Revision of Lophiotremataceae (Pleosporales, Dothideomycetes): Aquasubmersaceae, Cryptocoryneaceae, and Hermatomycetaceae fam. nov.

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Key words

coelomycetes holomorph hyphomycetes systematics

Abstract The family Lophiotremataceae (Pleosporales, Dothideomycetes) is taxonomically revised on the basis of morphological observations and phylogenetic analyses of sequences of nuclear rDNA SSU, ITS, and LSU regions and tef1 and rpb2 genes. A total of 208 sequences were generated from species of Lophiotremataceae and its relatives. According to phylogenetic analyses, Lophiotremataceae encompasses the genus Lophiotrema and five new genera: Atrocalyx, Crassimassarina, Cryptoclypeus, Galeaticarpa, and Pseudocryptoclypeus. These genera are characterised by ascomata with or without a slit-like ostiole and pycnidial conidiomata. Three new families, Aquasubmersaceae, Cryptocoryneaceae, and Hermatomycetaceae, are proposed. Two genera previously recognised as members of Lophiotremataceae, namely, Aquasubmersa having ascomata with a papillate ostiolar neck and pycnidial conidiomata and Hermatomyces possessing sporodochial conidiomata and dimorphic (lenticular and cylindrical) conidia, are included in Aquasubmersaceae and Hermatomycetaceae, respectively. Cryptocoryneum, characterised by the presence of stromatic sporodochia, cheiroid conidia, and conidial arms developed downward from the cap cells, is placed in Cryptocoryneaceae. Two new genera, Antealophiotrema and Pseudolophiotrema, are established, but their familial placements remain unresolved. Antealophiotrema bears ascomata morphologically similar to those of Lophiotrema, but is differentiated from the latter by having ascomata with a well-developed peridium and a monodictys-like asexual morph. Pseudolophiotrema is also similar to Lophiotrema, but can be distinguished by ascomata with a thin peridium. A total of three new families, seven new genera, eight new species, and two new combinations are described and illustrated.

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INTRODUCTION

Lophiotremataceae was originally established as a monotypic family comprising the type genus Lophiotrema (Hirayama & Tanaka 2011). This genus is relatively common, known from throughout the world, and new species are continually being discovered (Mathiassen 1993, Tanaka & Harada 2003, Zhang et al. 2009, Liu et al. 2015, Hyde et al. 2016). Species in Lophiotrema are characterised by immersed ascomata with a crestlike ostiolar neck and cylindrical asci. Although the taxonomic validity of Lophiotremataceae has been confirmed by molecular studies, only a few of its members have been confirmed based on DNA studies (Hyde et al. 2013).

Several major taxonomic and phylogenetic investigations have been carried out to discover missing lineages in Ascomycota (Zhang et al. 2012, Ariyawansa et al. 2015, Liu et al. 2015, Doilom et al. 2016, Hashimoto et al. 2016, Hyde et al. 2016). In these studies, three interesting genera, Aquasubmersa, Cryptocoryneum, and Hermatomyces, were analysed phylogenetically suggesting a close relationship to *Lophiotremataceae*. Aquasubmersa was initially described as a coelomycetous genus (Zhang et al. 2012). Later, a second species of this genus, A. japonicum with sexual and asexual morphs, was reported (Ariyawansa et al. 2015). The asexual genus Hermatomyces is characterised by having sporodochial conidiomata and dimorphic, i.e., lenticular and cylindrical, conidia (Chang 1995). Phylogenetic analyses using sequences of nuclear rDNA small subunit (18S; SSU), internal transcribed spacer (ITS) and large subunit (28S; LSU) regions and translation elongation factor 1-α (tef1) and DNA-directed RNA polymerase II second largest subunit (rpb2) genes have suggested that Aquasubmersa and Hermatomyces are closely related to Lophiotrema (Doilom et al. 2016, Tibpromma et al. 2016). On the basis of their phylogenetic studies, Doilom et al. (2016) and Tibpromma et al. (2016) proposed that Aquasubmersa and Hermatomyces are additional members of Lophiotremataceae. Another genus, Cryptocoryneum, is characterised by having stromatic sporodochia and cheiroid conidia (Schoknecht & Crane 1977). Recently, Hashimoto et al. (2016) assessed the taxonomic placement of Cryptocoryneum species on the basis of morphological observations and the results of a molecular phylogenetic analysis. Although their BLAST search of NCBIs GenBank nucleotide database (http://www.ncbi.nlm.nih.gov/ genbank/) suggested that the genus is also related to Lophiotremataceae (Hashimoto et al. 2016), they could not resolve the familial position of Cryptocoryneum. Many genera related to the family *Lophiotremataceae* still need to be re-evaluated. During our on-going studies of ascomycetous fungi in Japan (Tanaka et al. 2010, 2011, 2015, Hashimoto et al. 2015a, b, 2016), we have collected 57 strains morphologically or phylogenetically related to Lophiotremataceae. The main objectives of the present study were to establish a taxonomic framework within Lophiotremataceae s.lat. and to evaluate the circumscription

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Table 1 Specimens, isolates and new sequence accessions used in this study.

| Species | Family | Original no. | Specimen no. ⁸ | Strain no. | | GenBar | GenBank Accession no. ^b | 0. ^b | |
|--|--|---|----------------------------------|--|--|-----------------------|------------------------------------|-----------------------|-----------------------|
| | | | | | SSU | rsn | tef1 | rpb2 | ITS |
| control of the contro | | | HCCCOC 11 343 | 30000 | 0.000 | 1 0404240 | 1 0404303 | 10404440 | 1 5404474 |
| Americanopinou ema pramieosporam Americanopinou | Adjasjihmersaceae | KT 2813 | HHIJE 30468P | MAFF 245218 | LC194236 | C0615861 | LC194383 | LC 1944 19 | LC0615911 |
| | Agussubmersaceae | KT 2862 | HHIF 30469" | MAFE 2452.19 | LC061581 | 1.00615871 | LC194384 | C194421 | LC061591 |
| | Aguasubmersaceae | KT 2863 | HHUF 30470P | MAFF 245220 | LC0615831 | LC0615881 | LC194385 | LC194422 | LC0615931 |
| Atrocalyx acutisporus | Lophiotremataceae | KT 2436 | HHUF 30504 [⊬] | MAFF 245613 = NBRC 112316 | LC194299 | LC194341 | LC194386 | LC194423 | LC194475 |
| A. lignicola | Lophiotremataceae | I | CBS H-20221 ^H | CBS 122364 | LC194300 | LC194342 | LC194387 | LC194424 | LC194476 |
| Crassimassarina macrospora | Lophiotremataceae | KH 152 | HHUF 30512 ^p | MAFF 245617 | LC194301 | LC194343 | LC194388 | LC194425 | LC194477 |
| | Lophiotremataceae | KT 1764 | HHUF 29084 [⊬] | JCM 13096 = MAFF 239606 | LC194302 | LC194344 | LC194389 | LC194426 | LC194478 |
| Cryptoclypeus oxysporus | Lophiotremataceae | KT 2772 | HHUF 30507 ^H | MAFF 245614 = NBRC 112317 | LC194303 | LC194345 | LC194390 | LC194427 | LC194479 |
| C. ryukyuensis | Lophiotremataceae | AH 342 | HHUF 30510F | MAFF 245616 | LC194304 | LC194346 | LC194391 | LC194428 | LC194480 |
| | Lopniotremataceae | K1 3534 | HHUF 30509" | MAFF 245615 = NBRC 112318 | LC194305 | LC194347 | LC194392 | LC194429 | LC194481 |
| Cryptocoryneum akitaense | Cryptocoryneaceae | K1 3018 | HHUF 30477: | MATE 245365 = NBRC 111758 | LC194306 | LC194348 | LC096136 ² | LC194430 | LC096154* |
| C. brevicondensarum | Cryptocoryneaceae | yone 152 | HHUF 30478" | MATE 245366 = NBRC 111759 | LC194307 | LC194349 | LC096137 ² | LC194431 | LC096155* |
| C. condensatum | Cryptocolyneaceae | I | UP 3 F-632909 | CDS 113838 | LC 194300 | LC194550 | LC096136- | 1049432 | LC090136- |
| | Cryptocoryneaceae | I | I | CBS 122629 | LC194309 | LC194351 | LC096139* | LC194433 | LC0961572 |
| | Cryptocoryneaceae | 1 7 COOC 1-7 | | UBS 122033 MARE 245267 - NBBC 444760 | LC194310 | LC194352 | LC096140 ² | LC194434 | LC096158 ² |
| | Cryptocolyneaceae | 7 2082 7 2084 | 1110F 304795 | MART 245367 - 1467.0 1117.60 | LC 194511 | LC194555 | LC096141 ⁻ | LC 194435 | LC096139- |
| C. Japonicum | Crytocoryneaceae | KT 3291 | HHIIF 30481 | MAFF 245360 MAFF 245369 | LC194312 | LC194354 | LC096142 ² | LC 194436 | LC096161 ² |
| | | KT 3300 | HHI IE 304821 | MAEE 245370 = NBDC 441764 | 1 5194314 | 1 0194356 | 1 00961442 | 1 C 194438 | 1 00061622 |
| | Contocopyneaceae | KT 3413 | HHIJE 30483P | MAFF 245371 | LC194315 | LC194357 | LC096144 | C 194439 | LC096162 |
| | Cryptocoryneaceae | vone 36 | HHUF 30484P | MAFF 245372 | LC194316 | LC194358 | LC096146 ² | LC194440 | LC096164 ² |
| | Cryptocoryneaceae | yone 157 | HHUF 30485 | MAFF 245373 | LC194317 | LC194359 | LC096147 ² | LC194441 | LC096165 ² |
| C. longicondensatum | Cryptocoryneaceae | KT 2913 | HHUF 30486 ^H | MAFF 245374 = NBRC 111762 | LC194318 | LC194360 | LC096148 ² | LC194442 | LC096166 ² |
|) | Cryptocoryneaceae | KT 3487 | HHUF 30487P | MAFF 245375 | LC194319 | LC194361 | LC096149 ² | LC194443 | LC096167 ² |
| C. paracondensatum | Cryptocoryneaceae | KT 3071 | HHUF 30488° | MAFF 245376 | LC194320 | LC194362 | LC096150 ² | LC194444 | LC096168 ² |
| | Cryptocoryneaceae | KT 3241 | HHUF 30489 ^H | MAFF 245377 = NBRC 111763 | LC194321 | LC194363 | LC096151 ² | LC194445 | LC096169 ² |
| C. pseudorilstonei | Cryptocoryneaceae | I | HHUF 30490 [⊬] | CBS 113641 | LC194322 | LC194364 | LC096152 ² | LC194446 | LC096170 ² |
| Cryptocoryneum sp. | Cryptocoryneaceae | I | I | CBS 114518 | LC194323 | LC194365 | LC096153 ² | LC194447 | LC096171 ² |
| Galeaticarpa aomoriensis | Lophiotremataceae | KT 2563 | HHUF 30505 [⊬] | MAFF 245618 = NBRC 112319 | LC194324 | LC194366 | LC194393 | LC194448 | LC194482 |
| Hermatomyces iriomotensis | Hermatomycetaceae | KH 361 | HHUF 30518 ^H | MAFF 245730 = NBRC 112471 | LC194325 | LC194367 | LC194394 | LC194449 | LC194483 |
| H. tectonae | Hermatomycetaceae | KT 2450 | HHUF 30520 | MAFF 245731 | LC194326 | LC194368 | LC194395 | LC194450 | LC194484 |
| | Hermatomycetaceae | AH 276 | HHUF 30521 | MAFF 245732 | LC194327 | LC194369 | LC194396 | LC194451 | LC194485 |
| | Hermatomycetaceae | AH 314 | HHUF 30522 | MAFF 245733 | LC194328 | LC194370 | LC194397 | LC194452 | LC194486 |
| | Hermatomycetaceae | KH 329 | HHUF 30523 | MAFF 245/34 | LC194329 | LC194371 | LC194398 | LC194453 | LC194487 |
| | nemaiomycetaceae | 000 17 | HHUF 30324 | MATH Z45/55 | LC194330 | LC194372 | LC194399 | LC194454 | LC194488 |
| | Hermatomycetaceae | KH 409 | HHUF 30525 | MAFF 245736 MAFF 245737 | LC194331 | LC1943/3 | LC194400 | LC194455 | LC194489 |
| 'I onhiotrema' horage | incertae sedis | 201 | 02000 | CBS 114422 = JCM 14136 | LC 194332 | LC194374 | 1 0194402 | LC194450 | LC 194490 |
| L. eburnoides | Lophiotremataceae | KT 1424-1 | HHUF 30079 ^H | JCM 17826 = MAFF 242970 | LC0017063 | LC0017073 | LC194403 | LC194458 | LC0017093 |
| L. fallopiae | Lophiotremataceae | KT 2748 | HHUF 30506 ^H | MAFF 245612 | LC149911 ⁴ | LC149915 ⁴ | LC194404 | LC194459 | LC149913 ⁴ |
| L. neoarundinaria | Lophiotremataceae | KT 856 | HHUF 27547 | MAFF 239461 | AB524455⁵ | AB524596 ⁵ | AB539109 ⁶ | AB539096 ^e | AB524786 ⁵ |
| | Lophiotremataceae | KT 1034 | HHUF 30015 | NBRC 106239 | AB524457 ⁵ | AB524598 ⁵ | LC194405 | LC194460 | LC194492 |
| | Lophiotremataceae | KT 2200 | HHUF 30014 | NBRC 106238 | AB5244565 | AB524597 ⁵ | AB539110 ⁶ | AB5390976 | AB524787 ⁵ |
| | Lophiotremataceae | KH 17 | HHUF 30511 | MAFF 245619 | LC194334 | LC194376 | LC194406 | LC194461 | LC194493 |
| | Lophiotremataceae | KT 588 | HHUF 27368 | MAFF 245620 | LC194335 | LC194377 | LC194407 | LC194462 | LC194494 |
| | Lophiotremataceae | KT 713 | HHUF 27328 | JCM 17673 | AB618701 ⁷ | AB6190197 | LC194408 | LC194463 | LC194495 |
| elizaria. | Lopniotremataceae | KI /56 | HHUF 2/330 | MATE 23945/ | AB6187027 | AB619020 | LC194409 | LC194464 | LC194496 |
| L. nucula | Lopniotremataceae | 1 2 | L | CBS 627.86 = JCM 14132 | AB6187037 | AB619021 | LC194410 | LC194465 | LC194497 |
| L. vagabundum | Lopniotremaraceae | 7 7 7 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 | HHUF 30077 | JCM 17674 | AB018/04 | AB6190227 | LC194411 | LC194466 | LC194498 |
| | l ophiotremataceae | KT 664 | HHIJE 27323 | MAFF 239456 | AB6187067 | AB619023 | C 194413 | C194468 | C194500 |
| | Lophiotremataceae | KT 3310 | HHUF 30508 | MAFF 245621 | LC194336 | LC194378 | LC194414 | LC194469 | LC194501 |
| | Lophiotremataceae | 1 | F-634236 | CBS 113975 = JCM 14138 | AB6187077 | AB6190257 | LC194415 | LC194470 | LC194502 |
| 'Massarina albocarnis' | Lophiotremataceae | ı | I | CBS 119345 | LC194337 | LC194379 | LC194416 | LC194471 | LC194503 |
| Pseudocryptoclypeus yakushimensis | Lophiotremataceae | KT 2186 | HHUF 30503 ^H | MAFF 245622 = NBRC 112320 | LC194338 | LC194380 | LC194417 | LC194472 | LC194504 |
| Pseudolophiotrema elymicola | incertae sedis | KT 1450 | HHUF 28984 [⊬] | JCM 13090 = MAFF 239600 | LC194339 | LC194381 | LC194418 | LC194473 | LC194505 |
| ^a H = holotype; P = paratype. | | | | | | | | | |
| Sequences obtained in this study are s | hown in bold . Sequences with 1 from, | Ariyawansa et al. (2015), ² | from Hashimoto et al. (2016), 3. | . Sequences obtained in this study are shown in bold . Sequences with 1 from Anyawansa et al. (2015), 2 from Hashimoto et al. (2015), 4 from Liu et al. (2015), 4 from Hyde et al. (2016), 5 from Tanaka et al. (2009), 6 from Schoch et al. (2009), 7 from Hirayama & Tanaka (2011). | ınaka et al. (2009), ⁶ froı | m Schoch et al. (| (2009), ⁷ from Hin | ayama & Tanak | a (2011). |
| | | • | | | | | | , | |

 Table 2
 Specimens, isolates and GenBank accession numbers of species used in the phylogenetic study.

| Species | Family | Specimen no. ^a | Strain no. | | GenBan | GenBank Accession no | | |
|--|----------------------------|---------------------------|------------------------------|----------|----------|----------------------|----------|----------|
| | | | | SSU | rsn | tef1 | rpb2 | ITS |
| Amniculicola immersa | Amniculicolaceae | CBS H-20226 [⊬] | CBS 123083 | GU456295 | FJ795498 | GU456273 | GU456358 | ı |
| A. parva | Amniculicolaceae | CBS H-20227 [⊬] | CBS 123092 | GU296134 | GU301797 | GU349065 | I | ı |
| Anteaglonium abbreviatum | Anteagloniaceae | ANM 925.1 | I | I | GQ221877 | GQ221924 | I | ı |
| A. globosum | Anteagloniaceae | ANM 925.2 ^H | I | I | GQ221879 | GQ221925 | I | I |
| | Anteagloniaceae | SMH 5283 ^P | I | I | GQ221911 | GQ221919 | I | ı |
| A. parvulum | Anteagloniaceae | MFLU 16-0473 | MFLUCC 14-0815 | KU922912 | KU922911 | KU922919 | ı | ı |
| | Anteagloniaceae | MFLU 16-0472 | MFLUCC 14-0817 | KU922914 | KU922913 | ı | I | I |
| | Anteagloniaceae | MFLU 16-0474 | MFLUCC 14-0821 | KU922916 | KU922915 | KU922921 | ı | ı |
| | Anteagloniaceae | MFLU 16-0470 | MFLUCC 14-0823 | KU922918 | KU922917 | KU922922 | ı | ı |
| A. thailandicum | Anteagloniaceae | MFLU 16-0471 ^H | MFLUCC 14-0816 | KU922910 | KU922909 | KU922920 | I | I |
| Aquasubmersa mircensis | Aquasubmersaceae | MFLU 111001 ^H | MFLUCC 11-0401 = IFRDCC 2572 | JX276956 | JX276955 | ı | I | JX276954 |
| Byssolophis sphaerioides | incertae sedis | I | IFRDCC 2053 | GU456296 | GU456318 | GU456263 | GU456348 | ı |
| Hermatomyces tectonae | Hermatomycetaceae | MFLU 15-3437 ^H | MFLUCC 14-1140 | KU712465 | KU764695 | KU872757 | KU712486 | KU144917 |
| | Hermatomycetaceae | MFLU 15-3438P | MFLUCC 14-1141 | KU712466 | KU764696 | KU872758 | I | KU144918 |
| | Hermatomycetaceae | MFLU 15-3439P | MFLUCC 14-1142 | KU712467 | KU764697 | I | KU712487 | KU144919 |
| H. thailandica | Hermatomycetaceae | MFLU 15-3440 [⊬] | MFLUCC 14-1143 | KU712468 | KU764692 | KU872754 | KU712488 | KU144920 |
| | Hermatomycetaceae | MFLU 15-3441P | MFLUCC 14-1144 | KU712469 | KU764693 | KU872755 | KU712489 | KU144921 |
| | Hermatomycetaceae | MFLU 15-3442P | MFLUCC 14-1145 | KU712470 | KU764694 | KU872756 | KU712490 | KU144922 |
| Lepidosphaeria nicotiae | Testudinaceae | I | CBS 101341 | I | DQ678067 | DQ677910 | DQ677963 | 1 |
| Lophiostoma arundinis | Lophiostomataceae | I | CBS 621.86 | DQ782383 | DQ782384 | DQ782387 | DQ782386 | AJ496633 |
| 'Lophiotrema' bambusae | Lophiotremataceae | MFLU 11-0150 | MFLUCC 10-0558 | KX672159 | KX672154 | KX672162 | KX672161 | KX672149 |
| L. crenatum | Lophiostomataceae | I | CBS 629.86 | DQ678017 | DQ678069 | DQ677912 | DQ677965 | 1 |
| Polyplosphaeria fusca | Tetraplosphaeriaceae | HHUF 29399 [⊬] | JCM 13175 = MAFF 239685 | AB524463 | AB524604 | I | I | AB524789 |
| Pseudoastrosphaeriella bambusae | Pseudoastrosphaeriellaceae | MFLU 11-0155 ^H | MFLUCC 11-0205 | KT955455 | KT955475 | KT955437 | KT955414 | ı |
| P. Iongicolla | Pseudoastrosphaeriellaceae | MFLU 11-0207 ^H | MFLUCC 11-0171 | I | KT955476 | KT955438 | KT955420 | 1 |
| P. thailandensis | Pseudoastrosphaeriellaceae | MFLU 11-0145 ^H | MFLUCC 10-0553 | KT955456 | KT955477 | KT955439 | KT955411 | 1 |
| Pseudotetraploa curviappendiculata | Tetraplosphaeriaceae | HHUF 28582 ^н | JCM 12852 = MAFF 239495 | AB524467 | AB524608 | I | I | AB524792 |
| Quadricrura septentrionalis | Tetraplosphaeriaceae | HHUF 28781P | CBS 125429 | AB524474 | AB524615 | I | ı | AB524799 |
| Tetrapla sasicola | Tetraplosphaeriaceae | HHUF 27566 [⊬] | JCM 13167 = MAFF 239677 | AB524490 | AB524631 | ı | ı | AB524807 |
| Triplosphaeria maxima | Tetraplosphaeriaceae | HHUF 29390 [⊬] | JCM 13172 = MAFF 239682 | AB524496 | AB524637 | I | I | AB524812 |
| Ulospora bilgramii | Testudinaceae | I | CBS 101364 | DQ678025 | DQ678076 | DQ677921 | DQ677974 | ı |
| Verruculina enalia | Testudinaceae | ı | BCC 18402 | GU479771 | GU479803 | GU479864 | GU479836 | 1 |
| ^a H = holotype, P = paratype. | | | | | | | | |

of this family based on morphological observations and phylogenetic analyses of SSU, ITS, and LSU nuclear rDNA and *tef1* and *rpb2* gene sequences.

MATERIALS AND METHODS

Isolation and morphological observation

All fungal structures were observed in preparations mounted in distilled water. Morphological characters were observed by differential interference and phase contrast microscopy (Olympus BX53), with images captured with an Olympus digital camera (DP21). A total of 57 single-spore isolates were used for morphological observations and phylogenetic analyses (Table 1). Colony characteristics of cultures grown on potato dextrose agar (PDA; Difco) were observed after 3 wk cultivation at 20 °C in the dark. Colours were noted as described by Rayner (1970). To induce sexual or asexual fructification in culture, 5-mm squares of mycelial agar were placed on water agar including sterilised natural substrate, such as rice straws and pine needles, and the plates were incubated at 20 °C for 2 wk in the dark. When the substrate was colonised, the plates were incubated at 20 °C under blacklight blue illumination for 2 mo to observe sporulation. Cultures were deposited in the Japan Collection of Microorganisms (JCM), the NITE Biological Resource Centre (NBRC), and the GeneBank Project, NARO, Japan (MAFF). Specimens were deposited in the fungus herbarium of Hirosaki University (HHUF).

Phylogenetic analyses

DNA extraction was carried out with an ISOPLANT II kit (Nippon Gene, Japan) in accordance with the manufacturer's protocol. Partial SSU, complete ITS, and partial LSU nuclear rDNA regions and partial tef1 and partial rpb2 genes were amplified by PCR with the primer pairs NS1/NS4, ITS1/ITS4 (White et al. 1990), LR0R/LR7 (Rehner & Samuels 1994, Vilgalys & Hester 1990), EF1-983F/EF1-2218R (Rehner & Buckley 2005), and fRPB2-5F/fRPB2-7cR (Liu et al. 1999), respectively. Amplifications were performed in 25 µL volumes consisting of 2 μ L DNA extract, 2.5 μ L of 10× TEMPase Buffer I, 10 mM dNTP mix, 1 µL of each 20-pM primer, 25 mM MgCl₂, 14.5 µL MilliQ water, and 0.5 µL TEMPase Hot Start DNA polymerase (Ampliqon, Denmark). PCRs were carried out on a PC 320 thermocycler (ASTEC, Japan) as follows: 95 °C for 15 min, followed by 35 cycles of 1 min at 94 °C, 1 min at the designated annealing temperature (42.2 °C for SSU, 61.5 °C for ITS, 46 °C for LSU, 60 °C for tef1, and 58 °C for rpb2), and 1 min at 72 °C, with a final denaturation step of 7 min at 72 °C. The PCR products were sequenced directly at SolGent (South Korea).

Newly generated sequences were deposited in GenBank (Table 1). Sequences of 32 taxa from Amniculicolaceae, Anteagloniaceae, Aquasubmersaceae, Hermatomycetaceae, Lophiotremataceae, Pseudoastrosphaeriellaceae, Testudinaceae, and Tetraplosphaeriaceae were also phylogenetically analysed (Table 2). Lophiostoma crenatum and L. arundinis (Lophiostomataceae) were used as outgroups. All sequences were aligned using the MUSCLE algorithm as implemented in MEGA v. 5 (Tamura et al. 2011). Phylogenetic analyses were conducted using maximum likelihood (ML) and Bayesian methods. The optimum substitution models for each dataset were estimated by Kakusan4 (Tanabe 2011) based on the Akaike information Criterion (AIC; Akaike 1974) for the ML analysis and the Bayesian information Criterion (BIC; Schwarz 1978) for the Bayesian analysis. The ML analysis was performed using TreeFinder Mar 2011 (Jobb 2011) based on the models selected with the AICc4 parameter (proportional model among genes and proportional model among codons), namely, TN93ef+G for SSU, TN93+G for LSU, J2ef+G for ITS, F81+G for the tef1 first codon position,

TIMef+G for the *tef1* second codon position, J2+G for the *tef1* third codon position, J2+G for the rpb2 first codon position, JC69+G for the rpb2 second codon position, and TN93ef+G for the *rpb2* third codon position. Bootstrap proportions (BPs) were obtained by 1 000 bootstrap replications. Bayesian analysis was performed with MrBayes v. 3.2.2 (Ronquist et al. 2012) with substitution models for different regions selected with the BIC4 parameter (i.e., proportional model among loci and among codons): K80+G for SSU, SYM+G for LSU, SYM+G for ITS, F81+G for the tef1 first codon position, GTR+G for the tef1 second codon position, GTR+G for the *tef1* third codon position, GTR+G for the rpb2 first codon position, HKY85+G for the rpb2 second codon position, and SYM+G for the rpb2 third codon position. Two simultaneous, independent Metropolis-coupled Markov chain Monte Carlo (MCMC) runs were performed for 2 M generations with trees sampled every 1 000 generations. Convergence of the MCMC procedure was assessed from the average standard deviation of split frequencies (< 0.01) and effective sample size scores (all > 100) using MrBayes and Tracer v. 1.6 (Rambaut et al. 2014), respectively. The first 25 % of trees were discarded as burn-in, and the remaining trees were used to calculate 50 % majority rule trees and to determine posterior probabilities (PPs) for individual branches. The alignment was submitted to TreeBASE under study number S19310.

The ML and Bayesian phylogenetic analyses were conducted

using an aligned sequence dataset comprising 970 nucleotide

RESULTS

positions from SSU, 1 275 from LSU, 340 from ITS, 909 from tef1, and 1 023 from rpb2. The alignment contained a total of 89 taxa, which consisted of 84 taxa (94.4 %) in SSU, 89 (100 %) in LSU, 71 (79.8 %) in ITS, 81 (91 %) in tef1, 73 (82 %) in rpb2 (Table 1, 2). No significant conflict was observed among individual gene phylogenies, allowing the five genes to be combined into a single dataset. This combined dataset provided higher confidence values for the familial level than did those of the individual gene trees (data not shown). ITS1 was excluded from the analyses because it contained too many ambiguously aligned regions. Of the 4 517 characters included in the alignment, 1 307 were variable and 3 189 were conserved. The ML tree with the highest log likelihood (-31261.0019) is shown in Fig. 1. The Bayesian likelihood score was -31334.0529. The topology recovered by the Bayesian analysis was almost identical to that of the ML tree except for the position of *Anteagloniaceae*. In the phylogenetic tree shown in Fig. 1, Lophiotremataceae forms a moderately supported clade (77 % ML BP/1.00 Bayesian PP) (Fig. 1). The clade corresponding to Lophiotrema s.str. includes L. eburnoides, L. fallopiae, L. neoarundinaria, L. neohysterioides, L. nucula, and L. vagabundum. The results of the phylogenetic analyses suggest that four species previously placed in Lophiotrema, i.e., L. boreale, L. brunneosporum, L. lignicola, and L. bambusae (Mathiassen 1993, Zhang et al. 2009, Hyde et al. 2016), should be excluded from the genus. The first two species, L. boreale and L. brunneosporum, lie outside of Lophiotremataceae, but their familial placements are unresolved (Fig. 1). On the basis of these results, a new genus, Antealophiotrema, is established for L. brunneosporum. Lophiotrema lignicola is transferred to the new genus Atrocalyx, and a new combination, *Atrocalyx lignicola*, is proposed. The generic placement of L. bambusae and L. boreale remains unresolved because we were unable to examine any material of these species.

Four families, including the three new families, *Aquasubmersaceae*, *Cryptocoryneaceae*, and *Hermatomycetaceae*, 10 genera including seven new genera, nine species including seven new species, and two new combinations are described below.

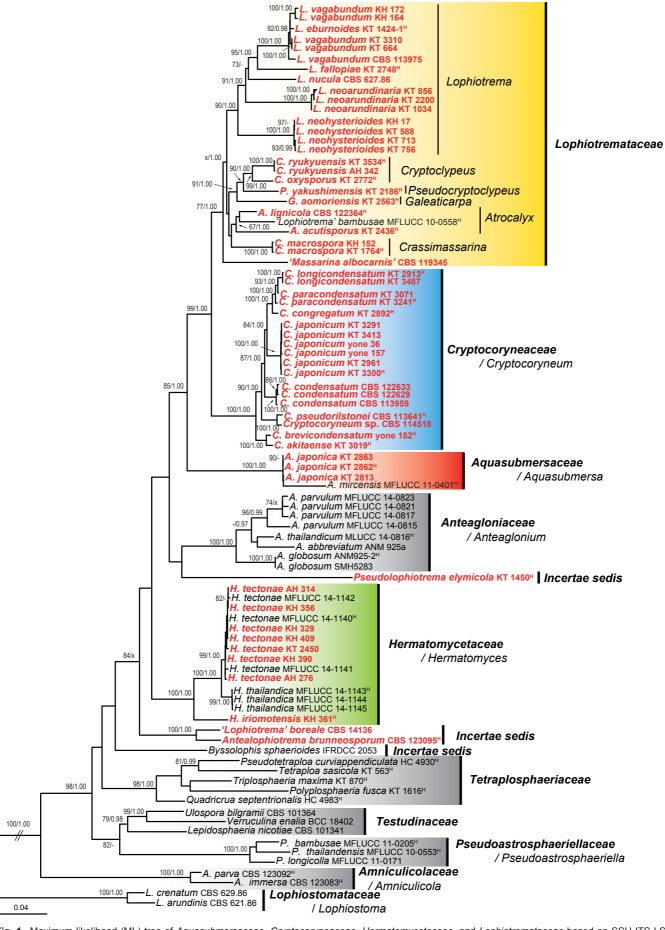


Fig. 1 Maximum-likelihood (ML) tree of *Aquasubmersaceae*, *Cryptocoryneaceae*, *Hermatomycetaceae*, and *Lophiotremataceae* based on SSU-ITS-LSU-*tef1-rpb2* sequences. ML bootstrap proportion (BP) > 60 % and Bayesian posterior probabilities (PP) > 0.95 are presented at the nodes as ML BP/Bayesian PP. A hyphen ('-') indicates values < 60 % BP or 0.95 PP, and a node not present in the Bayesian analysis is shown with 'x'. Ex-holotype strains are indicated with a superscript H. The newly obtained sequences are shown in **bold** and red. The scale bar represents nucleotide substitutions per site.

Aquasubmersaceae A. Hashim. & Kaz. Tanaka, fam. nov. — MycoBank MB819235

Type genus. Aquasubmersa K.D. Hyde & Huang Zhang, Cryptog. Mycol. 33: 340. 2012.

Saprobic on woody plants.

Sexual morph: Ascomata scattered to grouped, subglobose, semi-immersed, with a papillate ostiolar neck. Peridium composed of flattened, thin-walled, polygonal cells. Pseudoparaphyses septate, branched. Asci bitunicate, cylindrical, with a short pedicel, 8-spored. Ascospores hyaline, broadly fusiform with rounded ends, septate.

Asexual morph: Conidiomata pycnidial, globose to subglobose, scattered, semi-immersed, ostiolate. Conidiophores absent. Conidiogenous cells holoblastic, lageniform, hyaline, smooth. Conidia hyaline, ellipsoidal, smooth.

Notes — Previous phylogenetic studies using SSU and LSU rDNA sequences placed Aquasubmersa in Pleosporales (Zhang et al. 2012, Ariyawansa et al. 2015). This genus was treated as a member of Lophiotremataceae in a recent study (Doilom et al. 2016). The results of our phylogenetic analyses based on SSU-ITS-LSU-tef1-rpb2 sequences strongly support (99 % ML BP/1.00 Bayesian PP) this genus as sister to a clade comprising Lophiotremataceae and Cryptocoryneaceae (Fig. 1). To accommodate the genus Aquasubmersa, we introduce a new family, Aquasubmersaceae. Species in Aquasubmersaceae share several common features, such as ascomata having a papillate ostiolar neck and pycnidial conidiomata. Lophiotremataceae differs from Aquasubmersaceae in having ascomata with a compressed, slit-like ostiole. Cryptocoryneaceae and Hermatomycetaceae, whose sexual morphs are still undetermined, can be easily distinguished from Aquasubmersaceae, which has ellipsoidal conidia, because two families have sporodochial conidiomata and cheiroid or lenticular conidia, respectively.

Aquasubmersa K.D. Hyde & Huang Zhang, Cryptog. Mycol. 33: 340. 2012

Type species. Aquasubmersa mircensis Huang Zhang & K.D. Hyde, Cryptog. Mycol. 33: 340. 2012.

Notes — The type species of this genus, *A. mircensis*, was reported only as an asexual morph (Zhang et al. 2012). Ariyawansa et al. (2015) subsequently introduced the second species, *A. japonica*, with sexual and asexual morphs. Both species occurred on submerged woody plants.

Cryptocoryneaceae A. Hashim. & Kaz. Tanaka, fam. nov. — MycoBank MB819237

Type genus. Cryptocoryneum Fuckel, Fungi Rhenani Exsicc. XV-XVI: 25. 1865.

Saprobic on various plants.

Sexual morph: Undetermined.

Asexual morph: Conidiomata sporodochial, pulvinate, often confluent, dark brown to black. Conidiophores arising from stromatic cells, simple, septate, hyaline to pale brown. Conidiogenous cells monoblastic, cylindrical to oblong, terminal, determinate, hyaline to pale brown. Conidia solitary, acrogenous, branched, cheiroid, with dark brown to black cap cells firmly united together, multi-armed; basal cells brown, cuneiform, smooth, thin-walled; arms cylindrical, pale brown, branched at base, smooth, multi-septate.

Notes — *Cryptocoryneum* is characterised by having stromatic sporodochia, cheiroid conidia, and conidial arms that are developed downward from the cap cells (Schoknecht & Crane 1977). This genus was recently taxonomically and phylogeneti-

cally reassessed by Hashimoto et al. (2016). A BLAST search with ITS sequences of species in this genus suggested a close relationship with *Lophiotremataceae*, but their familial position remains unresolved (Hashimoto et al. 2016). According to our results, *Cryptocoryneum* forms a lineage that is phylogenetically distinct from *Lophiotremataceae* (Fig. 1). Although the sexual morph of *Cryptocoryneum* has not been determined, asexual morphs of this genus are phenotypically different from the coelomycetous asexual morph of *Lophiotremataceae* (Leuchtmann 1985, this study). We therefore introduce the new family *Cryptocoryneaceae* to accommodate the genus *Cryptocoryneum*.

Cryptocoryneum Fuckel, Fungi Rhenani Exsicc. XV–XVI: 25. 1865

Type species. Cryptocoryneum fasciculatum Fuckel, Fungi Rhenani Exsicc. XV–XVI: 25. Fung. Rhen. 1527. 1865. (= Cryptocoryneum hysterioides (Corda) Peyronel, Nuovo Giorn. Bot. Ital. 25: 449. 1918.)

Notes — For further information on this genus, see Hashimoto et al. (2016).

Hermatomycetaceae Locq. ex A. Hashim. & Kaz. Tanaka, fam. nov. — MycoBank MB819238

Hermatomycetaceae Locq., Mycologie générale et structural: 202. 1984. nom. inval. (Art. 36.1, Melbourne Code).

Type genus. Hermatomyces Speg., Anales Mus. Nac. Buenos Aires, ser. 3, 13: 445. 1911.

Saprobic on various plants.

Sexual morph: Undetermined.

Asexual morph: Conidiomata sporodochial, pulvinate, dark brown to black. Conidiophores mononematous, pale brown. Conidiogenous cells monoblastic, integrated, terminal, cylindrical. Conidia dimorphic; lenticular conidia ellipsoidal, muriform; cylindrical trans-septate, hyaline to brown.

Notes — The family *Hermatomycetaceae* sensu Locq. was informally proposed as a provisional name (Locquin 1984). Hermatomycetaceae is established here to accommodate the single genus *Hermatomyces*. Two of the most striking features of Hermatomyces are the sporodochial conidiomata and the dimorphic conidia, the latter existing in lenticular and cylindrical forms (Chang 1995). As a sexual morph of this genus has not been reported and a phylogenetic study has not been performed, this genus has long been treated as 'incertae sedis' within Ascomycota (Wijayawardene et al. 2012). Doilom et al. (2016) and Tibpromma et al. (2016) have suggested that this genus belongs to Lophiotremataceae on the basis of their phylogenetic analyses using SSU, LSU, tef1, and rpb2. Our analyses using additional species of Lophiotremataceae and its related taxa revealed that Hermatomyces is distantly related to Lophiotremataceae (Fig. 1).

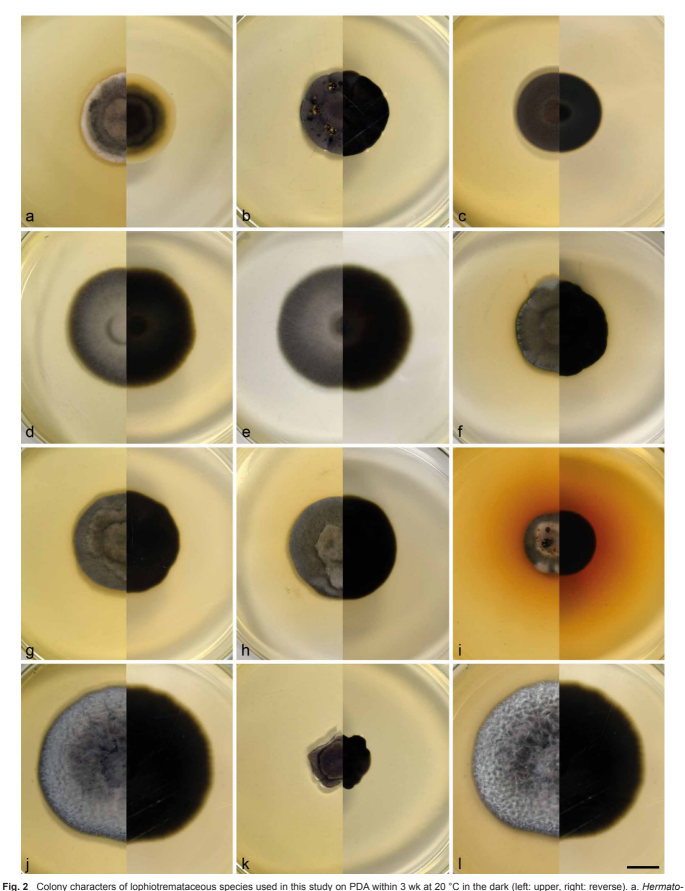
Hermatomyces Speg., Anales Mus. Nac. Buenos Aires, ser. 3, 13: 445. 1911

Type species. Hermatomyces tucumanensis Speg., Anales Mus. Nac. Buenos Aires, ser. 3, 13: 446. 1911.

Saprobic on various plants.

Sexual morph: Undetermined.

Asexual morph: Conidiomata sporodochial, pulvinate, often confluent, dark brown to black. Conidiophores mononematous, septate, pale brown. Conidiogenous cells monoblastic, integrated, terminal, cylindrical. Conidia dimorphic; lenticular conidia ellipsoidal, muriform, dark brown to black at central cells, pale brown at peripheral cells; cylindrical conidia composed of black peripheral cells and dark brown upper cells, hyaline at lower cells.



myces iriomotensis (MAFF 245730 = NBRC 112471, ex-holotype culture); b. Atrocalyx acutisporus (MAFF 245613 = NBRC 112316, ex-holotype culture); c. A. lignicola (CBS 122364, ex-holotype culture); d. Crassimassarina macrospora (JCM 13096 = MAFF 239606, ex-holotype culture); e. C. macrospora (MAFF 245617, ex-paratype culture); f. Cryptoclypeus oxysporus (MAFF 245614 = NBRC 112317, ex-holotype culture); g. C. ryukyuensis (MAFF 245615 = NBRC 112318, ex-holotype culture); h. C. ryukyuensis (MAFF 245616, ex-paratype culture); i. Galeaticarpa aomoriensis (MAFF 245618 = NBRC 112319, ex-holotype culture); j. Pseudocryptoclypeus yakushimensis (MAFF 245622 = NBRC 112320, ex-holotype culture); k. Antealophiotrema brunneosporum (CBS 123095, ex-holotype culture); l. Pseudolophiotrema elymicola (JCM 13090 = MAFF 239600, ex-holotype culture). — Scale bar = 1 cm.

Notes — This genus was established by Spegazzini (1911) to accommodate *H. tucumanensis*. Since its establishment, 10 additional taxa have been described worldwide (Spegazzini 1911, Hughes 1953, Rao & De Hoog 1986, Castañeda-Ruiz & Heredia 2000, Leão-Ferreira et al. 2013, Prasher & Prasher 2014, Doilom et al. 2016, Tibpromma et al. 2016). Most species of this genus occur on angiosperms and monocots, with a few rarely reported from ferns (Castañeda-Ruiz & Heredia 2000) or gymnosperms (Mel'nik 2000). The most widespread species in the genus, *H. sphaericus*, has been reported from Africa, Eurasia, and Central and South America (Spegazzini 1911, Hughes 1953, Chang 1995, Matsushima 1993, Mel'nik 2000, Barbosa & Gusmão 2011).

Hermatomyces iriomotensis A. Hashim. & Kaz. Tanaka, sp. nov. — MycoBank MB819239; Fig. 2a, 3

Etymology. Referring to the collection site.

Saprobic on woody plants. Sexual morph: Undetermined.

Asexual morph: Conidiomata sporodochial, pulvinate, often confluent, dark brown to black. Conidiophores mononematous, short, pale brown, smooth, $2.5-3~\mu m$ wide, $17.5-22~\mu m$ high, sometimes reduced to conidiogenous cells. Conidia dimorphic, lenticular and cylindrical. Lenticular conidia solitary, elliptical to almost round in one plane, smooth, muriform, with pale peripheral cells surrounding central dark brown to black cells, $30-36\times20-27~\mu m$ (av. = $33.6\times23.5~\mu m$, n = 50), $18-24~\mu m$ thick (av. = $21.9~\mu m$, n = 30) in lateral view. Cylindrical conidia

septa, $20.5-33 \times 7-12.5 \ \mu m$ (av. = $28.7 \times 9.6 \ \mu m$, n = 20). Culture characteristics — Colonies on PDA attaining 24–28 mm diam within 21 d at 20 °C in the dark, floccose, centrally

straight to curved, with one to two columns, 3-7-septate, hya-

line, sometimes pale brown at apical cell, constricted at the

raised, straw (46: Rayner 1970) to grey olivaceous (107); reverse smoke grey (105) to grey olivaceous (107) (Fig. 2a); no sporulation observed.

Specimen examined. Japan, Okinawa, Isl. Iriomote, near Tropical botanic garden, on dead twigs of woody plant, 13 July 2011, *K. Hirayama & K. Tanaka*, KH 361 (HHUF 30518 holotype designated here, ex-holotype living culture MAFF 245730 = NBRC 112471).

Notes — Lenticular conidia of this species resemble those of *H. uniseriatum*, but *H. iriomotensis* can be distinguished from the latter species by the presence of 3–7-septate cylindrical conidia arranged in 1–2 columns (vs 2–3-septate and one row in the latter; Leão-Ferreira et al. 2013). *Hermatomyces iriomotensis* differs from *H. krabiensis*, which has cylindrical conidia with swollen lower cells (Tibpromma et al. 2016). Additionally, ITS sequences between these two taxa differed at seven positions.

Lophiotremataceae K. Hiray. & Kaz. Tanaka, Mycoscience 52: 405. 2011

Type genus. Lophiotrema Sacc., Michelia 1 (no. 3): 338. 1878.

Saprobic on various plants.

Sexual morph: Ascomata immersed, erumpent at the apex, subglobose. Ostiolar neck crest-like or rarely papillate, mostly elongated and laterally compressed. Peridium composed of rectangular to globose cells. Pseudoparaphyses septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, with a short stipe, 8-spored. Ascospores fusiform to broadly fusiform, hyaline, smooth.

Asexual morph: Conidiomata pycnidial, globose to subglobose, scattered, semi-immersed, ostiolate. Peridium composed of subglobose to angular, brown cells. Conidiophores absent. Conidiogenous cells holoblastic or phialidic, cylindrical to ampliform, hyaline. Conidia ellipsoidal to cylindrical with rounded ends, hyaline, smooth, aseptate or multi-septate.

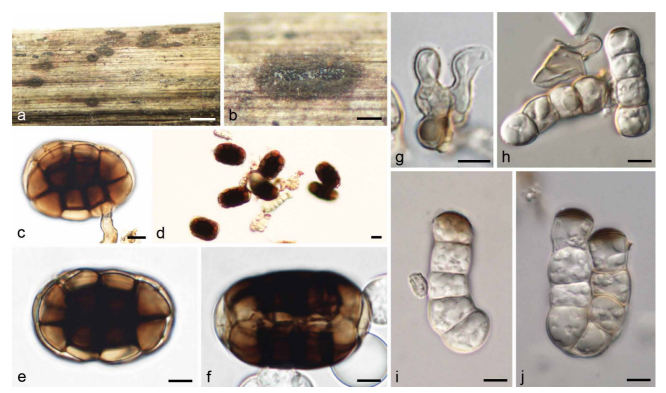


Fig. 3 Hermatomyces iriomotensis. a-b. Conidiomata on substrate; c. conidiogenous cell and immature lenticular conidium; d-f. lenticular conidia (f. lateral view); g. conidiogenous cells and immature cylindrical conidium; h-j. cylindrical conidia (all: HHUF 30518, holotype). — Scale bars: a=1 mm; b=200 μ m; c, e-j=5 μ m; d=10 μ m.

Notes — As originally circumscribed *Lophiotremataceae* was a monotypic family comprising the genus *Lophiotrema* (Zhang et al. 2009, Hirayama & Tanaka 2011, Hyde et al. 2013). A somewhat broader familial concept for *Lophiotremataceae* was adopted by Doilom et al. (2016) and Tibpromma et al. (2016), who considered the family to comprise *Aquasubmersa*, *Hermatomyces*, and *Lophiotrema* on the basis of phylogenetic studies. However, the results of our morphological examination and phylogenetic analyses using SSU, ITS, LSU, *tef1*, and *rpb2* sequences suggest that this family encompasses *Lophiotrema* and five new genera.

Because Aquasubmersa and Hermatomyces were placed outside of Lophiotremataceae in our phylogenetic tree (Fig. 1), we treat these genera as belonging to the families Aquasubmersaceae and Hermatomycetaceae, respectively. One species of Lophiotrema (L. lignicola) grouped with Atrocalyx, a new genus in Lophiotremataceae, while two species (Lophiotrema boreale and L. brunneosporum) were placed outside of Lophiotremataceae entirely.

Marincowitz et al. (2008) have suggested that 'Massarina albocarnis' (CBS 119345) has a phylogenetic affinity with Lophiotrema based on BLAST results involving ITS and LSU sequences. However, Beier et al. (2015), who observed the holotype specimen of M. albocarnis, has indicated that this species belongs to Diaporthe (Sordariomycetes). The isolate CBS 119345 may thus be misidentified. Unfortunately, we were unable to examine any morphological features of CBS 119345 because it did not sporulate in culture.

Lophiotrema Sacc., Michelia 1 (no. 3): 338. 1878

Type species. Lophiotrema nucula (Fr.) Sacc., Michelia 1 (no. 3): 338. 1878.

Notes — For further information on this genus, see Holm & Holm (1988), Tanaka & Harada (2003), Zhang et al. (2009), and Hirayama & Tanaka (2011). In the present study, we found that *Lophiotrema* s.str. should be limited to species having ascomata with a slit-like ostiole and an ascomatal wall of uniform thickness, asci with a short stipe, and pycnidial asexual morphs. Although *L. brunneosporum* with a monodictys-like asexual morph was sister to '*Lophiotrema*' boreale in our phylogenetic tree (Fig. 1), these two taxa were distinct from *Lophiotremataceae* s.str. (Fig. 1). Consequently, they should be treated as distinct lineages in *Dothideomycetes*. We were unable to morphologically examine *L. boreale* from an isolate of this species (CBS 114422) or the original specimen used for isolation. Further examination is required to clarify the taxonomic placement of this species.

Atrocalyx A. Hashim. & Kaz. Tanaka, gen. nov. — MycoBank MB819240

Etymology. From the Latin atro-, meaning black, and calyx, meaning cap.

Type species. Atrocalyx acutisporus A. Hashim. & Kaz. Tanaka.

Saprobic on woody plants.

Sexual morph: Ascomata solitary to grouped, semi-immersed to immersed. Ostiolar neck crest-like, elongated and laterally compressed, surrounded by dark brown hyphae. Peridium composed of 2 zones at side. Pseudoparaphyses septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 8-spored. Ascospores broadly fusiform, hyaline, 1-septate, smooth.

Asexual morph: Conidiomata pycnidial, globose to subglobose, superficial, black, ostiolate. Peridium composed of elongated, brown cells. Conidiophores absent. Conidiogenous cells holoblastic, ampliform to cylindrical, hyaline. Conidia ellipsoidal, hyaline, smooth, aseptate.

Notes — The new genus *Atrocalyx* is established to accommodate *A. lignicola* (formerly *L. lignicola*) and a new species, *A. acutisporus*. These two species are characterised by a crest-like, elongated and laterally compressed ostiolar neck (Fig. 4b, 5b) surrounded by a well-developed peridium (up to 62.5 µm) (Fig. 4c, 5c). *'Lophiotrema' bambusae*, which was recently introduced as a species in *Lophiotrema* (Hyde et al. 2016), was nested within a moderately supported clade (67 % ML BP/1.00 Bayesian PP) along with these two species in our phylogenetic tree (Fig. 1). Because we could not observe any material of *'Lophiotrema' bambusae*, the generic placement of this species is pending.

The genus is morphologically similar to *Lophiotrema*, but can be distinguished from the latter by its well-developed peridium around the ostiolar neck and base (vs a poorly developed peridium up to $25 \mu m$ thick; Holm & Holm 1988).

Atrocalyx acutisporus A. Hashim. & Kaz. Tanaka, sp. nov. — MycoBank MB819241; Fig. 2b, 4

Etymology. Referring to the ascospores with acute ends.

Saprobic on dead twigs of woody plants.

Sexual morph: Ascomata ellipsoidal, solitary to 2-4-grouped, immersed, 110–140 µm high, 190–210 µm diam. Ostiolar neck crest-like, elongated and laterally compressed, surrounded by dark brown hyphae. Peridium 20-30 µm thick at side, composed of 2 zones; outer zone 11-17 µm thick, composed of elongated, thin-walled, $8.5-12.5 \times 2.5-3 \mu m$, brown cells; inner zone 15-17 µm thick, composed of globose to rectangular, $7.5-9 \times 4-5 \mu m$, hyaline cells; near the ostiole 37.5-45 μm thick, composed of globose, brown to black cells; 19-37.5 µm thick at the base, composed of globose to rectangular, 1.8–2.5 µm diam cells. Pseudoparaphyses numerous, 1–1.5 µm wide, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, $(66.5-)75-89.5 \times 8-11 \mu m$ (av. = 80.8 \times 9.2 μ m, n = 10), with a short stipe (4–7.5 μ m long, av. = 5.5 μ m, n = 10), apically rounded with an ocular chamber, 8-spored. Ascospores broadly fusiform with acute ends, 13.5–18(–20) × $3-4(-5.5) \mu m$ (av. = $15.9 \times 3.7 \mu m$, n = 50), I/w 3.7-5.1 (av. = 4.4, n = 50), hyaline, with a septum nearly median (0.44–0.59, av. = 0.51, n = 50), slightly constricted at the septum, smooth, with an entire gelatinous sheath up to 2 µm thick.

Asexual morph: Conidiomata pycnidial, globose to subglobose, up to 145 μm high, 60-130 μm diam, 2-3-grouped, superficial, black, with a papillate ostiolar neck. Peridium 7.5–10 μm thick, composed of 3-4 layers of $7.5-13 \times 1.5-3$ μm, elongated, brown cells. Conidiophores reduced to conidiogenous cells. Conidiogenous cells holoblastic, $7.5-12 \times 1.5-3$ μm, ampliform to cylindrical, hyaline, smooth. Conidia ellipsoidal, $3-4 \times 1.9-2$ μm (av. = 3.5×2.0 μm, n = 50), l/w 1.5-2.1 (av. = 1.8, n = 50), hyaline, smooth, aseptate, guttulate when young.

Culture characteristics — Colonies on PDA attaining 24–25 mm diam within 21 d at 20 °C in the dark, floccose, radiately, smoke grey (105); reverse olivaceous grey to olivaceous black (108) (Fig. 2b); asexual morph formed.

Specimen examined. Japan, Okinawa, Isl. Iriomote, Sono trail, on dead twigs of woody plant, 29 Sept. 2007, K. Tanaka & H. Yonezawa, KT 2436 (HHUF 30504 holotype designated here, ex-holotype living culture MAFF 245613 = NBRC 112316).

Notes — *Atrocalyx acutisporus* superficially resembles *A. lignicola*, but can be distinguished from the latter by its smaller ascospores $(13.5-18(-20)\times3-4(-5.5)\ \mu m$ vs $20-26\times4-5.5(-6)\ \mu m$, respectively). ITS sequence differences between these two species were found at 16 of 529 nucleotide positions, with two gaps.

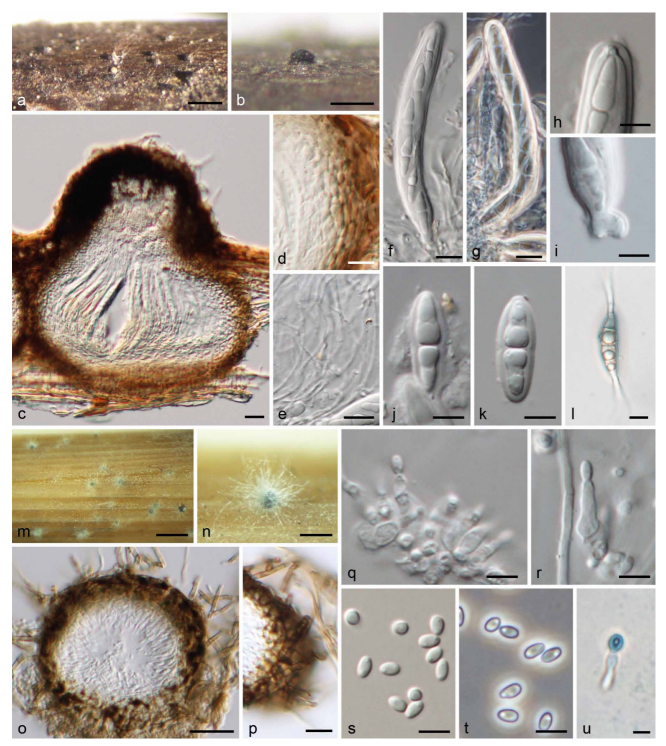


Fig. 4 Atrocalyx acutisporus. a-b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma; e. pseudoparaphyses; f-g. asci; e. ascus apex; e. ascus stipe; e. ascospores; e. germinating ascospore; e. n. conidiomata in culture; e. conidioma in longitudinal section; e. peridium of conidioma; e. conidiogenous cells; e. conidia; e. germinating conidium (e-e): HHUF 30504, holotype; e0. MAFF 245613, ex-holotype culture). — Scale bars: e0. e1. e2. e2. e3. e4. e4. e6. e9. e9.

Atrocalyx lignicola (Ying Zhang, J. Fourn. & K.D. Hyde)
A. Hashim. & Kaz. Tanaka, comb. nov. — MycoBank MB819242; Fig. 2c, 5

Basionym. Lophiotrema lignicola Ying Zhang, J. Fourn. & K.D. Hyde, Fung. Diversity 38: 238. 2009.

Saprobic on dead twigs of Populus sp.

Sexual morph: Ascomata ellipsoidal, 330–380 μ m high, 350–600 μ m long, 230–400 μ m wide, solitary to 4–5-grouped, semi-immersed. Ostiolar neck crest-like, elongated and laterally compressed, surrounded by dark brown hyphae. Peridium 27.5–42 μ m thick at side, composed of 2 zones; outer zone 15–20 μ m thick, composed of elongated, thin-walled, 10–12.5 \times

3.5–5 µm, brown cells; inner zone 15–22 µm thick, composed of globose to rectangular, 5–6.5 µm diam, hyaline cells; near the ostiole 57–62.5 µm thick, composed of dark brown cells; at base 35–46 µm thick, composed of globose to rectangular, 2.5–4(–6) µm diam, brown to black cells. *Pseudoparaphyses* numerous, 1–1.5 µm wide, septate, branched and anastomosed. *Asci* bitunicate, fissitunicate, cylindrical, 100–146 × 12.5–17 µm (av. = 125.2 × 13.4 µm, n = 11), with a short stipe (4–18 µm long, av. = 8.4 µm, n = 11), apically rounded with an ocular chamber, 8-spored. *Ascospores* broadly fusiform with rounded ends, 20–26 × 6.5–9.5 µm (av. = 22.3 × 7.4 µm, n = 50), hyaline, with a septum nearly median (0.45–0.54, av. = 0.50, n = 50), slightly constricted at the septum, smooth, with an

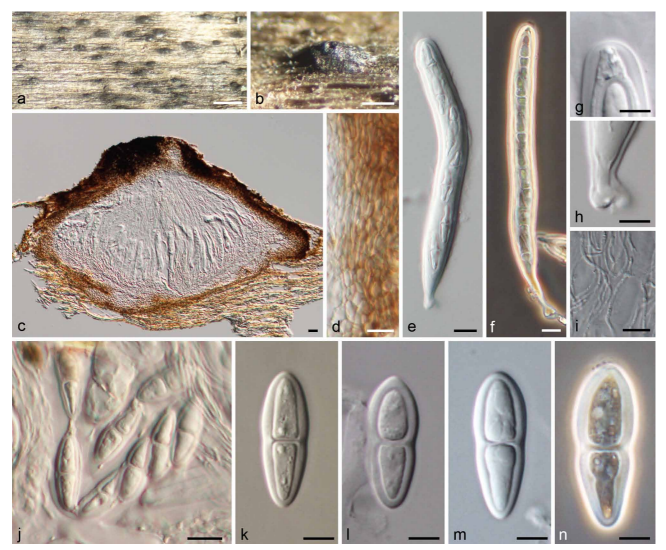


Fig. 5 Atrocalyx lignicola. a – b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma; e – f. asci; g. ascus apex; h. ascus stipe; i. pseudoparaphyses; j – n. ascospores (all: CBS H-20221, holotype). — Scale bars: a = 1 mm; b = 250 μm; c = 20 μm; d – f, i = 10 μm; g, h, j – n = 5 μm.

entire gelatinous sheath up to 2 μm thick. Senescent ascospores 3-septate, yellowish.

Asexual morph: Undetermined.

Culture characteristics — Colonies on PDA attaining 23–26 mm diam within 21 d at 20 °C in the dark, velvety, plane, smoke grey (105); reverse olivaceous black (108) (Fig. 2c); no sporulation observed.

Specimen examined. Belgium, Hainaut, Orval, ruisseau de Williers, on decorticated trunk of *Populus* sp., 29 Sept. 2006, *J. Fournier* (CBS H-20221 holotype, ex-holotype living culture CBS 122364).

Notes — *Atrocalyx lignicola* was first described as a species of *Lophiotrema* on the basis of its 1-septate, hyaline, strongly constricted ascospores (Zhang et al. 2009). According to our morphological observations of the holotype specimen, however, this species is not typical for the genus *Lophiotrema*; in particular, *A. lignicola* possesses ascomata with well-developed peridium (Fig. 5c).

Crassimassarina A. Hashim. & Kaz. Tanaka, gen. nov. — Myco-Bank MB819243

Etymology. After its morphological similarity to Massarina, but with well-developed ascomatal wall.

Type species. Crassimassarina macrospora A. Hashim. & Kaz. Tanaka. Saprobic on dead twigs of woody plants.

Sexual morph: Ascomata solitary to grouped, immersed to erumpent, subglobose. Ostiolar neck papillate, without slit-like ostiole, composed of carbonaceous, thick-walled, black cells. Peridium composed of rectangular, thin-walled cells. Pseudoparaphyses trabeculate, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 8-spored. Ascospores broadly fusiform with rounded ends, straight, 1-septate, hyaline, smooth.

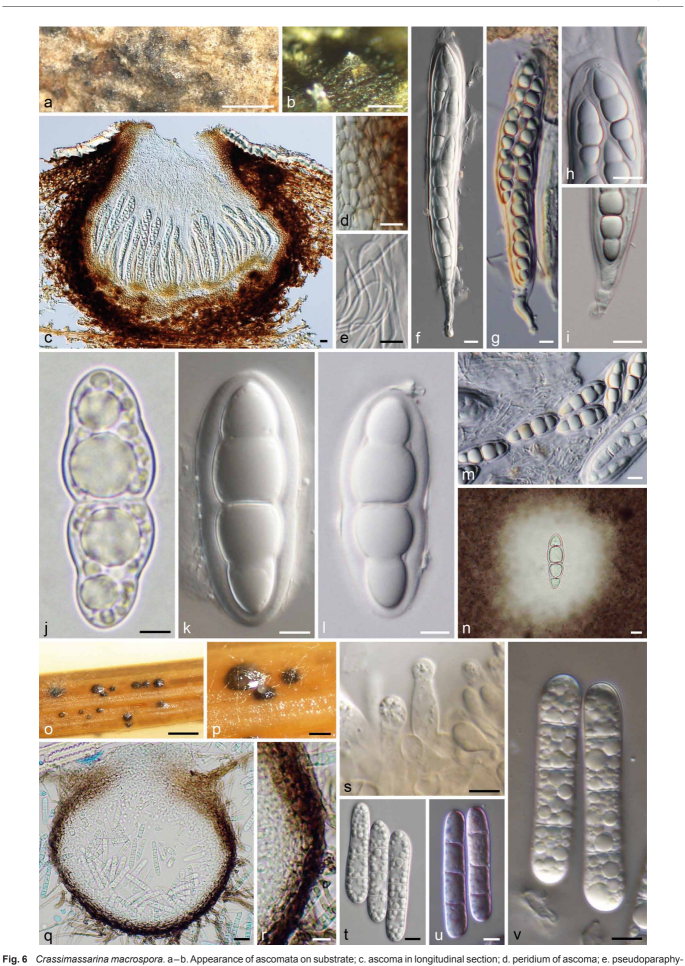
Asexual morph: Conidiomata pycnidial, globose to subglobose, scattered, semi-immersed, solitary, black, ostiolate. Peridium composed of subglobose to rectangular, brown cells. Conidiophores absent. Conidiogenous cells holoblastic, cylindrical, hyaline. Conidia cylindrical with rounded ends, hyaline, smooth, multi-septate.

Crassimassarina macrospora A. Hashim. & Kaz. Tanaka, *sp. nov.* — MycoBank MB819244; Fig. 2d-e, 6

Etymology. Referring to the large ascospores.

Saprobic on dead twigs of Cornus controversa.

Sexual morph: Ascomata solitary to 4–5-grouped, immersed, erumpent at the apex, subglobose in section, $450-620~\mu m$ high, $380-700~\mu m$ diam. Ostiolar neck papillate, without slit-like ostiole, composed of carbonaceous, thick-walled, black cells. Peridium $25-60~\mu m$ thick, composed of 6–8 layers of rectangular, thin-walled, $5-12.5~\mu m$ diam cells, surrounded by brown



ses; f-g. asci; h. ascus apex; i. ascus stipe; j-m. ascospores; n. ascospore with a gelatinous sheath (in India ink); o-p. conidiomata in culture; q. conidioma in longitudinal section; r. peridium of conidioma; s. conidiogenous cells; t-v. conidia (u in Trypan Blue) (a, c-f, j-k, n: HHUF 29084, holotype; b, g-i, l-m: HHUF 30512, paratype; o-s, u-v: JCM 13096 = MAFF 239606, ex-holotype culture; t: MAFF 245617, ex-paratype culture). — Scale bars: a, o = 1 mm; b, p = 250 μ m; c, q = 20 μ m; d-g, m-n, r = 10 μ m; h-l, s-v = 5 μ m.

hyphae (2–3 µm thick). *Pseudoparaphyses* numerous, trabeculate, 1–1.5 µm wide, septate, branched and anastomosed. *Asci* numerous, bitunicate, fissitunicate, cylindrical, 165–200 \times 25–33 µm (av. = 183.5 \times 27.9 µm, n = 7), with a short stipe (7.5–26 µm long, av. = 16.8 µm, n = 7), apically rounded with an ocular chamber, 8-spored. *Ascospores* broadly fusiform with rounded ends, straight, (29.5–)33–42 \times 9–15 µm (av. = 37.8 \times 13.0 µm, n = 55), I/w 2.1–4.0 (av. = 2.9, n = 55), 1(–3)-septate, with a submedian primary septum (0.51–0.58 (–0.71), av. = 0.53, n = 55), strongly constricted at the septum and midpoints of each cell, hyaline, smooth, guttulate when young, with an entire gelatinous sheath (1–3 µm wide at sides).

Asexual morph: Conidiomata pycnidial, globose to subglobose, up to 240 μm high, 190–250 μm diam, scattered, semi-immersed, solitary, black, with a papillate ostiolar neck. Peridium 15–20 μm thick, composed of 4–6 layers of 4–7 μm diam, subglobose to rectangular, brown cells. Conidiophores reduced to conidiogenous cells. Conidiogenous cells holoblastic, 6–14 \times 3–5 μm, cylindrical, hyaline, smooth. Conidia cylindrical with rounded ends, (32–)36–55(–58) \times 6–8 μm (av. = 44.1 \times 7.2 μm, n = 50), l/w 4.7–8.1 (av. = 6.2, n = 50), hyaline, smooth, 3(–7)-septate, guttulate when young.

Culture characteristics — Colonies on PDA attaining 36–40 mm diam within 21 d, velvety, plane, smoke grey (105), grey olivaceous (107) at margin; reverse olivaceous black (108) (Fig. 2d–e); asexual morph formed.

Specimens examined. Japan, Ibaraki, Tsukuba, Amakubo, Tsukuba botanical garden, on dead twigs of Cornus controversa, 20 Nov. 2004, Y. Ooki, KT 1764 (HHUF 29084 holotype designated here, ex-holotype living culture JCM 13096 = MAFF 239606); ibid., KT 1765 (HHUF 29085 paratype); Aomori, Minamitsugaru, Owani, on dead twigs of woody plant, 28 June 2008, K. Hirayama & K. Tanaka, KH 152 (HHUF 30512 paratype, ex-paratype living culture MAFF 245617).

Notes — This genus is morphologically similar to Massarina and genera in Pleomassariaceae s.lat. (Barr 1982, Tanaka et al. 2005, 2015) in having large, immersed ascomata with a short papillate ostiolar neck and relatively large ascospores. However, Crassimassarina is different from Massarina in having a well-developed ascomatal wall; it differs from pleomassariaceous genera in having trabeculate pseudoparaphyses and hyaline ascospores. The asexual morph of Crassimassarina resembles that of Stagonospora in regards to its pycnidial conidiomata, conidiophores with reduced conidiogenous cells, and multi-septate cylindrical conidia; however, the latter genus differs from Crassimassarina in having phialidic conidiogenous cells (Quaedvlieg et al. 2013, Tanaka et al. 2015). Massarina and Stagonospora (Pleosporales, Massarinaceae) are phylogenetically distinct lineages from Lophiotremataceae containing Crassimassarina.

Crassimassarina can be distinguished from other genera in Lophiotremataceae by its ascomata lacking a slit-like ostiole, an ascomatal peridium composed of carbonaceous cells (Fig. 6c), and multi-septate, large conidia (Fig. 6t–v).

ITS sequences of the two examined isolates of *C. macrospora* differed at only two positions, with two gaps. Morphological features and culture characteristics of these isolates were completely identical (Fig. 2d-e, 6k-l).

Cryptoclypeus A. Hashim. & Kaz. Tanaka, gen. nov. — Myco-Bank MB819245

 $\ensuremath{\textit{Etymology}}.$ Referring to the ascomata covered by a less-developed clypeus.

Type species. Cryptoclypeus ryukyuensis A. Hashim. & Kaz. Tanaka.

Saprobic on dead twigs of bamboo.

Sexual morph: Ascomata solitary to grouped, immersed. Ostiolar neck crest-like, elongated, laterally compressed, with a slit-like ostiole and less-developed clypeus. *Peridium* composed of rectangular, thin-walled, pale brown cells. *Pseudoparaphyses* septate, branched and anastomosed. *Asci* bitunicate, fissitunicate, cylindrical, 8-spored. *Ascospores* fusiform, multiseptate, hyaline, smooth.

Asexual morph: Conidiomata pycnidial, globose to subglobose, grouped, immersed, ostiolate. Peridium composed of subglobose to rectangular, brown cells. Conidiophores absent. Conidiogenous cells phialidic, ampliform to cylindrical. Conidia cylindrical with slightly angular ends, hyaline, smooth, 1-septate.

Notes — Two species of *Cryptoclypeus* share common characteristics, e.g., ascomata with a less-developed clypeus (up to 450 μ m wide), and a peridium composed of rectangular cells (Fig. 7c–d, 8c–d). In the phylogenetic tree, they formed a well-supported clade in *Lophiotremataceae* (99 % ML BP/1.00 Bayesian PP) (Fig. 1).

The sexual morph of *Cryptoclypeus* is similar to that of *Tetraploa* (*Tetraplosphaeriaceae*), which also has ascomata with a clypeus and narrowly fusiform ascospores (Tanaka et al. 2009). *Cryptoclypeus*, however, can be distinguished from the latter genus by having a well-developed neck with a slit-like ostiole. The asexual morph of *Cryptoclypeus* superficially resembles that of *Bambusicola* (*Bambusicolaceae*), but is easily distinguishable from the latter by having phialidic conidiogenous cells rather than annellidic ones (Dai et al. 2012).

Cryptoclypeus oxysporus A. Hashim. & Kaz. Tanaka, sp. nov.
— MycoBank MB819246; Fig. 2f, 7

Etymology. Referring to the sharp ascospores.

Saprobic on dead culm of Sasa sp.

Sexual morph: Ascomata ellipsoidal, solitary, immersed, 245–310 μm high, 180–240 μm diam. Ostiolar neck crest-like, elongated, laterally compressed, with less-developed clypeus (230–320 μm wide). Peridium uniform, 15–17.5 μm thick, composed of rectangular, thin-walled, 6.5–14 × 2.5–4.5 μm, pale brown cells. Pseudoparaphyses numerous, 0.5–1 μm wide, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 71–100 × 6–9 μm (av. = 85.8 × 7.7 μm, n = 10), with a short stipe (5.5–11 μm long, av. = 7.8 μm, n = 10), apically rounded with an ocular chamber, 8-spored. Ascospores fusiform with acute ends, straight, 19–27 × 3–4.5 μm (av. = $23.3 \times 3.2 \, \mu m$, n = 50), l/w 5.5–8.9 (av. = 7.2, n = 50), 3-septate, slightly constricted at the primary septum nearly median (0.46–0.55, av. = 0.50, n = 50), hyaline, smooth. Asexual morph: Undetermined.

Culture characteristics — Colonies on PDA attaining 24–28 mm diam within 21 d at 20 °C in the dark, velvety, radiately, centrally raised, smoke grey (105) to grey olivaceous (107); reverse olivaceous black (108) (Fig. 2f); no sporulation observed.

Specimen examined. Japan, Iwate, Hanamaki, near Dai spa, on dead culm of Sasa sp., 25 June 2011, K. Tanaka, KT 2772 (HHUF 30507 holotype designated here, ex-holotype living culture MAFF 245614 = NBRC 112317).

Notes — This species can be distinguished from *C. ryukyuensis* by its slightly larger ascospores with acute ends (Fig. 7k–n) (19–27 \times 3–4.5 µm vs 15–24 \times 3–4.5 µm, with rounded ends in the latter species; Fig. 8k–m). ITS sequences between these two species differed at 17–18 positions, with eight gaps.

Cryptoclypeus ryukyuensis A. Hashim. & Kaz. Tanaka, sp. nov. — MycoBank MB819247; Fig. 2g-h, 8

Etymology. Referring to the collection site.

Saprobic on dead twigs of Pleioblastus linearis.

Sexual morph: Ascomata ellipsoidal, solitary to 3–5-grouped, immersed, 220–240 μm high, 330–380 μm diam. Ostiolar neck

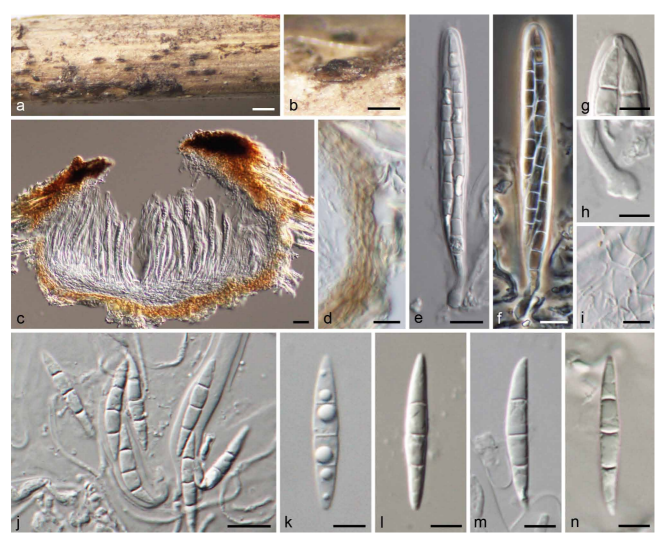


Fig. 7 Cryptoclypeus oxysporus. a-b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma; e-f. asci; g. ascus apex; h. ascus stipe; i. pseudoparaphyses; j-n. ascospores (all: HHUF 30507, holotype). — Scale bars: a = 1 mm; $b = 250 \text{ }\mu\text{m}$; $c = 20 \text{ }\mu\text{m}$; d-f, $i = 10 \text{ }\mu\text{m}$; q-h, $i-n=5 \text{ }\mu\text{m}$.

crest-like, elongated, laterally compressed, with less-developed black clypeus (380–450 µm wide). *Peridium* uniform, 22.5–25 µm thick at side, composed of rectangular, thin-walled, 10–22.5 \times 5–6.5 µm, pale brown cells. *Pseudoparaphyses* numerous, 1–1.5 µm wide, septate, branched and anastomosed. *Asci* bitunicate, fissitunicate, cylindrical, 72.5–100.5 \times 6–10 µm (av. = 87.0 \times 7.8 µm, n = 20), with a short stipe (4–9 µm long, av. = 6.5 µm, n = 20), apically rounded with an ocular chamber, 8-spored. *Ascospores* fusiform with rounded ends, straight, 15–24 \times 3–4.5 µm (av. = 19.7 \times 3.8 µm, n = 50), I/w (3.8–)4.3–6.6 (av. = 5.2, n = 50), 3-septate, slightly constricted at the primary septum nearly median (0.45–0.53, av. = 0.50, n = 50), hyaline, smooth, guttulate when young.

Asexual morph: Conidiomata pycnidial, globose to subglobose, up to 230 μm high, 230–340 μm diam, 3–5-grouped, immersed. Ostiolar neck up to 50 μm high, carbonaceous, papillate. Peridium 17.5–27.5 μm thick, composed of 7–10 layers of 4–8 × 2.5–6.5 μm, subglobose to rectangular, brown cells. Conidiophores reduced to conidiogenous cells. Conidiogenous cells holoblastic, 7–12 × 2–3 μm, ampliform to cylindrical, hyaline, smooth. Conidia cylindrical with slightly angular ends, 11.5–15 × 2–2.5 μm (av. = 12.9 × 2.1 μm, n = 50), l/w 5.1–7.0 (–8.3) (av. = 6.3, n = 50), hyaline, smooth, 1-septate, guttulate when young.

Culture characteristics — Colonies on PDA attaining 28–30 mm diam within 21 d at 20 °C in the dark, velvety, radiately, centrally raised, smoke grey (105) to grey olivaceous (107); reverse greenish black (124) (Fig. 2g–h); no sporulation observed.

Specimens examined. Japan, Okinawa, Kunigami-son, Okuma, Mt Yonaha, on dead culm of *Pleioblastus linearis*, 18 May 2015, *A. Hashimoto* et al., AH 342 (HHUF 30510 paratype, ex-paratype living culture MAFF 245616); Yona, Mt Fuenchiji, on dead culm of *Pleioblastus linearis*, 19 May 2015, *K. Tanaka* et al., KT 3534 (HHUF 30509 holotype designated here, exholotype living culture MAFF 245615 = NBRC 112318).

Notes — Ex-holotype and ex-paratype isolates of *C. ryukyuensis* are derived from sexual and asexual morphs, respectively. Unfortunately, a connection between sexual and asexual forms of this species could not be confirmed in culture. Nonetheless, *rpb2* sequences were completely identical between the two strains, while ITS sequences differed at only one position and *tef1* sequences differed at two, neither of which caused an amino acid substitution. Culture characteristics were also identical (Fig. 2g-h). We therefore regard these isolates as conspecific.

Galeaticarpa A. Hashim. & Kaz. Tanaka, gen. nov. — Myco-Bank MB819248

 $\label{eq:carpa} \textit{Etymology}. \ \text{From the Latin } \textit{galea}, \ \text{meaning helmet}, \ \text{and } \textit{carpa}, \ \text{meaning fruiting body}.$

Type species. Galeaticarpa aomoriensis A. Hashim. & Kaz. Tanaka.

Saprobic on woody plants.

Sexual morph: Ascomata solitary to grouped, immersed to erumpent, subglobose. Ostiolar neck elongated, laterally compressed, surrounded by well-developed clypeus. Peridium

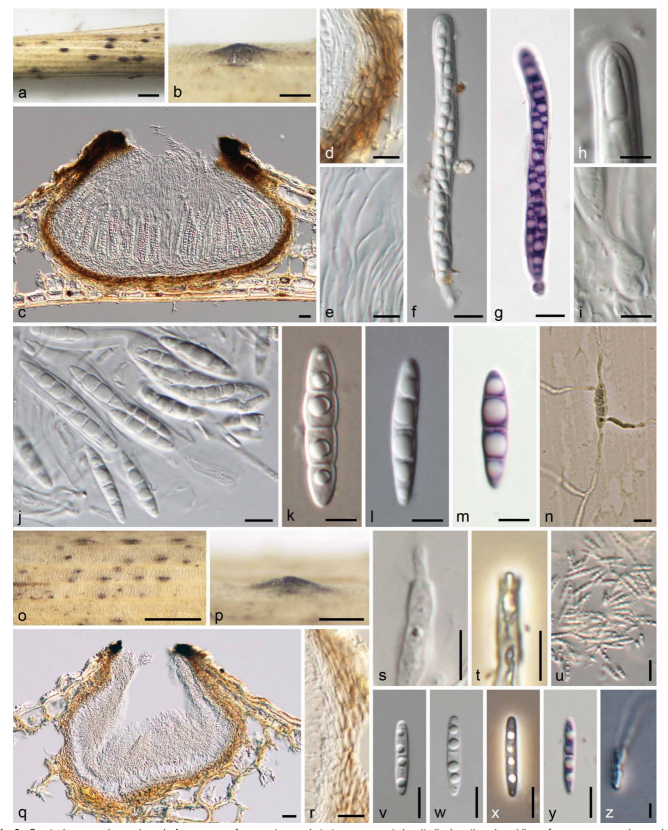


Fig. 8 Cryptoclypeus ryukyuensis. a—b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma; e. pseudoparaphyses; f—g. asci (g in Trypan Blue); h. ascus apex; i. ascus stipe; j—m. ascospores (m in Trypan Blue); n. germinating ascospore; o—p. conidiomata on substrate; q. conidioma in longitudinal section; r. peridium of conidioma; s—t. conidiogenous cells; u—y. conidia (y in Trypan Blue); z. germinating conidium (a—n: HHUF 30509, holotype; o—z: HHUF 30510, paratype). — Scale bars: a, o = 1 mm; b, p = 250 μ m; c, q = 20 μ m; d—g, j, n, r, u = 10 μ m; h—i, k—m, s—t, v—z = 5 μ m.

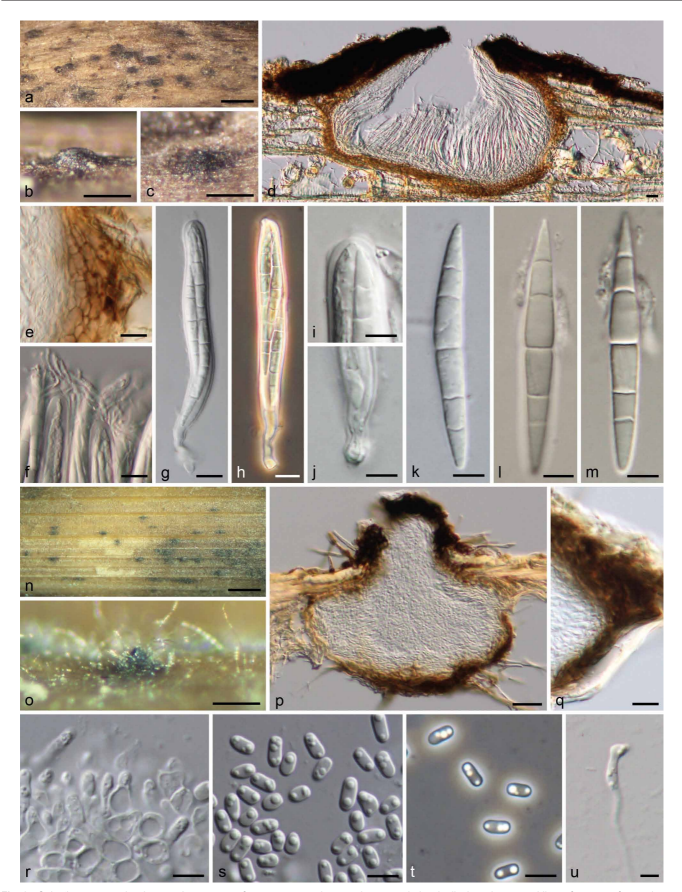


Fig. 9 *Galeaticarpa aomoriensis.* a–c. Appearance of ascomata on substrate; d. ascoma in longitudinal section; e. peridium of ascoma; f. pseudoparaphyses; g–h. asci; i. ascus apex; j. ascus stipe; k–m. ascospores; n–o. conidiomata in culture; p. conidioma in longitudinal section; q. peridium of conidioma; r. conidiogenous cells; s–t. conidia; u. germinating conidium (a–m: HHUF 30505, holotype; n–u: MAFF 245618 = NBRC 112319, ex-holotype culture). — Scale bars: a, n = 1 mm; b–c, o = 250 μ m; d, p = 20 μ m; e–h, q = 10 μ m; i–m, r–u = 5 μ m.

composed of rectangular, brown cells. *Pseudoparaphyses* septate, branched and anastomosed. *Asci* bitunicate, fissitunicate, cylindrical, 8-spored. *Ascospores* fusiform, multi-septate, hyaline, smooth.

Asexual morph: Conidiomata pycnidial, globose to subglobose, grouped, immersed. Ostiolar neck carbonaceous, papillate. Peridium composed of rectangular, pale brown to brown cells. Conidiophores absent. Conidiogenous cells holoblastic, cylindrical, hyaline, smooth. Conidia ellipsoidal with rounded ends, hyaline, smooth, aseptate.

Galeaticarpa aomoriensis A. Hashim. & Kaz. Tanaka, sp. nov. — MycoBank MB819249; Fig. 2i, 9

Etymology. Referring to the collection site.

Saprobic on dead twigs of woody plants.

Sexual morph: Ascomata ellipsoidal, solitary to 4-5-grouped, immersed to erumpent, subglobose, 265-285 µm high, 370–400 µm diam. Ostiolar neck crest-like, elongated, laterally compressed, surrounded by well-developed clypeus (620–750 μm wide). Peridium uniform, 15-26 μm thick, composed of 3–5 layers of rectangular, thin-walled, $7.5 \times 3-6.5 \mu m$, brown cells. Pseudoparaphyses numerous, 1–1.5 µm wide, septate, branched and anastomosed. Asci numerous, bitunicate, fissitunicate, cylindrical, $78-102 \times 8.5-11 \mu m$ (av. = 93.5×9.7 μ m, n = 11), with a short stipe (4–11 μ m long, av. = 7.2 μ m, n = 11), apically rounded with an ocular chamber, 8-spored. Ascospores fusiform with acute ends, straight, $31-47 \times 4-5.5$ μ m (av. = 40.6 × 4.7 μ m, n = 50), I/w (5.8–)7.2–10.5 (av. = 8.7, n = 50), 5-septate, with a primary septum nearly median (0.46-0.57, av. = 0.51, n = 50), slightly constricted at the primary septum, hyaline, smooth.

Asexual morph: Conidiomata pycnidial, flask-shaped, up to 135 μm high in section, 210-250(-420) μm diam, 3-5-grouped, semi-immersed. Ostiolar neck 32-50 μm high, carbonaceous, papillate. Peridium 10-12.5 μm wide, composed of 2-3 layers of $10-12\times3-4$ μm, rectangular, pale brown to brown cells. Conidiophores reduced to conidiogenous cells. Conidiogenous cells holoblastic, $5.5-11\times2.5-3$ μm, cylindrical, hyaline, smooth. Conidia ellipsoidal with rounded ends, $3.5-6\times1.8-2.2$ μm (av. = 4.5×2.0 μm, n = 50), l/w 1.8-3.0 (av. = 2.2, n = 50), hyaline, smooth, aseptate, guttulate when young.

Culture characteristics — Colonies on PDA attaining 17–19 mm diam within 21 d at 20 °C in the dark, velvety, plane, smoke grey (105); reverse chestnut (40), brown vinaceous (84) pigment produced (Fig. 2i); asexual morph formed.

Specimen examined. Japan, Aomori, Nishimeya, Shirakami, Ooshirosawa stream, on dead twigs of dead woody plant, 30 Aug. 2008, *K. Tanaka* et al., KT 2563 (HHUF 30505 holotype designated here, ex-holotype living culture MAFF 245618 = NBRC 112319).

Notes — A new monotypic genus, *Galeaticarpa*, is proposed here for species having ascomata with a clypeus, cylindrical asci with a short stipe, and fusiform, multi-septate, hyaline ascospores. These morphological characters are similar to those of *Astrosphaeriella*, but *Galeaticarpa* can be distinguished from *Astrosphaeriella* by its crest-like ostiolar neck and uniformly developed peridium (vs a poorly developed peridium at the base in *Astrosphaeriella*; Chen & Hsieh 2004, Phookamsak et al. 2015). These two genera are distantly related and belong to the families *Lophiotremataceae* and *Astrosphaeriellaceae*, respectively.

Cryptoclypeus, Galeaticarpa, and Pseudocryptoclypeus are morphologically similar to one another in having ascomata with a clypeus, but Galeaticarpa can be distinguished from these other genera by its most striking features – a well-developed clypeus (up to 750 µm wide) (Fig. 9a–d) and flask-shaped

conidiomata (Fig. 9p). In addition to phenotypic differences, *Cryptoclypeus* and *Pseudocryptoclypeus* tend to occur exclusively on bamboos, while *Galeaticarpa* occurs on woody plant hosts. Several bambusicolous fungi have been reported to be phylogenetically unrelated to genera and species on other host plants, even though they have morphological similarities with those groups (Tanaka et al. 2009, Hashimoto et al. 2015b). Further discovery of new lineages related to these genera is needed to clarify the relationship between their evolution and host preference.

Pseudocryptoclypeus A. Hashim. & Kaz. Tanaka, gen. nov. — MycoBank MB819250

Etymology. After its morphological similarity to Cryptoclypeus.

Type species. Pseudocryptoclypeus yakushimensis A. Hashim. & Kaz. Tanaka.

Saprobic on bamboo.

Sexual morph: Ascomata scattered to grouped, immersed. Ostiolar neck crest-like, elongated, laterally compressed, with less-developed clypeus. Peridium composed of compressed, rectangular, thin-walled, brown cells. Pseudoparaphyses septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 8-spored. Ascospores fusiform with acute ends, multi-septate, hyaline, smooth.

Asexual morph: Conidiomata pycnidial, globose to subglobose, grouped, immersed, ostiolate. Peridium composed of rectangular, pale brown cells. Conidiophores absent. Conidiogenous cells holoblastic, ampliform to cylindrical, hyaline. Conidia cylindrical with rounded ends, hyaline, smooth, 1-septate.

Pseudocryptoclypeus yakushimensis A. Hashim. & Kaz. Tanaka, *sp. nov.* — MycoBank MB819251; Fig. 2j, 10

Etymology. Referring to the collection site.

Saprobic on dead culms of bamboo.

Sexual morph: Ascomata ellipsoidal, 4–5-grouped, immersed, 325-380 µm high, 250-260 µm diam. Ostiolar neck crestlike, elongated, laterally compressed, with less-developed clypeus (200-315 µm wide). Peridium 17.5-25 µm thick of 2 zones at side; outer zone 10-17.5 µm thick of compressed, thin-walled, 12-17 × 1.5-4.2 µm, brown cells; inner zone of 7.5–10 μ m thick, rectangular, 5–9 × 4–6 μ m, pale brown cells. Pseudoparaphyses 0.5-1 µm wide, septate, branched, anastomosed. Asci bitunicate, fissitunicate, cylindrical, 92.5–127 x $7.5-10 \mu m$ (av. = $104.8 \times 8.2 \mu m$, n = 15), with a short stipe $(5-15 \mu m long, av. = 8.2 \mu m, n = 15)$, apically rounded with an ocular chamber, 8-spored. Ascospores fusiform with acute ends, straight, $20-32.5 \times 3-5 \mu m$ (av. = $25.4 \times 4.0 \mu m$, n = 50), I/w 4.7–8.1 (av. = 6.4, n = 50), 5-septate, with a primary septum nearly median (0.44-0.57, av. = 0.50, n = 50), slightly constricted at the primary septum, hyaline, smooth, with gelatinous pad at each end.

Asexual morph: Conidiomata pycnidial, globose to subglobose, up to 375 μm high, 400–480 μm diam, 5–6-grouped, immersed. Ostiolar neck carbonaceous, papillate, 47.5–77.5 μm high. Peridium 10–15 μm thick, composed of 3–4 layers of 7.5–10 \times 0.8–1.0 μm, rectangular, pale brown cells. Conidiophores absent. Conidiogenous cells holoblastic, 12–16 \times 2.5–3 μm, ampliform to cylindrical, hyaline, smooth. Conidia cylindrical with rounded ends, 9–14 \times 2.5–3.5 μm (av. = 12.0 \times 3.0 μm, n = 50), l/w 3.3–4.8 (av. = 4.0, n = 50), hyaline, smooth, 1-septate, guttulate when young.

Culture characteristics — Colonies on PDA attaining 32–52 mm diam within 21 d at 20 °C in the dark, floccose, plane, smoke grey (105); reverse grey olivaceous (107) to olivaceous black (108) (Fig. 2j); asexual morph formed.

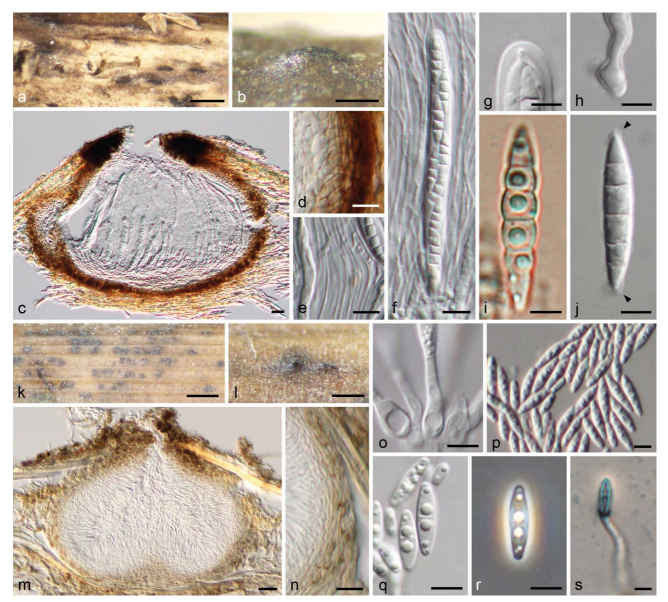


Fig. 10 Pseudocryptoclypeus yakushimensis. a – b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma; e. pseudoparaphyses; f. ascus; g. ascus apex; h. ascus stipe; i – j. ascospores (j arrowheads indicate gelatinous pad); k – l. conidiomata in culture; m. conidioma in longitudinal section; n. peridium of conidioma; o. conidiogenous cells; p – r. conidia; s. germinating conidium (a – j: HHUF 30503, holotype; k – s: MAFF 245622 = NBRC 112320, ex-holotype culture). — Scale bars: a, k = 1 mm; b = 200 μ m; c, m = 20 μ m; d – f, n = 10 μ m; g – j, o – s = 5 μ m; I = 250 μ m.

Specimen examined. Japan, Kagoshima, Isl. Yakushima, Nagata, on dead culms of bamboo, 16 Mar. 2007, *K. Tanaka & H. Yonezawa*, KT 2186 (HHUF 30503 holotype designated here, ex-holotype living culture MAFF 245622 = NBRC 112320).

Notes — *Pseudocryptoclypeus* is similar to *Cryptoclypeus* in having a less-developed clypeus in ascomata, multi-septate ascospores, pycnidial conidiomata, and 1-septate, hyaline conidia. *Pseudocryptoclypeus* has an ascomatal peridium composed of 2 zones and holoblastic conidiogenous cells (Fig. 10d, o) rather than the ascomatal wall with 1 zone and phialidic conidiogenous cells of *Cryptoclypeus* (Fig. 8d, s–t). Although these two genera constituted a highly supported clade (90 % ML BP/1.00 Bayesian PP) in our phylogenetic tree (Fig. 1), they had 43–45 base differences with 61–63 gaps in their ITS regions.

INCERTAE SEDIS

Antealophiotrema A. Hashim. & Kaz. Tanaka, gen. nov. — MycoBank MB819252

 ${\it Etymology}. \ {\it Formerly belonging to Lophiotrema}.$

Type species. Antealophiotrema brunneosporum (Ying Zhang, J. Fourn. & K.D. Hyde) A. Hashim. & Kaz. Tanaka.

Saprobic on woody plants.

Sexual morph: Ascomata subglobose to depressed ellipsoidal. Ostiolar neck crest-like, elongated, laterally compressed. Peridium composed of globose to rectangular, hyaline to brown cells, with brown hyphae at side. Pseudoparaphyses septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical to clavate, 4–8-spored. Ascospores narrowly fusiform, 1-septate, brown, smooth.

Asexual morph: Mycelium superficial, brown. Conidiophores absent. Conidiogenous cells holoblastic, integrated, terminal, brown, truncate, solitary, acrogenous. Conidia globose to obovoid, black, muriform.

Antealophiotrema brunneosporum (Ying Zhang, J. Fourn. & K.D. Hyde) A. Hashim. & Kaz. Tanaka, comb. nov. — Myco-Bank MB819253; Fig. 2k, 11

Basionym. Lophiotrema brunneosporum Ying Zhang, J. Fourn. & K.D. Hyde, Fung. Diversity 38: 240. 2009.

Saprobic on decorticated wood of Salix sp.

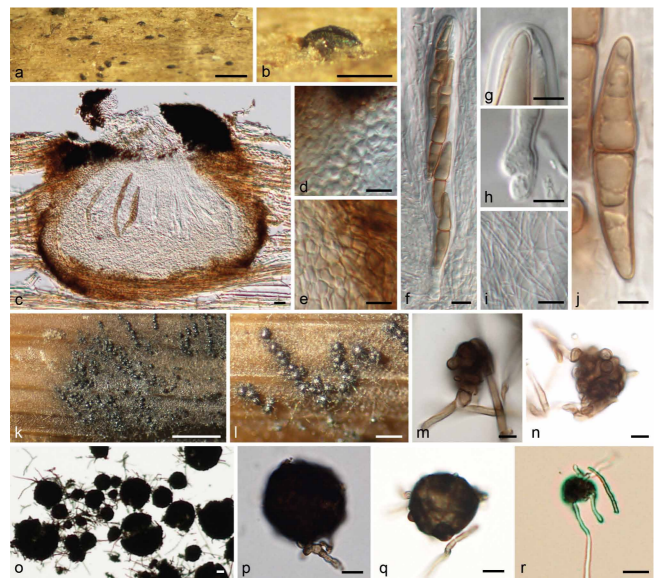


Fig. 11 Antealophiotrema brunneosporum. a–b. Appearance of ascomata on substrate; c. ascoma in longitudinal section; d. peridium of ascoma near ostiole; e. peridium of ascoma at side; f. ascus; g. ascus apex; h. ascus stipe; i. pseudoparaphyses; j. ascospore; k–l. conidia in culture; m–n. conidiogenous cells and immature conidia; o–q. conidia; r. germinating conidium (a–j: CBS H-20222, holotype; k–r: CBS 123095, ex-holotype culture). — Scale bars: a, k = 1 mm; b, l = 250 μ m; c, o, r = 20 μ m; d–f, i, p–q = 10 μ m; g–h, j, m–n = 5 μ m.

Sexual morph: Ascomata subglobose to depressed ellipsoidal, up to 380 µm high, 460-530 µm diam. Ostiolar neck crestlike, elongated, laterally compressed. Peridium ununiform, 42.5-62.5 µm thick, composed of 2 zones; outer zone 22.5-25 µm thick, composed of moderately thick-walled, rectangular, $(5-)7-9 \times 5-7.5 \mu m$, brown cells, with brown hyphae; inner zone 20–32.5 μ m thick of thin-walled, rectangular, 7–9 × 4–7.5 μm, hyaline to pale brown cells; at base 27.5-35 μm thick, of globose to rectangular, $3.5-7 \times 3.5-4 \mu m$, pale brown cells. Pseudoparaphyses numerous, 0.8-1.5 µm wide, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical to clavate, $119-148 \times 13-15 \mu m$ (av. = 138.2×14.3 μ m, n = 8), with a short stipe (8.5–16 μ m long, av. = 10.9 μ m, n = 8), apically rounded with an ocular chamber, with biseriate 4-8 ascospores. Ascospores narrowly fusiform with slightly rounded ends, $34.5-48 \times 6.5-10 \ \mu m$ (av. = $39.8 \times 8.0 \ \mu m$, n = 32), I/w 4.1–6.1 (av. = 5.0, n = 32), with a septum nearly median (0.44-0.54, av. = 0.49, n = 50), strongly constricted at the septum, brown, smooth.

Asexual morph: Mycelium superficial, brown. Conidiophores absent. Conidiogenous cells holoblastic, integrated, terminal,

brown, truncate, solitary, acrogenous. *Conidia* globose to obovoid, black, 27.5–85 µm diam (av. = 52.0 µm, n = 57), muriform.

Culture characteristics — Colonies on PDA attaining 12–17 mm diam within 21 d at 20 °C in the dark, floccose, centrally raised, grey olivaceous (107); reverse olivaceous black (108) (Fig. 2k); asexual morph formed.

Specimen examined. France, Ariège, Rimont, Las Muros, on decorticated wood of Salix sp., 24 Sept. 2006, J. Fournier & K.D. Hyde (CBS H-20222 holotype, ex-holotype culture CBS 123095).

Notes — The transfer of *L. brunneosporum* to *Antealophiotrema* is based on its morphological features, as it differs from *Lophiotrema* s.str. in having a well-developed peridium (up to 62.5 µm thick, Fig. 11c) as well as a monodictys-like asexual morph (Fig. 11k-r). In our phylogenetic tree, this species and *'Lophiotrema' boreale* (CBS 114422) formed a fully supported clade (100 % ML BP/1.00 Bayesian PP) outside of *Lophiotremataceae* and are recognised as a lineage distinct from *Lophiotrema* s.str. (Fig. 1). We treat *Antealophiotrema* as *'incertae sedis'* in *Pleosporales* at present; additional taxa related to this monotypic genus will be required to resolve its familial placement.

Pseudolophiotrema A. Hashim. & Kaz. Tanaka, gen. nov. — MycoBank MB819254

Etymology. After its morphological similarity to Lophiotrema.

Type species. Pseudolophiotrema elymicola A. Hashim. & Kaz. Tanaka.

Saprobic on herbaceous plants.

Sexual morph: Ascomata grouped, immersed, globose. Ostiolar neck crest-like, elongated, laterally compressed. Peridium composed of compressed, thin-walled, pale brown cells. Pseudoparaphyses numerous, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 8-spored. Ascospores fusiform, 1-septate, hyaline, smooth.

Asexual morph: Undetermined.

Pseudolophiotrema elymicola A. Hashim. & Kaz. Tanaka, sp. nov. — MycoBank MB819255; Fig. 2I, 12

Etymology. Referring to the generic name of the host plant.

Saprobic on dead leaves of Leymus mollis.

Sexual morph: Ascomata grouped, immersed, globose, 200–300 μm high, 190–340 μm diam. Ostiolar neck crest-like, elongated, laterally compressed. Peridium uniform, 10–12.5 μm thick composed of compressed, thin-walled, 5–12 × 2–3.5 μm, pale brown cells. Pseudoparaphyses numerous, 1.5–2 μm wide, septate, branched and anastomosed. Asci bitunicate, fissitunicate, cylindrical, 82–108 × 10–17 μm (av. = 92.4 × 13.9 μm, n = 11), with a short stipe (5–8 μm long, av. = 6.3 μm, n = 10), apically rounded with an ocular chamber, 8-spored. Ascospores fusiform with acute ends, straight, 20–28 × 4.5–7 μm (av. = 22.7 × 5.5 μm, n = 50), l/w 3.2–5.1 (av. = 4.2, n = 50), with a septum nearly median (0.47–0.56, av. = 0.51, n = 50), strongly constricted at the septum and midpoint of each cell, hyaline, smooth, with a gelatinous sheath.

Asexual morph: Undetermined.

Culture characteristics — Colonies on PDA attaining 18–19 mm diam within 21 d at 20 °C in the dark, velvety, plane, radiate-

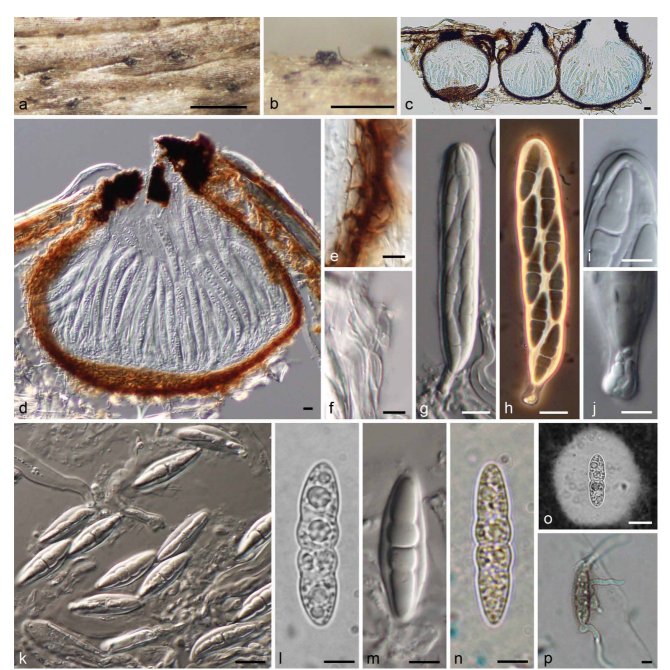


Fig. 12 Pseudolophiotrema elymicola. a-b. Appearance of ascomata on substrate; c-d. ascomata in longitudinal section; e. peridium of ascoma; f. pseudoparaphyses; g-h. asci; i. ascus apex; j. ascus stipe; k-n. ascospores; o. ascospore with a gelatinous sheath (in India ink); p. germinating ascospore (a-m, o-p: HHUF 28984, holotype; n: JCM 13090 = MAFF 239600, ex-holotype culture). — Scale bars: $a = 500 \mu m$; $b = 250 \mu m$; $c-d = 20 \mu m$; e-h, e-h,

ly, smoke grey (105); reverse grey olivaceous (107) (Fig. 2l); sexual morph formed.

Specimen examined. Japan, Hokkaido, Yufutsu, on dead leaves of Leymus mollis, 1 Sept. 2003, Y. Harada, KT 1450 (HHUF 28984 holotype designated here, ex-holotype culture JCM 13090 = MAFF 239600).

Notes — Pseudolophiotrema is quite similar to Lophiotrema in having a crest-like ostiolar neck, a peridium of uniform thickness, cylindrical asci with a short stipe, and 1-septate hyaline ascospores with a gelatinous sheath. However, it can be separated from the latter genus by its thinner ascomatal wall (10–12.5 µm thick; Fig. 12d–e) composed of compressed cells vs the 20–30 µm thick wall made up of rectangular cells in Lophiotrema (Holm & Holm 1988).

In our phylogenetic tree, the genus *Pseudolophiotrema* was placed completely outside of *Lophiotremataceae*, and was also separate from the families *Cryptocoryneaceae* and *Aquasubmersaceae* (Fig. 1). The familial placement of *Pseudolophiotrema* remains unclear. Additional taxa belonging to this genus are needed to fully understand its taxonomic affiliations within *Pleosporales*.

DISCUSSION

Lophiotremataceae was established by Hirayama & Tanaka (2011) to accommodate Lophiotrema. Subsequently, Aquasubmersa and Hermatomyces were recognised as additional members of this family on the basis of molecular phylogenetic analyses (Ariyawansa et al. 2015, Doilom et al. 2016, Hyde et al. 2016). A phylogenetic relationship between Cryptocoryneum and Lophiotremataceae was also suggested by a BLAST search of ITS sequences of species in that genus (Hashimoto et al. 2016). Although tree topologies generated in previous molecular studies have suggested that the above-mentioned genera might be members of Lophiotremataceae, statistical support for a Lophiotremataceae s.lat. clade has been relatively weak (Ariyawansa et al. 2015, Doilom et al. 2016, Hashimoto et al. 2016, Hyde et al. 2016). Our results do not support the monophyly of Lophiotremataceae s.lat. as recognised in several previous studies (Fig. 1). Instead, the various asexual morphs found in Aquasubmersa, Cryptocoryneum, and Hermatomyces belong to separate families whose monophyly is strongly supported (Fig. 1). We thus consider *Lophiotremataceae* to be restricted to Lophiotrema and five new genera (Atrocalyx, Crassimassarina, Cryptoclypeus, Galeaticarpa, and Pseudocryptoclypeus), all of which have ascomata with or without slit-like ostioles and pycnidial conidiomata. We also conclude that Aquasubmersa (a freshwater lineage having papillate ascomatal ostioles and pycnidial conidiomata with 1-celled conidia), Cryptocoryneum (having sporodochial conidiomata with cheiroid conidia), and Hermatomyces (having sporodochial conidiomata with lenticular and/or cylindrical conidia), which have provisionally been regarded as genera of Lophiotremataceae, should be placed in their own families given their different asexual morphs.

Except for *Crassimassarina*, genera accepted in the present study as members of *Lophiotremataceae* (i.e., *Lophiotrema* s.str. and the newly introduced genera *Atrocalyx*, *Cryptoclypeus*, *Galeaticarpa*, and *Pseudocryptoclypeus*) have features that are generally consistent with traditional generic concepts of *Lophiotrema* s.lat. (Holm & Holm 1988, Tanaka & Harada 2003, Zhang et al. 2009, Hirayama & Tanaka 2011). In contrast, *Crassimassarina* has some characteristics atypical of *Lophiotremataceae*: large, immersed ascomata with a papillate ostiolar neck, resembling those of genera in *Pleomassariaceae* s.lat. (Barr 1982, Tanaka et al. 2005), as well as pycnidial conidiomata with multi-septate cylindrical conidia resembling

those of Stagonospora (Quaedvlieg et al. 2013, Tanaka et al. 2015). The phylogenetic placement of this morphologically distinct lineage in Lophiotremataceae led us to conduct detailed observations of the above lophiotremataceous genera. These observations revealed morphological variations among ascomata with slit-like ostioles in Lophiotremataceae, such as ascomata with a distinct clypeus around the ostiolar neck (in Cryptoclypeus, Galeaticarpa and Pseudocryptoclypeus, Fig. 7c, 8c, 9d, 10c), ascomata with a well-developed peridium (Atrocalyx, Fig. 4c, 5c), and ascomata with a less-developed peridium of a uniform thickness up to 25 µm (Lophiotrema; Holm & Holm 1988). These morphologically distinguishable groups constitute phylogenetically distinct lineages that also correspond to their differing asexual morphs (Fig. 4o, 6q, 8q, 9p, 10m). For example, the sexual morphs of *Cryptoclypeus*, Galeaticarpa, and Pseudocryptoclypeus have similar morphological features, but these genera can be separated from one another according to the morphology of their conidiomata or conidiogenous cells (see Notes on each genus). Consequently, we treat these five genera as separate from Lophiotrema s.str. on the basis of holomorph morphology and phylogenetic relationships. Antealophiotrema brunneosporum, originally misidentified as a species in *Lophiotrema* because of its morphological resemblance to that genus (Zhang et al. 2009), is only distantly related to Lophiotremataceae. This species is clearly different from Lophiotrema in regard to the anatomical characteristics of its peridium and its monodictys-like asexual morph. The result of our present study strongly confirms that the presence of a slit-like ostiole is an unreliable character for delimitation of generic relationships.

Traditionally, the slit-like ostiole of the ascomata has been especially emphasised as a useful character for familial circumscription in ascomycetes (Chesters & Bell 1970). In earlier studies, Lophiotrema was placed in Lophiostomataceae according to this criterion (Chesters & Bell 1970, Leuchtmann 1985, Holm & Holm 1988, Barr 1992). Zhang et al. (2009) suggested that Lophiotrema is phylogenetically distinct from Lophiostomataceae but could not find any morphological differences between their sexual morphs. Hirayama & Tanaka (2011) re-evaluated the phylogenetic significance of several morphological features used for characterisation of each genus in previous studies. After determining that ascus shape and length of ascus stipe are reliable taxonomic indicators to delineate these two genera, they established Lophiotremataceae to accommodate Lophiotrema (Hirayama & Tanaka 2011). Our results, however, strongly confirm that several lophiotrema-like species, i.e., Antealophiotrema brunneosporum, 'Lophiotrema' boreale, and Pseudolophiotrema elymicola, deviate from Lophiotremataceae. These species were originally misidentified or provisionally identified as species in Lophiotrema on the basis of their morphological resemblance to the genus (Mathiassen 1989, 1993, Zhang et al. 2009), but were found to be phylogenetically separate from Lophiotrema s.str. in this study. Previous morphological circumscriptions of Lophiotremataceae and Lophiotrema (Holm & Holm 1988, Mathiassen 1989, 1993, Tanaka & Harada 2003, Zhang et al. 2009, Hirayama & Tanaka 2011, Ariyawansa et al. 2015, Doilom et al. 2016, Hyde et al. 2016) were obviously fairly broad concepts that did not reflect their phylogenetic relationships. We additionally discovered several morphological variants of ascomata having slit-like ostioles (e.g. Atrocalyx, Cryptoclypeus, Galeaticarpa, and Pseudocryptoclypeus), revealed the phylogenetic position of a genus without slit-like ostioles in Lophiotremataceae (i.e., Crassimassarina), and observed asexual morphs of genera belonging to Lophiotremataceae. More precise morphological examination of both sexual and asexual morphs in this family will be needed to define familial concepts of Lophiotremataceae and its relatives.

The phylogenetic significance of slit-like ostioles of ascomata should also be re-evaluated because this phenotypic character is now known in several families, such as *Aigialaceae* (Suetrong et al. 2009), *Ligninsphaeriaceae* (Zhang et al. 2016), and *Lophiostomataceae* (Thambugala et al. 2015). The slit-like ostiole should be regarded as a character that has evolved multiple times independently within *Dothideomycetes*, similar to the parallel evolution of hysterothecial ascomata in *Anteagloniaceae* (Mugambi & Huhndorf 2009), *Gloniaceae* (Boehm et al. 2009), *Hysteriaceae* (Boehm et al. 2009), *Lophiostomataceae* (Thambugala et al. 2015), and *Mytilinidiaceae* (Boehm et al. 2009).

Recent molecular studies on *Dothideomycetes* have revealed hidden lineages and prompted revision of several families in this class (Crous et al. 2015, Guatimosim et al. 2015, Knapp et al. 2015, Tanaka et al. 2015, Jaklitsch & Voglmayr 2016, Van Nieuwenhuijzen et al. 2016). Although multiple molecular systematic studies, mainly of pathogenic fungi of woody plants (Phillips et al. 2013, Slippers et al. 2013, Alves et al. 2014, Fan et al. 2015, Trakunyingcharoen et al. 2015), have generated a robust phylogeny for Botryosphaeriales, many additional new lineages in this group, including endophytes and saprophytes, have been discovered from various niches (Thambugala et al. 2014, Crous et al. 2015, 2016, Wyka & Broders 2016, Osorio et al. 2017, Yang et al. 2017). Drawing on many recent collections, our study has revealed previously unrecognised diversity within Lophiotremataceae and its relatives and the potential taxonomic importance of the asexual morphs in defining families. To build a comprehensive taxonomic framework, further morphological surveys based on additional collections together with more molecular data are needed.

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