyaline to pale brown. *Conidia* (40–)43–54(–57) \times (20–)23–32(–36) μm (\bar{x} = 49.5 \times 26 μm , n = 40), acrogenous,

solitary, cheiroid, smooth-walled, complanate, yellowish-brown to light brown, consisting of 39–68 cells arranged in 4–6 (mostly 5) closely compact columns, 9–11-euseptate in each column, guttulate; the inner columns nested within the outer columns, the outer columns derived from the basal cell of the conidium; the intermediate columns are derived from the first or second cell of the outer columns; the inner columns derived from the first or second cell of the intermediate columns; usually with 2–3 central columns longest and of equal length, 2–3 peripheral columns shorter and of equal length; sometimes with hyaline globose appendages at the apical cells of outer columns with hyaline cloud-shaped mucilaginous sheath.

Sexual morph: Undetermined.

Cultural characteristics. Conidia germinating on PDA within 24 h and germ tubes produced from basal cell. Colonies on MEA reaching 5–10 mm diam. in a week at 25 °C, in natural light, circular, with fluffy, dense, pale yellow mycelium in the middle and sparse mycelium in the outer ring on the surface with irregular margin; in reverse, dark yellow to brown in the middle and pale yellow at the margin.

Material examined. THAILAND. Trat Province, Amphoe Ko Chang, 12°08'N, 102°38'E, on decaying wood submerged in a freshwater stream, 27 April 2017, Y.Z. Lu, YJT 6-2 (MFLU 18-1042 **holotype**, HKAS 102133 **isotype**), ex-type living culture MFLUCC 17-2052 (Additional SSU sequence GenBank MH381761).

Notes. Phylogenetic analyses indicated *Dictyosporium tratense* nested within *Dictyosporium* and close to *D. wuyiense*. It is distinguished from the other species in the genus in having a mucilaginous sheath. Morphologically, *D. tratense* is most comparable to *D. elegans* in conidial colour and shape, but conidia of the new taxon (40–57 × 20–36 µm) are smaller than those of *D. elegans* (40–80 × 24–36 µm) (Goh et al. 1999).

Dictyosporium sp.

Figure 7

Material examined. THAILAND. Prachuap Khiri Khan Province, near 12°30.19'N, 99°31.35'E, on decaying wood submerged in a freshwater stream, 25 December 2014, J. van Strien, Site 5-5-1 (MFLU 15-1164), living culture MFLUCC 15-0629 (Additional SSU sequence GenBank MH381760).

Notes. Phylogenetic analyses indicated the isolate *Dictyosporium* sp. (MFLUCC 15-0629) was placed as sister taxon to *D. digitatum* (KH 401), *D. aquaticum* (MF 1318) and *D. stellatum* (CCFC 241241) with good support. The strain *D. digitatum* (KH 401), *D. aquaticum* (MF 1318) and our strain MFLUCC 15-0629, showed the same nucleotide (490 bp) between them for ITS gene regions, while there is only one nucleotide difference between our strain and *D. stellatum* (CCFC 241241). However, the strain *Dictyosporium* sp. (MFLUCC 15-0629) showed seven nucleotides different from *D. digitatum* (yone 280) for ITS gene regions. Morphologically, *D. digitatum* and *D. aquaticum* share the character in having appendages borne at the terminal cells of each conidial arm (Chen et al. 1991, Liu et al. 2015). *Dictyosporium stellatum* dif-

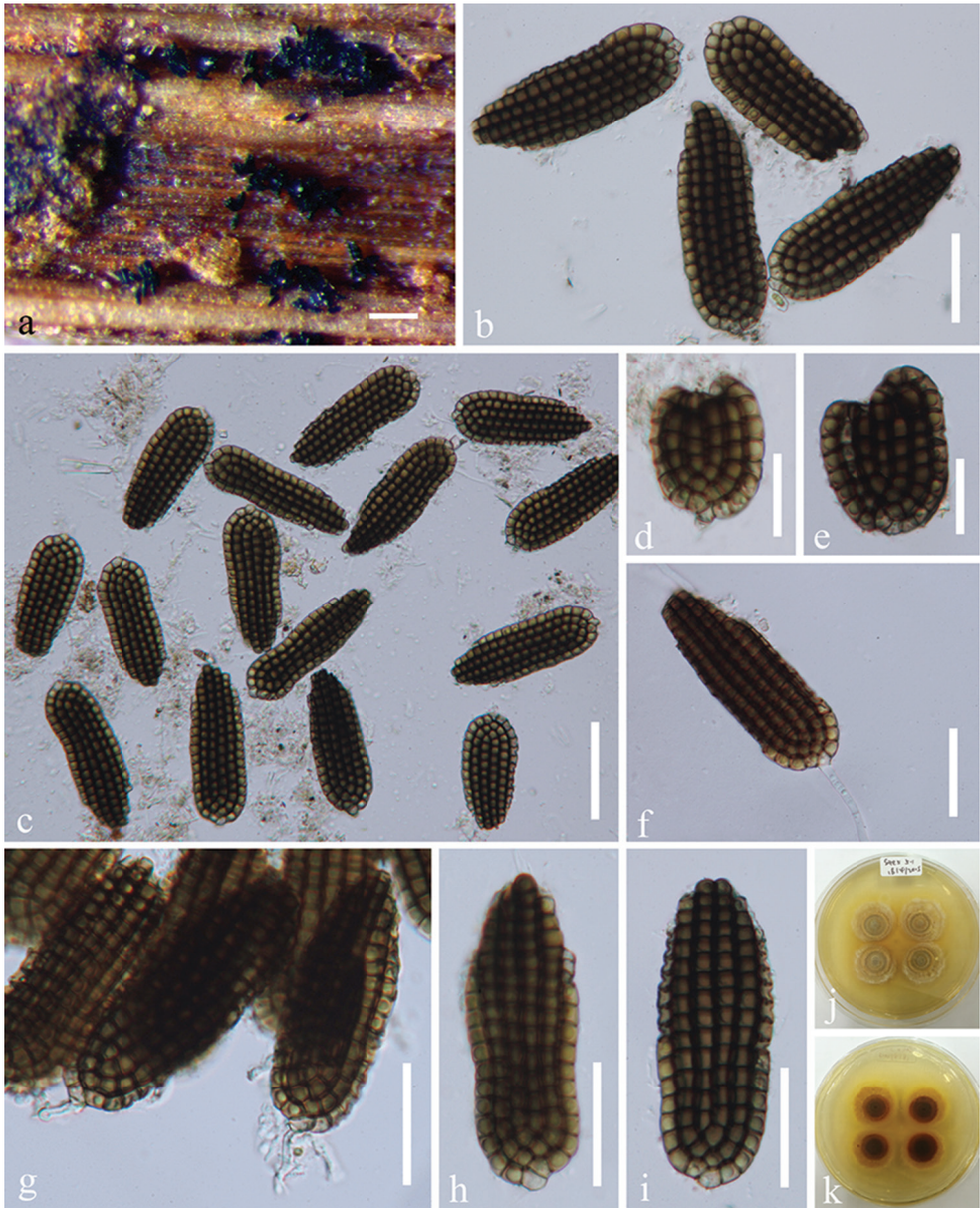


Figure 7. *Dictyosporium* sp. (MFLU 15-1164). **a** Colonies on submerged wood **b** Squash mount of a sporodochium; **c** Germinated conidium **b–e, h, i** Conidia **f** Germinated conidium **g** Conidia with conidiophores **j, k** Culture, j from above, k from reverse. Scale bars: **a** = 200 μm , **b, f–i** = 30 μm , **c** = 50 μm , **d, e** = 20 μm .

fers from *D. digitatum* and *D. aquaticum* in lacking conidial appendages (Crous et al. 2011). In this case, it is difficult to identify our collection based on the recommendations advocated by Jeewon and Hyde (2016) for differentiating species or establishing

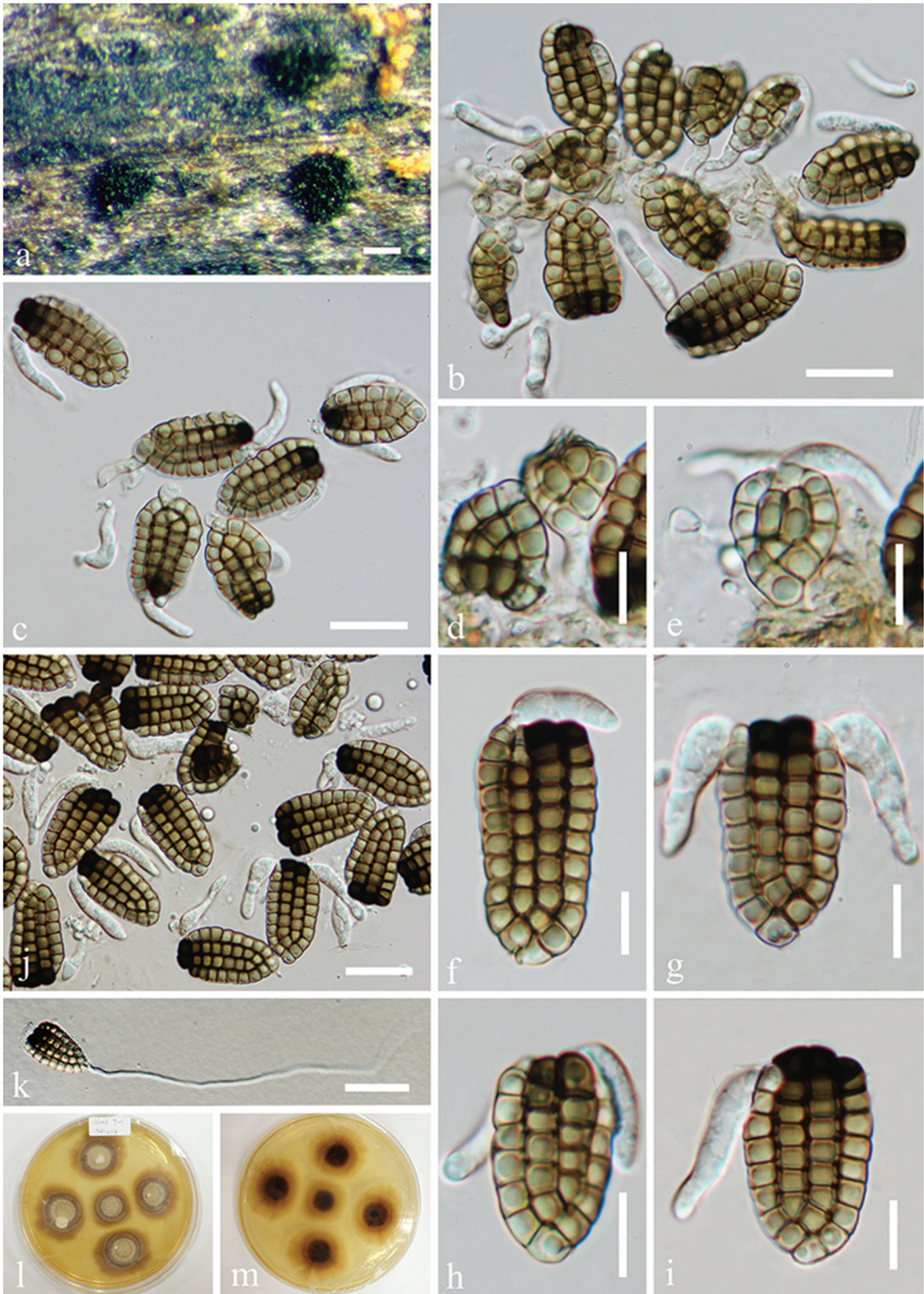


Figure 8. *Dictyosporium nigroapice* (MFLU18-1043). **a** Colonies on submerged wood **b, c** Conidia and conidiophores **d–j** Conidia **k** Germinated conidium **l, m** Culture, **l** from above, **m** from reverse. Scale bars: **a** = 100 μm , **b, c, j** = 20 μm , **d–i** = 10 μm , **k** = 30 μm .

new species. Thus, we recommend designating this collection as unknown species until enough evidence is available for its identification.

***Dictyosporium nigroapice* Goh, W.H. Ho & K.D. Hyde, Fungal Diversity 2: 83 (1999)**

Index Fungorum number: IF450470

Facesoffungi number: FoF04683

Figure 8

Material examined. THAILAND. Trat Province, Amphoe Ko Chang, 12°08'N, 102°38'E, on decaying wood submerged in a freshwater stream, 27 April 2017, Y.Z. Lu, YJT 7-1 (MFLU 18-1043, HKAS 102134), living culture MFLUCC 17-2053 (Additional SSU sequence GenBank MH381762).

Notes. Conidia in *Dictyosporium nigroapice* are characterised by conspicuously darker apical cells of the two inner arms, rarely darker at the apex of the outer arms. Morphological characters of this collection well agree with the original diagnosis of the holotype of *D. nigroapice* (Goh et al. 1999).

Discussion

Dictyosporiaceae accommodates a holomorphic group of Dothideomycetes, including 12 genera with nine being dictyosporous (Wijayawardene et al. 2017b, Wijayawardene et al. 2018). *Dictyocheiropora* and *Dictyosporium* are the two largest genera in the family. *Dictyosporium* has cheiroid, digitate and complanate conidia without separating arms, while *Dictyocheiropora* is characterised by non-complanate conidia with arms arising from the basal cell and closely gathered at the apex and compact. Thus, *Dictyosporium hydei*, *D. indicum*, *D. musae* and *D. tetraploides* are transferred to *Dictyocheiropora* based on the clear morphological characters. Phylogenetic analyses revealed the placement of *Dictyocheiropora indica* (MFLUCC 15-0056 reference specimen) within *Dictyocheiropora*. We believe that the other three species belong to *Dictyocheiropora* in having similar conidia and appendages to *Dictyocheiropora indica*, although molecular data are unavailable for them.

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specialist fungi associated with ants, *Rhododendron* species and *Dracaena* species” (grant no: DBG6080013) and “Impact of climate change on fungal diversity and biogeography in the Greater Mekong Subregion” (grant no: RDG6130001) for supporting this study. Jing Yang thanks Shaun Pennycook for the corrections to the Latin names. Zong-Long Luo is acknowledged for the help with phylogenetic analyses.

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