**Taxonomic Paper**

*Paraeutypella guizhouensis* gen. et sp. nov. and *Diatrypella longiasca* sp. nov. (*Diatrypaceae*) from China

Lakmali S. Dissanayake‡, Nalin N. Wijayawardene§∥, Monika C. Dayarathne¶, Milan C. Samarakoon#, Dong-Qin Dai¶, Kevin D. Hyde#, Ji-Chuan Kang‡

‡ Engineering Research Centre of the Utilization for Characteristic Bio-Pharmaceutical Resources in Southwest, Ministry of Education, Guizhou University, Guiyang, Guizhou Province 550025, China
§ Center for Yunnan Plateau Biological Resources Protection and Utilization, College of Biological Resource and Food Engineering, Qujing Normal University, Qujing, Yunnan 655011, China
∥ State Key Laboratory of Functions and Applications of Medicinal Plants, Guizhou Medical University, Guiyang 550014, China
¶ Department of Plant Pathology, Agriculture College, Guizhou University, Guiyang, Guizhou Province, 550025, China
# Center of Excellence in Fungal Research, Mae Fah Luang University, Chiang Rai, 57100, Thailand

Corresponding author: Ji-Chuan Kang (jckang@gzu.edu.cn)

Academic editor: Renan Barbosa

Received: 02 Feb 2021 | Accepted: 07 Mar 2021 | Published: 26 Mar 2021

Citation: Dissanayake LS, Wijayawardene NN, Dayarathne MC, Samarakoon MC, Dai D-Q, Hyde KD, Kang J-C (2021) *Paraeutypella guizhouensis* gen. et sp. nov. and *Diatrypella longiasca* sp. nov. (*Diatrypaceae*) from China. Biodiversity Data Journal 9: e63864. [https://doi.org/10.3897/BDJ.9.e63864](https://doi.org/10.3897/BDJ.9.e63864)

**Abstract**

**Background**

In this study, we introduce a novel genus, *Paraeutypella*, of the family *Diatrypaceae* comprising three species viz. *Paraeutypella guizhouensis* sp. nov. and *P. citricola* (basionym: *Eutypella citricola*) and *P. vitis* (basionym: *Sphaeria vitis*). *Diatrypella longiasca* sp. nov. is also introduced, which forms a distinct clade in *Diatrypella* sensu stricto. The discovery of this new genus will contribute to expanding the knowledge and taxonomic framework of *Diatrypaceae* (Xylariales).
New information

Generic delimitations in *Diatrypaceae* are unsettled because the phylogeny has yet to be resolved using extensive taxon sampling and sequencing of ex-type cultures. During an investigation of xylarialean fungi, we collected eutypella-like fungi which is distinct from *Eutypella* sensu stricto in our phylogenetic analyses (ITS and β-tubulin), thus, introduced as *Paraeutypella guizhouensis* gen. et sp. nov. *Paraeutypella* is characterised by having 4–25 perithecia in a stroma each with 3–6 sulcate, long ostiolar necks. *Paraeutypella citricola* comb. nov. (basionym: *Eutypella citricola*) is introduced on *Acer* sp. from China. *Diatrypella longiasca* sp. nov. is introduced as a new species in *Diatrypella* sensu stricto. which has 2–5 ascomata per stroma and long ascospores, unusual when compared to other *Diatrypella* species and distinct phylogenetically.

Keywords

*Acer*, morphology, novel taxa, phylogeny, Xylariales

Introduction

*Diatrypaceae* Nitschke (Ascomycota, Xylariales) comprises 21 genera and more than 1,500 species (Senwanna 2017, Mehrabi et al. 2019, Dayarathne et al. 2020b, Wijayawardene et al. 2020). Species of this family are characterised by erumpent to immersed, rarely superficial, black or dark brown, eustromatic or pseudostromatic stromata and 8-spored or polysporous asci with hyaline to light brown, allantoid ascospores (Konta et al. 2020) in their sexual morph. Several asexual morph genera have been linked to the family *Diatrypaceae*, including *Cytosporina* Sacc. and *Libertella* Desm. (Glawe and Rogers 1984). *Cytosporina* Sacc. includes species with pycnidial and filiform conidia; *Libertella* Desm. includes species with acervula and filiform conidia (Glawe and Rogers 1984).

Members of *Diatrypaceae* are saprobes, pathogens or endophytes, associated with a wide range of hosts in terrestrial and aquatic environments (Mehrabi et al. 2019, Dayarathne et al. 2020a, Dayarathne et al. 2020b Konta et al. 2020). Dayarathne et al. (2020a), Dayarathne et al. (2020b) introduced two novel genera, *Halocryptosphaeria* Dayar. et al. and *Halocryptovalsa* Dayar. & K.D. Hyde from marine environments. Species of *Anthostoma* Nitschke, *Cryptosphaeria* Ces. & De Not., *Crytovalsa* Ces. & De Not. ex Fuckel, *Diatrype* Fr., *Diatrypella* (Ces. & De Not.) De Not. and *Eutypella* (Nitschke) Sacc. have been reported as causal agents of canker diseases on a wide range of host plants worldwide (Hyde et al. 2020). The taxonomy and phylogeny of *Diatrypaceae* need to be resolved, as many genera are polyphyletic. Hence, fresh collections and sequences are required to define genera and establish their phylogenetic placement within the family.

*Diatrypella* was introduced by Cesati and De Notaris (1863) with *D. verruciformis* (Ehrh.) Nitschke as the type. The genus is characterised by conical to truncate, cushion-like or discoid stromata usually delimited by a black zone in host tissues, umbilicate or sulcate
Ostiolar necks, cylindrical, polysporous, long-stalked asci and allantoid, hyaline or yellowish ascospores in their sexual morph and a libertella-like coelomycetes asexual morph (Kirk et al. 2008, Hyde et al. 2020). Both Cryptovalsa and Diatrypella have polysporous asci and cannot easily be distinguished, based only on morphological comparisons (Acero et al. 2004, Vasilyeva and Stephenson 2005). Therefore, molecular data are essential for defining genera in Diatrypaceae (Mehrabi et al. 2015). There are 65 names of Diatrypella in Species Fungorum (2020) (http://www.indexfungorum.org/names/names.asp), but only 15 have molecular data in GenBank (Hyde et al. 2020).

In this study, we introduce a new genus, Paraeutypella, which shows eutypella-like morphology, but is distinct phylogenetically. Paraeutypella comprises three species viz. Paraeutypella guizhouensis sp. nov. and P. citricola (basionym: Eutypella citricola) and P. vitis (basionym: Sphaeria vitis). Diatrypella longiasca sp. nov. is also introduced, which forms a distinct clade in Diatrypella sensu stricto. Species novelties are confirmed by morphological comparisons along with micro-photographs and the phylogeny of combined ITS and β-tubulin sequence data.

Materials and methods

Sample collection and morphological observations

Dead twigs of Acer palmatum and undetermined plants were collected from China (Guiyang, Guizhou Province) during September to October 2019. Samples were observed with a stereomicroscope (SZX16, Olympus). Hand sections of the ascomata were mounted in distilled water and the following characters were measured: diameter and height of ascomata, width of the peridium, diameter and height of ostiolar necks, length and width of asci and ascospores. Melzer’s Reagent was used for testing the ascal apical ring reaction. Images were captured with a Canon EOS70D digital camera fitted to a compound microscope. Measurements were made with the Tarosoft (R) Image Frame Work programme and images used for figures processed with Adobe Photoshop CS6 software (Adobe Systems, USA). Single spore isolation was performed according to Chomnunti et al. (2014) and germinating spores were transferred to potato dextrose agar (PDA-Shanghai Bio-way Technology Co. Ltd.). The pure cultures were incubated at 18–20°C for four weeks. The type specimens were deposited in the Cryptogamic Herbarium, Kunming Institute of Botany, Academia Sinica (HKAS), Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium (HMAS), Beijing, China. Ex-type cultures were deposited in the Kunming Institute of Botany Culture Collection (KUMCC). Facesoffungi and Index Fungorum numbers are provided as mentioned in Jayasiri et al. (2015) and Index Fungorum (http://www.indexfungorum.org) respectively.

DNA extraction, PCR amplifications and sequencing

Fungal isolates were grown on PDA for 3–4 weeks at 25°C and total genomic DNA was extracted from 50 to 100 mg of axenic mycelium scraped from the edges of the growing cultures (Wu et al. 2001). EZgene™ fungal gDNA extraction kit (BIOMIGA, Hangzhou City,
Zhejiang Province, China) was used to extract DNA by following the manufacturer’s protocol. DNA extracts were stored at – 4°C for use in regular work and duplicated at – 20°C for long term storage.

DNA sequence data were obtained from the internal transcribed spacer (ITS) and partial β-tubulin gene. ITS and β-tubulin were amplified by using the primers ITS5/ITS4 (White et al. 1990) and T1/T22 (O'Donnell and Cigelnik 1997), respectively. Polymerase chain reaction (PCR) was carried out in a volume of 25 μl, which contained 9.5 μl of ddH₂O, 12.5 μl of 2× PCR Master Mix (2× Bench Top™ Taq Master Mix, BIOMIGA, China), 1 μl of DNA template and 1 μl of forward and reverse primers (10 μM each) in each reaction. The PCR thermal cycle programme for all gene amplifications was as follows: initialisation at 95°C for 5 min, followed by 35 cycles of denaturation at 95°C for 30 s, annealing at 55°C for 50 s, elongation at 72°C for 90 s and final extension at 72°C for 10 min. Purification and sequencing of PCR products were done by Sangon Biotech, Shanghai, China.

**Molecular phylogenetic analyses**

**Sequence alignment**

The sequence data generated in this study were analysed with closely-related taxa retrieved from GenBank (Table 1), based on BLASTn searches (https://www.ncbi.nlm.nih.gov) and recently published data (Mehrabi et al. 2019, Dayaratne et al. 2020b, Konta et al. 2020). ITS and β-tubulin were used for the analyses according to the previous studies listed above. Sequences (ITS and β-tubulin) were aligned using MAFFT v. 6.864b (Katoh et al. 2019) and manually improved when necessary in BioEdit v. 7.0 (Hall 1999). The single gene alignments were used to perform model test in MrModeltest 2.3 to estimate the best-fit evolutionary model under the Akaike Information Criterion (AIC) (Nylander 2004) and resulted in a GTR+I+G substitution model for each. Ambiguously aligned areas of each gene region were excluded and gaps were treated as missing data. Missing characters were assessed to be unordered and equally weighted.

| Species                      | Strain no.        | GenBank Accession no. | Reference             |
|------------------------------|-------------------|-----------------------|-----------------------|
| Allocryptovalsa cryptovalsoidea T | HVFIG02          | HQ692573              | Trouillas et al. (2011) |
| A. elaeidis                  | MFLUCC 15-0707   | MN308410 MN308415     | Konta et al. (2020)   |
| Allocotrypta arengae T       | MFLUCC 15-0713   | MN308411 MN308415     | Konta et al. (2020)   |
| A. elaeidicola T             | MFLUCC 15-0737   | MN308415 MN308419     | Konta et al. (2020)   |
| A. elaeidis                  | MFLUCC 15-0708b  | MN308413 NA           | Konta et al. (2020)   |
| Species                  | Strain no. | GenBank Accession no. | Reference                  |
|--------------------------|------------|-----------------------|----------------------------|
|                          |            | ITS                   | β-tubulin                  |
| **Anthostoma decipiens** | IPV-FW349  | AM399021              | AM920693                   | Nitschke (1867) |
| A. decipiens             | JL567      | JN975370              | JN975407                   | Luque et al. (2012) |
| **Cryptosphaeria eunomia** | CBS 216.87 | AJ302417              | NA                         | Acero et al. (2004) |
| C. var. eunomia          | CBS 223.87 | AJ302421              | NA                         | Acero et al. (2004) |
| **Cryptovals ampelina**  | A001       | GG293901              | GG293972                   | Trouillas et al. (2015) |
| C. ampelina              | KHJ20      | KJ767718              | KY352426                   | Mehrabi et al. (2015) |
| **Diatrypasimilis australiensis** | ATCC MYA 3540 | FJ430590              | NA                         | Chalkley et al. (2010) |
| D. bullata               | UCDDCh400  | DQ006946              | DQ007002                   | Rolshausen et al. (2006) |
| **D. disciformis**       | MFLUCC 15-0538 | KR092795              | NA                         | Senanayake et al. (2015) |
| D. lijiangensis          | MFLU 19-0717 | MK852582              | MK852583                   | Thyagaraja et al. (2019) |
| D. stigma                | DCASH200   | GQ293947              | GQ294003                   | Trouillas et al. (2015) |
| **Diatrypella atlantica** | HUEFS 194228 | KM396615              | KR363998                   | de Almeida et al. (2016) |
| D. atlantica             | HUEFS 192148 | KM396633              | KT175563                   | de Almeida et al. (2016) |
| D. delonicis             | MFLUCC 15-1014 | MH812994              | MH847790                   | Hyde et al. (2019) |
| D. delonicis             | MFLU 16-1032 | MH812995              | MH847791                   | Hyde et al. (2019) |
| D. elaeidis              | MFLUCC 15-0279 | MN308417              | MN340300                   | Konta et al. (2020) |
| D. favacea               | ANM 96     | KJ320616              | NA                         | de Almeida et al. (2016) |
| D. frostii               | UFMGCB 1917 | HQ377280              | NA                         | Vieira et al. (2012) |
| D. heveae                | MFLUCC 17-0368 | MF959501              | MG334557                   | Senwanna (2017) |
| D. heveae                | MFLUCC 15-0274 | MN308418              | MN340301                   | Konta et al. (2020) |
| D. iranensis             | IRAN 2280C | KM245033              | KY352429                   | Mehrabi et al. (2015) |
| **D. longiasca**         | KUMCC 20-0021 | MW039349              | MW239658                   | This study |
| **D. longiasca**         | KUMCC 20-0022 | MW036141              | MW239659                   | This study |
| D. macrospora            | IRAN 2344C | KR605648              | KY352430                   | Mehrabi et al. (2016) |
| D. major                 | ANM 1947   | KJ320613              | NA                         | de Almeida et al. (2016) |
| D. pulvinata             | H048       | FR715523              | FR715495                   | Pazoutova et al. (2012) |
| D. verruciformis         | UCROK1467  | JX144793              | JX174093                   | Lynch et al. (2013), Luque et al. (2012) |
| D. vulgaris              | HVFRA02    | HQ692591              | HQ692503                   | Trouillas et al. (2015) |
| D. vulgaris              | HVGRF03    | HQ692590              | HQ692502                   | Trouillas et al. (2015) |
| Eutypa laevesa           | CBS 291.87 | AJ302449              | NA                         | Acero et al. (2004) |
| E. lata                  | ATCC 28120 | DQ006948              | DQ006975                   | Rolshausen et al. (2006) |
| E. lata                  | EP18       | HQ692611              | HQ692501                   | Trouillas et al. (2011) |
| E. lata                  | RGA01      | HQ692614              | HQ692497                   | Trouillas et al. (2011) |
| Species                  | Strain no. | GenBank Accession no. | Reference                        |
|--------------------------|------------|-----------------------|----------------------------------|
| **ITS**                  |            |                       |                                  |
| E. lata var. aceri       | CBS 290.87 | HM164736              | HM164770                         | Trouillas et al. (2010)       |
| Eutypella caricae        | EL51C      | AJ302460              | NA                               | Acero et al. (2004)           |
| E. cerviculata           | EL59C      | AJ302468              | NA                               | Acero et al. (2004)           |
| E. cerviculata           | M68        | JF340269              | NA                               | Arhipova et al. (2012)        |
| E. leprosa               | EL54C      | AJ302463              | NA                               | Acero et al. (2004)           |
| E. leprosa               | ANM 85     | KU320622              | NA                               | de Almeida et al. (2016)      |
| E. microtheca            | ADEL200    | HQ692559  HQ692527   | Trouillas et al. (2011)          |
| E. microtheca            | BCMX01     | KC405566  KC405560   | Paolinelli-Alfonso et al. (2015) |
| E. parasitica            | CBS 210.39 | MH855984              | NA                               | Vu et al. (2019)              |
| E. persica T             | IRAN 2540C | KX828144  KY352451   | Mehrabi et al. (2019)            |
| E. quercina T            | IRAN 2543C | KX828139  KY352449   | Mehrabi et al. (2019)            |
| E. semicircularis T      | MP4669     | JQ517314              | NA                               | Chacón et al. (2013)          |
| E. tamaricis             | MFLUCC 14-0445 | NA | KX453302 | Thambugala et al. (2016) |
| E. virescens             | CBS 205.36 | MH855778  MH867286   | Vu et al. (2019)                 |
| Halocryptovalsa salicorniae | MFLUCC 15-0185 | MH304410  MH370274 | Dayarathne et al. (2020b)        |
| Halodiatrype avicenniae  | MFLUCC 15-0948 | MH304414  MH370278 | Dayarathne et al. (2020b)        |
| H. salinicola T          | MFLUCC 15-1277 | KX573915  KX573932 | Dayarathne et al. (2016)         |
| H. salinicola            | MFLUCC17-2468 | MN047113 | NA | Dayarathne et al. (2016) |
| Kretzschmaria deusta T   | CBS 826.72 | KU683767  KU684190   | U’Ren et al. 2016                |
| Monosporascus cannonballus T | ATCC 26931 | FJ430598 | NA | Unpublished                  |
| M. cannonballus          | CMM3646    | JX971617              | NA                               | Sales et al. (2010)           |
| Neoeutypella baoshanensis | MFLUCC 16-1002 | MT310662 | NA | Phukhamsakda et al. (2020)   |
| N. baoshanensis T        | LC 12111   | MH822887  MH822888   | Hyde et al. (2019)               |
| Paraeutypella citricolca | HVGRF01    | HQ692579  HQ692512   | Trouillas et al. (2011)          |
| P. citricola             | HVVIT07    | HQ692589  HQ692521   | Trouillas et al. (2011)          |
| P. citricola             | IRAN 2340C | KR605647  KY352439   | Mehrabi et al. (2016)            |
| P. citricola             | KUMCC 20-0023 | MW040050  MW239663  | This study                       |
| P. citricola             | KUMCC 20-0024 | MW040049  MW239662  | This study                       |
| P. guizhouensis T        | KUMCC 20-0016 | MW036142  MW239660  | This study                       |
| P. guizhouensis          | KUMCC 20-0017 | MW039348  MW239661  | This study                       |
| P. vitis                 | UCD2291AR  | HQ288224  HQ288303   | Úrbez-torres et al. (2012)       |
| P. vitis                 | UCD2428TX  | FJ790851  GU294726   | Úrbez-Torres and Gubler (2009)   |
| Pedumispora rhizophorae  | BCC44877   | KJ888853 | NA | Klaysuban et al. (2014)      |
| P. rhizophorae           | BCC44878   | KJ888854 | NA | Klaysuban et al. (2014)      |
| Species               | Strain no. | GenBank Accession no. | Reference                      |
|----------------------|------------|-----------------------|--------------------------------|
| *Peroneutypa curvispora* | HUEFS 136877 | KM396641               | de Almeida et al. (2016)       |
| *P. rubiformis* T    | MFLUCC 17-2142  | MG873477               | Shang et al. (2018)            |
| *P. scoparia*        | MFLUCC 11-0478  | KU940151               | Dai et al. (2016)              |
| *Quaternaria quaternata* | CBS 278.87       | AJ302469               | Acer et al. (2004)             |
| *Q. quaternata*      | GNF13        | KR605645 KX352464      | Mehrabi et al. (2016)          |
| *Xylaria hypoxylon*  | CBS-122620  | KY610407 KY271279      | Peršoh et al. (2009)           |

T: Types strains, newly-generated sequences are indicated in bold, NA: No sequence available in GenBank, ATCC: American Type Culture Collection, Manassas, USA, BCC: BIOTEC Culture Collection, Bangkok, Thailand, CBS: Centra albureau voor Schimmel cultures, Utrecht, The Netherlands, MFLU: Mae Fah Luang University, Chiang Rai, Thailand, CCMB: Bahia Culture Collection of Microorganisms, CMM: Culture Collection of Phytopathogenic Fungi “Prof. Maria Menezes,” Federal Rural University of Pernambuco, Brazil, MFLUCC: Mae Fah Luang University Culture Collection, Chiang Rai, Thailand, HKAS: The Herbarium Mycologium, Institute of Microbiology Chinese Academy of Sciences, Beijing, China, HUEFS: Herbarium of the State University of Feira de Santana, IRAN: Iranian Fungal Culture Collection, Iranian Research Institute of Plant Protection, Tehran, Iran, KUMCC: Kunming Institute of Botany Culture Collection, Kunming, China.

### Phylogenetic Analyses

Maximum Likelihood (ML) analysis was performed using RAxML-HPC2 on XSEDE (8.2.8) (Stamatakis 2014) in the CIPRES Science Gateway platform (Miller et al. 2010) using the GTR+I+G model of evolution. Bootstrap supports were obtained by running 1,000 pseudo-replicates. Bayesian analysis was conducted with MrBayes v. 3.1.2 (Huelsenbeck and Ronquist 2001) to evaluate Bayesian posterior probabilities (BYPP) (Rannala and Yang 1996, Zhaxybayeva and Gogarten 2002) by Markov Chain Monte Carlo sampling (BMCMC). GTR+I+G was used as the substitution model. Six simultaneous Markov chains were run for 2,000,000 generations and trees were sampled every 200th generation. The distribution of log-likelihood scores was examined to determine the stationary phase for each search and to decide if extra runs were required to achieve convergence, using the programme Tracer 1.5. The first 10% of generated trees were discarded and remaining 90% of trees were used to calculate posterior probabilities of the majority rule consensus tree. All trees were visualised in FigTree v.1.4.0 (Rambaut 2012) and the final layout (Fig. 1) was done with Microsoft PowerPoint (2013). The final alignment and tree were registered in TreeBASE under the submission ID: 27435. The final layout and tree were registered in TreeBASE under the submission ID: 27435 (http://purl.org/phylo/treebase/phylows/study/TB2:S27435?x-access-code=3101b93c442e7aa253174d89df7a500c&format=html).
Figure 1. Phylogram generated from Maximum Likelihood (RAxML) analysis, based on ITS-β-tubulin matrix. ML bootstrap supports (≥ 70%) and Bayesian posterior probability (≥ 0.95) are indicated as ML/BYPP. The tree is rooted to Kretzschmaria deusta (CBS 826.72) and Xyliaria hypoxylon (CBS 122620). Newly-generated strains are in red and type strains are in bold. The asterisks represent unstable species.
Taxon treatments

*Diatrypella* (Ces. & De Not.) De Not. 1863

Type species

*Diatrypella verruciformis* (Ehrh.) Nitschke (Fr.)

Description

Notes – *Diatrypella* was introduced by Cesati & De Notaris (1863) and is typified as *Diatrypella verruciformis* (Ehrh.) Nitschke. There are 146 epithets listed in Index Fungorum (2020). This genus was established to accommodate members of stromatic Sphaeriales which were characterised by ovoid and numerous ascospores and we introduce a new species viz. *Diatrypella longiasca*, based on new collections from China.

*Diatrypella longiasca* L.S. Dissan., J.C. Kang & K.D. Hyde, sp. nov.

- IndexFungorum [IF557952](#)
- Facesoffungi number [FoF09151](#)

Materials

Holotype:

- kingdom: Fungi; phylum: Ascomycota; class: Sordariomycetes; order: Xylariales; family: Diatrypaceae; taxonRank: species; genus: *Diatrypella*; specificEpithet: longiasca; scientificNameAuthorship: L.S. Dissanayake, J.C. Kang & K.D. Hyde, sp. nov.; country: China; stateProvince: Guizhou Province; county: Guiyang; locality: Guizhou University Garden (South); identifiedBy: L.S. Dissanayake; institutionID: HMAS 290656; collectionID: HMAS 290658; institutionCode: Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium; collectionCode: Kunming Institute of Botany Culture Collection; datasetName: CLD 42

Other material:

- type: isotype; institutionID: HMAS 290658; collectionID: KUMCC 20-0022; institutionCode: Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium; collectionCode: Kunming Institute of Botany Culture Collection

Description

Saprobic on dead twigs (Fig. 2). Sexual morph: Stromata 0.5–0.7 mm in diam., well-developed, solitary to gregarious, erumpent, black, immersed, globose to subglobose. Ascomata 525–540 μm high, 470–510 μm diam. (x̄ = 532 × 490 μm, n = 15), perithecial, surrounded by white entostroma, immersed in stromata, 2–5 perithecia arranged in a valsoid configuration, subglobose, individual ostiole with a long neck. Neck 180–190 μm long (x̄ = 185 μm, n = 15), cylindrical, with periphyses. Peridium 36–45 μm wide (x̄ = 40.5 μm, n = 20), composed of two layers: outer layer of black, thick-
walled cells; inner layer; hyaline, thick-walled cells forming textura angularis. Hamathecium 259–287 μm wide (\( \bar{x} = 273 \mu m, n = 20 \)), composed of cells 3–5 μm wide (\( \bar{x} = 4 \mu m, n = 20 \)), paraphyses arising from base of perithecia, hyaline, long, narrow, unbranched, septate, guttulate, narrowing and tapering towards apex. Asci 105–155 × 10–16 μm (\( \bar{x} = 130 \times 14 \mu m, n = 30 \)), polysporous, unitunicate, clavate, apically pointed, with a J-apical ring, long pedicellate (40–50 μm). Ascospores 4–8 × 1–2 μm (\( \bar{x} = 6 \times 1.5 \mu m, n = 30 \)), overlapping, hyaline, yellowish in mass, allantoid, aseptate, guttulate, smooth-walled. Asexual morph: Undetermined.

Culture characteristics – Colonies on PDA reaching 21 mm diam. after 2 weeks at 20–25°C, medium dense, circular to slightly irregular, slightly raised, cottony surface; colony from above: at first white, becoming buff; from below: yellowish white at margin, yellow to brown at centre; mycelium yellowish.

Etymology

The specific epithet longiasca refers to the long asci.

Notes

_Diatrypella longiasca_ shares similar characters with _D. vulgaris_ in having erumpent stromata through the bark often surrounded by remaining adherent epidermis or woody fragments and asci with many ascospores. However, _D. vulgaris_ is different from _D. longiasca_ in having longer ascospores (8–10 × 2–2.5 μm vs. 4–8 × 1–2 μm) (Trouillas et al. 2011). _Diatrypella vulgaris_ has 4–8 ascomata per stroma, while _D. longiasca_
comprises 2–5 ascomata per stromata. Comparison of the ITS 12% (73/570) and β-tubulin 13% (56/432) nucleotide differences, phylogenetic analyses and significant morphological differences indicate that *D. longiasca* and *D. vulgaris* are distinct taxa. Thus, *D. longiasca* is introduced as a new species in *Diatrypella*, based on its morphology coupled with high support values from the phylogenetic analysis (96% ML, 0.99 BYPP, Fig. 1). A key to species related to *Diatrypella longiasca* is given below.

**Paraeutypella** L.S. Dissan., J.C. Kang, Wijayaw. & K.D. Hyde, gen. nov.

- IndexFungorum [IF557954](#)
- Facesoffungi [FoF09231](#)

**Type species**

*Paraeutypella guizhouensis* L.S. Dissan., J.C. Kang & K.D. Hyde, sp. nov.

**Description**

Saprobic on dead twigs. **Sexual morph**: Stromata immersed in bark of dead branches, erumpent, solitary or aggregated. Ascomata with groups of 4–25 perithecia arranged in a valsoid configuration, surrounded by white, powdery entostroma, perithecial, black or brown, subglobose, clustered, immersed in stromata. Necks papillate, with an elongated ostiolar neck, central ostiolar canal filled with periphyses, 3–6 sulcate. Peridium composed of two layers of cells of textura angularis; inner layers cells hyaline or light brown, outer layers cells dark brown to black. Hamathecium composed of paraphyses arising from the base of perithecia, elongate, filiform, narrow, unbranched, septate, guttulate, narrowing and tapering towards apex. Asci 8-spored, unitunicate, thin-walled, clavate to cylindrical clavate or spindle-shaped, long pedicellate, apical rings J-. Ascospores overlapping biseriate, allantoid, slightly to moderately curved, allantoid, several oil droplets in each end, hyaline to light brown, sometimes yellow in mass, aseptate. **Asexual morph**: Coelomycetous. Conidiomata black, subconic, multiloculate, largely prosenchymatous, producing yellowish conidial masses. Conidiophores not recorded. Conidigenous cells cylindrical, tapering, arising from pseudoparenchyma or interwoven hyphae, proliferating percurrently or sympodially, rarely both ways. Conidia hyaline, single-celled, slightly to moderately curved, with flattened bases, becoming guttulate (description of asexual morph adapted from Glawe and Jacobs 1987).

**Etymology**

With reference to the morphological resemblance to *Eutypella*.

**Notes**

*Paraeutypella* is introduced to accommodate three species viz. *P. guizhouensis* sp. nov., as well as *P. citricola* and *P. vitis*, two species previously placed in *Eutypella*.
sensu lato. *Paraeutypella* is typified by *P. guizhouensis*, which was collected from undetermined dead twigs. *Paraeutypella* can be distinguished from *Eutypella* species by stromata with perithecia in groups of 4–25 arranged in a valsoid configuration, 3–6 sulcate, long ostiolar necks, while stromata of *Eutypella* comprise groups of 20–70 perithecia having comparatively shorter ostiolar necks with sulcate or smooth ostiolar necks. Strains of both genera appear in distinct clades in a phylogeny based on ITS and Beta tubulin data (Fig. 1), thereby justifying the erection of the new genus *Paraeutypella*. However, sequence data are not available for the type of *P. citricola* and *P. vitis*. A co-elomycetous asexual morph has been recorded for *P. vitis* as *Eutypella vitis* in culture (Glawe and Jacobs 1987).

**Paraeutypella guizhouensis** L.S. Dissan., J.C. Kang & K.D. Hyde, sp. nov.

- IndexFungorum IF557953
- Facesoffungi number FoF09148

**Materials**

**Holotype:**
- kingdom: Fungi; phylum: Ascomycota; class: Sordariomycetes; order: Xylariales; family: Diatrypaceae; genus: *Paraeutypella*; specificEpithet: *guizhouensis*; country: China; stateProvince: Guizhou Province; county: Guiyang; locality: Guizhou University Garden (North); habitat: Saprobic on dead twigs.; fieldNumber: CLD018; identifiedBy: L.S.Dissanayake; type: Holotype; institutionID: HMAS 290654; collectionID: KUMCC 20–0016; institutionCode: Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium; collectionCode: Kunming Institute of Botany Culture Collection; datasetName: CLD018

**Other material:**
- type: isotype; institutionID: HKAS 290655; collectionID: KUMCC 20-0017; institutionCode: Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium; collectionCode: Kunming Institute of Botany Culture Collection

**Description**

Saprobic on dead twigs (Fig. 3). **Sexual morph:** Stromata immersed in bark of dead branches, erumpent, aggregated, circular to irregular, superficial, carbonaceous. Ascomata 590–600 × 470–480 μm (x̄ = 595 × 475 μm, n = 10), perithecial, with groups of 6–12 perithecia arranged in a valsoid configuration, subglobose, clustered, immersed in stromata, ostiolate. Neck 400–418 μm long (x̄ = 409 μm, n = 10), papillate, central ostiolar canal filled with periphyses, 3–4 sulcate. Peridium 22–35 μm wide, composed of two layers of textura angularis; inner layer cells light brown to hyaline, outer layers cells dark brown to black. Hamathecium hyaline. Paraphyses 1–2 μm wide (x̄ = 1.5 μm, n = 10), arising from base of perithecia, long, narrow, unbranched, septate, guttulate, narrowing and tapering towards apex. Asci 55–80 × 5–9 μm (x̄ = 67.5 × 7 μm, n = 20), 8-spored, unitunicate, thin-walled, clavate to cylindrical clavate, long pedicellate (25–30 μm), with a J- apical ring. Ascospores 7–11 × 1–3 μm.
(\(\bar{x} = 9 \times 2 \mu m, n = 30\)), overlapping biseriate, allantoid, hyaline to light brown, smooth, aseptate, usually with 2–3 guttules. **Asexual morph**: Undetermined.

Culture characteristics – Colonies on PDA, reaching 21 mm diam. after 2 weeks at 20–25°C, medium dense, circular to slightly irregular, slightly raised, cottony surface; colony from above: at first white, becoming buff; from below: yellowish-white at margin, yellow to brown at centre; mycelium yellowish.

**Etymology**

The specific epithet *guizhouensis* refers to the locality in which the fungus was collected.

**Notes**

*Paraeutypella guizhouensis* resembles *P. vitis*, which comprises stromata that are erumpent through bark, with elongated perithecial necks and allantoid, slightly to moderately curved ascospores (Glawe and Jacobs 1987). However, *P. guizhouensis* differs from *P. vitis* in having comparatively longer ostiolar necks and longer asci (55–80 × 5–9 \(\mu m\)), while *P. vitis* has comparatively shorter ostiolar necks and shorter asci (40–46 × 6–8 \(\mu m\)) (Glawe and Jacobs 1987). *Paraeutypella vitis* (UCD2428TX) differs phylogenetically from our new taxon in 14% (80/576) base pairs in the ITS and 10% (42/405) base pairs in β-tubulin. Thus, *P. guizhouensis* is introduced as a new species in *Paraeutypella*, based on its morphology, base pair differences and phylogenetic analyses (94% ML, Fig. 1).
**Paraeutypella citricola** (Speg.,) L.S. Dissan., Wijayaw., J.C. Kang & K.D. Hyde, comb. nov. ≡ *Eutypella citricola* Speg., in Anales del Museo Nacional de Buenos Aires 6: 245, (1898)

- IndexFungorum [IF558003](https://www.indexfungorum.org/Names/NameRecord.aspx?id=107548968)
- Facesoffungi number [FoF09150](https://www.facesoffungi.org/)

**Nomenclature**

= *Eutypella citricola* Syd. & P. Syd., Hedwigia 49: 80 (1909), nom. illegit., Art. 53.1

**Materials**

**Holotype:**
- institutionID: LPS-2120

**Paratype:**
- kingdom: Fungi; phylum: Ascomycota; class: Sordariomycetes; order: Xylariales; family: Diatrypaceae; genus: *Paraeutypella*; specificEpithet: *citricola*; country: China; county: Guiyang; locationAccordingTo: Guizhou University Garden (South); year: 2019; month: October; day: 5; habitat: on a dead branch of *Acer* sp.; recordedBy: Nalin N. Wijayawardene; identifiedBy: L.S. Dissanayake; type: paratype; institutionID: HMAS 290660, HMAS 290659; collectionID: culture KUMCC 20–0024, KUMCC 20–0023; institutionCode: Chinese Academy of Science, Kunming and Chinese Academy of Science Herbarium; collectionCode: Kunming Institute of Botany Culture Collection

**Description**

Saprobic on dead twigs of *Acer palmatum* (Fig. 4). **Sexual morph**: Stromata immersed in bark of dead branches, erumpent, solitary or aggregated, circular to irregular in shape, superficial, carbonaceous. Ascomata 410–430 × 430–470 μm (x̄ = 420 × 450 μm, n = 10), perithecial, with groups of 4–6 perithecia arranged in a valsoid configuration, black, subglobose, clustered, immersed in ascostroma with ostiolar neck. Necks 360–390 μm long (x̄ = 375 μm, n = 10), papillate, sulcate, central ostiolar canal filled with paraphyses. Peridium 27–40 μm wide, composed of two layers of textura angularis; inner layer cells hyaline, outer layer cells dark brown to black. Hamathecium composed of 3–7 μm wide (x̄ = 5 μm, n = 10), hyaline, paraphyses arising from base of perithecium, composed of long, narrow, unbranched, septate, guttulate, narrowing and apically truncate. Asci 70–75 × 5–8 μm (x̄ = 72.5 × 6.5 μm, n = 20), 8-spored, unitunicate, thin-walled, clavate to cylindrical clavate, long pedicellate (40–50 μm), J-apical ring. Ascospores 7–10 × 2–3 μm (x̄ = 8.5 × 2.5 μm, n = 30), overlapping biseriate, allantoid, hyaline to light brown, smooth, aseptate, usually with guttules.

**Asexual morph**: Undetermined.
Culture characteristics – Colonies on PDA, reaching 21 mm diam. after 2 weeks at 20–25°C, medium dense, circular to slightly irregular, slightly raised, cottony surface; colony from above: at first white, becoming buff; from below: yellowish-white at margin, yellow to brown at centre; mycelium yellowish.

Notes

*Eutypella citricola* was described by Spegazzini (1898) from *Citrus* in Argentina and has since been reported to cause dieback on various woody plants in warm temperate and tropical regions (Farr and Rossman 2020). *Eutypella citricola* strains have previously been isolated from hosts such as *Citrus limon*, *C. sinensis*, *C. paradisi*, *Salix* spp., *Schinus molle*, *Ulmus procera* and *Vitis vinifera* (Trouillas et al. 2011, Mehrabi et al. 2016). In our study, we provide additional information for *P. citricola* from dead stems of *Acer* (Sapindaceae) in China. In morphology, our collection (HMAS 290660) resembles *Eutypella*, thus having pustulate stromata with stout, converging ostiolar necks and asci with eight spores. According to phylogenetic analysis, KUMCC 20–0024 closely groups with a collection of *E. citricola* (IRAN 2349C), which was collected on dead branches of *Salix* sp. (Salicaceae) in Gilan Province, Iran (Mehrabi et al. 2016) (Fig. 1). However, the IRAN 2349C strain is slightly different from our strain in having stromata with groups of 6–25 perithecia in a valsoid configuration and short ostiolar necks (100–300 µm), while our collection comprises stromata with groups of 4–6 perithecia in a valsoid configuration with a longer neck (356–385 µm). Based on phylogenetic analysis, both strains grouped in *Paraeutypella* sensu stricto (Fig. 1). Hence, the name *Eutypella citricola* is placed in *Paraeutypella* as *P. citricola*. 

Figure 4. doi

*Paraeutypella citricola* (HMAS 290660) a–c. stromata on substrate; d. cross section of stroma; e. vertical section through stroma showing ostiolar necks and perithecia; f. ostiolar canal; g. peridium; h. paraphyses; i–k. asci; l–n. ascospores; o. germinating ascospore; p, q. culture on PDA after 6 weeks from above and below. Scale bars: 500 µm (b–d), 200 µm (e–g), 20 µm (g–l), 5 µm (m–o).
Paraeutypella guizhouensis, the type of Paraeutypella, morphologically resembles P. citricola both having immersed stromata, perithecia each with a long ostiolar neck and allantoid, aseptate ascospores with an oil droplet at each end. However, Paraeutypella citricola differs from P. guizhouensis by the number of perithecia within a stroma (4–6 vs. 6–12). A comparison of the ITS 1.0% (6/576) and β-tubulin 1.2% (5/406) between KUMCC 20-0024 and IRAN 2340C, ITS 1.0% (6/576) and β-tubulin 1.0% (5/406) between KUMCC 20-0024 and HVGRF01, HVVIT07 has been made.

Paraeutypella vitis (Schwein.,) L.S. Dissan., J.C. Kang & K.D. Hyde, comb. nov. ≡ Sphaeria vitis Schwein., in Schr. Naturf. Ges. Leipzig 1: 39 (1822)

- IndexFungorum IF558004
- Facesoffungi number FoF09426

Nomenclature

= Valsa vitis (Schwein.) Fuckel, Jb. Nassau. Ver. Naturk. 23-24: 199 (1870)

= Engizostoma vitis (Schwein.) Kuntze, Revis. Gen. pl. (Leipzig) 3 (3): 475 (1898)

= Eutypella vitis (Schwein.) Ellis & Everh., The North American Pyrenomycetes: 490 (1892)

Notes

Eutypella vitis was collected from young shoots of grape vines in New York and was introduced by Ellis and Everhart (1982). According to our phylogenetic analyses, our new strain which represents Eutypella vitis (UCD 2291AR, USE2428TX) grouped as the sister clade (bootstrap support: 78% ML) to Paraeutypella citricola within Paraeutypella sensu stricto. Hence, in this study, we introduce the new combination, Paraeutypella vitis. Paraeutypella vitis shares similar morphologies to Paraeutypella species, such as having erumpent stromata through bark, 3–4 sulcate, long ostiolar necks, clavate asci, allantoid, slightly to moderately curved ascospores with several oil droplets in each end.

Identification keys

**Key to species similar to Diatrypella longiasca**

|   |   |   |
|---|---|---|
| 1 | Ascospores 4–5 μm long on average | Diatrypella major |
| – | Ascospores 6–10 μm long on average | 2 |
| 2 | Entostroma yellowish or olive-green | 3 |
Key to species of *Paraeutypella*

|   |   |   |
|---|---|---|
| 1 | Stromata immersed | *Paraeutypella citricola* |
|   | stromata erumpent | 2 |
| 2 | Short ostiolar neck and longer asci (55–80 × 5–9 μm) | *P. vitis* |
|   | Long ostiolar neck and shorter asci (40–46 × 6–8 μm) | *P. guizhouensis* |

Analysis

Phylogenetic analyses

The combined ITS and β-tubulin matrix comprises 79 sequences that represents the genera in *Diatrypaceae* including the outgroup taxa. The best scoring RAxML tree is shown (Fig. 1) with a final ML optimisation likelihood value of -12709.069416. The matrix had 784 distinct alignment patterns, with 28.77% undetermined characters or gaps. Estimated base frequencies were: A = 0.226868, C = 0.263622, G = 0.232845, T = 0.27666; substitution rates AC = 1.218567, AG = 2.693651, AT = 1.272423, CG = 0.850048, CT = 3.427431, GT = 1.000000; proportion of invariable sites I = 0.100328; gamma distribution shape parameter α = 0.775027. All trees (ML and BYPP) were similar in topology and did not differ in generic relationships, which are in agreement with multi-gene phylogenies of previous studies.

Species of *Eutypella* are polyphyletic in our phylogram, while new isolates KUMCC 20-0023 and KUMCC 20-0024 grouped in a clade that comprises *Eutypella citricola* Syd. &
P. Syd. and *Eutypella vitis* (Schwein.) Ellis & Everh. (Fig. 1). KUMCC 20-0016 and KUMCC 20-0017 formed a separate clade basal to *E. vitis* with high statistical support (94% ML) (Fig. 1). These species form a separate clade from the *Eutypella* clade. A novel genus is needed to accommodate these species, hence we introduce *Paraeutypella*.

Our new strains KUMCC 20-0021 and KUMCC 20-0022 are accommodated within *Diatrypella* with high statistical support (96% ML, 1.00 BYPP) as a distinct lineage.

**Discussion**

This study introduces a new genus, *Paraeutypella* and accepts 22 genera in Diatrypaceae. According to the previous analyses of combined ITS and β-tubulin sequence data, the genus *Eutypella* has been often identified as polyphyletic in *Diatrypaceae* (Trouillas et al. 2011, Mehrabi et al. 2016, Mehrabi et al. 2019, Dayarathne et al. 2016, Dayarathne et al. 2020a, Dayarathne et al. 2020b) and determined in our study as well (Fig. 1). The type of *Eutypella*, *E. cerviculata* (Fr.) Sacc. grouped with *E. semicircularis* S. Chacón & M. Piepenbr., *E. persica* Mehrabi et al. and *E. quercina* Mehrabi et al.

*Eutypella citricola* groups separately from *Eutypella* sensu stricto with *Eutypella vitis* and our newly-generated strains. These new strains are introduced as a new genus, *Paraeutypella* with three species viz. *P. citricola*, *P. guizhouensis* and *P. vitis*. We studied the morphological characteristics of the species belonging to this clade and found considerable morphological differences from *Eutypella* sensu stricto. The differences include stromata with 4–25 groups of perithecia in a valsoid configuration, 3–6 sulcate, long ostiolar necks; thus, we consider them to belong in a distinct genus from the *Eutypella* and hence, we introduce the novel *Paraeutypella*.

There does not appear to be any host-specificity since members of Diatypaceae are found on a wide range of hosts in various habitats. Diatypaceae species frequently have been identified as saprobics on the decaying wood of angiosperms. Few endophytes, such as *Diatrypella frostii* Peck and *Peroneutypa scoparia* (Schwein.) Carmarán & A.I. Romero, have been reported (de Errasti et al. 2010, Vieira et al. 2011, Grassi et al. 2014). Therefore, the family may have the potential for switching nutritional modes during the degradation of plant material (de Errasti et al. 2010, Grassi et al. 2014). Several species have been reported as pathogens, such as *Cryptosphaeria populina* (Pers.) Sacc., *C. pullmanensi* Glawe and *Eutypella parasitica* R.W. Davidson & R.C. Lorenz, causing canker disease (Glawe and Rogers 1984, Rappaz 1987, Ma et al. 2016), *Crytovalsa ampelina* (Nitschke) Fuckel causing grapevine trunk disease (Luque et al. 2006), *Eutypa lata* (Pers.) Tul. & C. Tul. causing canker and dieback disease (Lardner et al. 2005) and *E. leptoplaca* (Durieu & Mont.) Rappaz contributing to the dieback of grapevines (Trouillas and Gubler 2004, Catal et al. 2007).

In our phylogenetic analyses, some species of *Diatrypella*: *D. favacea* (Fr.) Ces. & De Not., *D. iranensis* Mehrabi & Hemmati, *D. macrospora* Mehrabi et al. and *D. pulvinata* Nitschke formed a distinct lineage (Fig. 1) in *Diatrypella*. Similarly, some species of *Eutypella* (*E.*
caricae (De Not.) Berl., E. parasitica R.W. Davidson & R.C. Lorenz and E. virescens Wehm.) often form distinct lineages within Diatrypaceae (Fig. 1). This may be due to lack of single-copy nuclear genes like β-tubulin or misidentified species.

Acknowledgements

This work was funded by grants of the National Natural Science Foundation of China (NSFC Grants Nos. 31670027 & 31460011). Dr. Shaun Pennycook is thanked for the nomenclatural advice. Nalin N. Wijayawardene gratefully acknowledges Natural Science Foundation of China (grant No. NSFC 31950410558) and grant FAMP201906K provided by the State Key Laboratory of Functions and Applications of Medicinal Plants, Guizhou Medical University. Dong-Qin Dai thanks the National Natural Science Foundation of China (NSFC 31760013) and the Thousand Talents Project of Yunnan Provinces for support. Monika C. Dayarathne would like to thank National Natural Science Foundation of China (No. 31972222, 31560489). Lakmali S. Dissanayake would like to thank Ms. D.S. Marasinghe and Ms. S. N. Wijesinghe for valuable suggestions and guidance.

References

• Acero FJ, Gonzalez V, Sanchez-Ballesteros J, Rubio V, Checa J, Bills G, Salazar O, Platas G, Pelaez F (2004) Molecular phylogenetic studies on the Diatrypaceae based on rDNA-ITS sequences. Mycologia 96 (2). https://doi.org/10.2307/3762061
• Arhipova N, Gaitnieks T, Donis J, Stenlid J, Vasaitis R (2012) Heart-rot and associated fungi in Alnus glutinosae stands in Latvia. Scandinavian Journal of Forest Research 27 (4): 327-336. https://doi.org/10.1080/02827581.2012.670727
• Catal M, Jordan SA, Butterworth SC, Schilder AM (2007) Detection of Eutypa lata and Eutypella vitis in grapevine by nested multiplex polymerase chain reaction. Phytopathology 97: 737-747. https://doi.org/10.1094/PHYTO-97-6-0737
• Cesati V, De Notaris G (1863) Schema di classificazione degli sferiacei italici aschigeri: più o meno appartenenti al genere Sphaerianell’antico significato attribuitogli da Persoo. Commentario della Società Crittogamologica Italiana 1: 205.
• Chacón S, Dörge D, Weisenborn J, Piepenbring M (2013) A new species and a new record of Diatrypaceae from Panama. Mycologia 105: 681-688. https://doi.org/10.3852/12-131
• Chalkley DB, Su SO, Kohlmeye BV, Kohlmeyer J, Zhou JJ (2010) Diatrypasimilis australiensis, a novel xylarialean fungus from mangrove. Mycologia 102: 430-437. https://doi.org/10.3852/09-142
• Chomnunti P, Hongsanan S, Hudson BA, Tian Q, Peršoh D, Dhami MK, Alias AS, Xu JC, Liu XZ, Stadler M, Hyde KD (2014) The sooty moulds. Fungal Diversity 66: 1-36. https://doi.org/10.1007/s13225-014-0278-5
• Dai D, Phookamsak R, Wijayawardene NN, Li WJ, Bhat J, Chukeatirote E, Bahkali AH, Zhao RL, Xu JC, Hyde KD (2016) Bambusicolous fungi. Fungal Diversity 82: 1-105. https://doi.org/10.1007/s13225-016-0367-8
• Dayarathne MC, Phookamsak R, Hyde KD, Manawasinghe IS, To-anun C, Jones EB (2016) Halodiatrype, a novel diatrypaceous genus from mangroves with H. salinicola and H. avicenniae spp. nov. Mycosphere 7 (5): 612-627. https://doi.org/10.5943/mycosphere/7/5/7

• Dayarathne MC, Jones EBG, Maharachchikumbura SSN, Devadatha B, Sarma VV, Khongphinittibunjong K, Chomnunti P, Hyde KD (2020a) Morpho-molecular characterization of microfungi associated with marine based habitats. Mycosphere 11 (1): 1-188. https://doi.org/10.5943/mycosphere/11/1/1

• Dayarathne MC, Wanasinghe D, Devadatha B, Abeywickrama P, Gareth Jones EB, Chomnunti P, Sarma VV, Hyde K, Lumyong S, Mckenzie EC (2020b) Modern taxonomic approaches to identifying diatrypaceous fungi from marine habitats, with a novel genus Halocryptovalsa Dayarathne & K.D.Hyde, gen. nov. Cryptogamie, Mycologie 41 (3). https://doi.org/10.5252/cryptogamie-mycologie2020v41a3

• de Almeida DAC, Gusmão LFP, Miller AN (2016) Taxonomy and molecular phylogeny of Diatrypaceae (Ascomycota, Xylariales) species from the Brazilian semi-arid region, including four new species. Mycological Progress 15 (6). https://doi.org/10.1007/s11557-016-1194-8

• de Errasti A, Carmarán CC, Novas MV (2010) Diversity and significance of fungal endophytes from living stems of naturalized trees from Argentina. Fungal Diversity 41 (1): 29-40. https://doi.org/10.1007/s13225-009-0012-x

• Ellis JB, Everhart BM (1982) Mycologic botany, Phrenomygetes. Published by Ellis and Everhart New Field, New Jersey, 490 pp.

• Farr DF, Rossman AY (2020) Fungal databases, U.S. National Fungus Collections, ARS, USDA. https://nt.ars-grin.gov/fungaldatabases/. Accessed on: 2020-12-10.

• Glawe A, Jacobs KA (1987) Taxonomic notes on Eutypella vitis, Cryptosphaeria populina, and Diatrype stigma. Mycologia 79: 135-139. https://doi.org/10.1080/00275514.1987.12025379

• Glawe DA, Rogers JD (1984) Diatrypaceae in the Pacific Northwest. Mycotaxon 20: 401-460.

• Grassi E, Belen Pildain M, Levin L, Carmaran C (2014) Studies in Diatrypaceae: the new species Eutypa microasca and investigation of ligninolytic enzyme production. Sydowia 66: 99-114. https://doi.org/10.12905/0380.sydowia66(1)2014-0099

• Hall TA (1999) BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series. 41: 95-98.

• Huelsenbeck JP, Ronquist F (2001) MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics 17: 754-755. https://doi.org/10.1093/bioinformatics/17.7.754

• Hyde KD, Tennakoon DS, Jeewon R, Bhat DJ, Maharachchikumbura SSN, Rossi W, Leonardi M, Lee HB, Mun HY, Houbraken J, Nguyen TTT, Jeon SJ, Frisvad JC, Dhanushka N, Wanasinghe DN, Luücking R, Aptroot A, Cáceres MES, Karunarathna SC, Hongsanan S, Phookamsak R, de Silva NI, Thambugala KM, Jayawardena RS, Senanayake IC, Boonmee S, Chen J, Luo ZL, Phukhamsakda C, Pereira OL, Abreu VP, Rosado AW, Bart B, Randrianjohany E, Hofstetter V, Gibertoni TB, da Silva Soares AM, Plautz HL Jr, Sotão HMP, Xavier WKS, Bezerra JDP, de Oliveira TGL, de Souza-Motta CM, Magalhães OMC, Bundhun D, Harishchandra D, Manawasinghe IS, Dong W, Zhang SN, Bao DF, Samaraoke MC, Pem D, Karunarathna A, Lin CG, Yang J, Perera RH, Kumar V, Huang SK, Dayarathne MC, Ekanayaka AH, Jayasiri SC, Xiao YP, Konta S, Niskanen T, Liimatainen K, Dai YC, Ji XH, Tian XM, Mešić A, Singh SK,
Phutthacharoen K, Cai L, Sorvongxay T, Thiyagaraja V, Norphanphoun C, Chaiwan N, Lu YZ, Jiang HB, Réblová M, Fournier J, Nekvindová J, do Nascimento Barbosa R, dos Santos JEF, de Oliveira NT, Li GJ, Ertz D, Phillips AJL, Kuo CH, Camporesi E, Bulgakov TS, Lumyong S, Jones EBG, Chomnunti P, Gentekaki E, Bungartz F, Zeng XY, Fryar S, Tkalčec Z, Liang J, Li GS, Wen TC, Singh PN, Gafforov Y, Promputtha I, Yasanthika E, Goonasekara ID, Zhao Q, Kirk PM, Liu JK, Yan JY, Mortimer PE, Xu JC (2019) Fungal diversity notes 1036–1150: taxonomic and phylogenetic contributions on genera and species of fungal taxa. Fungal Diversity 96: 1-242. https://doi.org/10.1007/s13225-019-00429-2

• Hyde KD, Norphanphoun C, Maharachchikumbura SSN, Bhat DJ, Jones EBG, Bundhun D, Chen YJ, Bao DF, Boonmee S, Calabon MS, Chaiwan N, Chethana KWT, Dai DQ, Dayarathe MC, Devadatha B, Dissanayake AJ, Dissanayake LS, Doilom M, Dong W, Fan XL, Goonasekara ID, Hongsanan S, Huang SK, Jayawardena RS, Jeewon R, Karunarathna A, Konta S, Kumar V, Lin CG, Liu JK, Liu N, Luangsa-ard J, Lumyong S, Luo ZL, Marasinghe DS, McKenzie EHC, Niego AGT, Nirankan M, Perera RH, Phukhamsakda C, Rathnayaka AR, Samarakoon MC, Samarakoon SMBC, Sarma VV, Senanayake IC, Shang QJ, Stadler M, Tibpromma S, Wei DP, Wijayawardene NN, Xiao YP, Xiang MM, Yang J, Zeng XY, Zhang SN (2020) Refined families of Sordariomycetes. Mycosphere 11: 305-1059. https://doi.org/10.5943/mycosphere/11/1/7

• Jayasiri SC, Hyde KD, Jeewon R, Jones EBG, Kirk P, Lei C, Liu JK, Maharachchikumbura SSN, McKenzie E, GhabadNejhad M, Nilsson H, Pang KL, Phookamsak R, Rollins AW, Romero AI, Stephenson S, Suotrong S, Tsui CKM, Vizzini A, Wen TC, De Silva NI, Promputtha I, Kang JC (2015) The faces of fungi database: fungal names linked with morphology, molecular and human attributes. Fungal Diversity 74: 3-18. https://doi.org/10.1007/s13225-015-0351-8

• Konta S, Maharachchikumbura SSN, Senanayake IC, McKenzie EHC, Stadler M, Boonmee S, Phookamsak R, Jayawardena RS, Senwanna C, Hyde KD, Elgorban AM, Eungwanichayapant PD (2020) A new genus Allodiatrype, five new species and a new host record of diatrypaceous fungi from palms (Arecaceae). Mycosphere 11 (1): 239-268. https://doi.org/10.5943/mycosphere/11/1/4

• Lardner R, Stummer BE, Sosnowski MR, Scott ES (2005) Molecular identification and detection of Eutypa lata in grapevine. Mycological Research 109: 799-808. https://doi.org/10.1017/S0953756205002893

• Luque J, Sierra D, Torres E, Garcia F (2006) Cryptovalsa ampelina on grapevines in NE Spain: identification and pathogenicity. Phytopathologia Mediterranea 45: 101-109. https://doi.org/10.1520/STP37480S

• Luque J, Garcia-Figuieres F, Legorburu FJ, Muruamendiaraiz A, Armengol J, Trouillas FP (2012) Species of Diatrypaceae associated with grapevine trunk diseases in Eastern...
Spain. Phytopathologia Mediterranea 51: 528-540. https://doi.org/10.14601/Phytopathol_Mediterr-9953

- Lynch SC, Eskalen A, Zambino PJ, Mayorquin JS, Mayorquin JS, Wang DH (2013) Identification and pathogenicity of Botryosphaeriaceae species associated with coast live oak (Quercus agrifolia) decline in southern California. Mycologia 105: 125-140. https://doi.org/10.3852/12-047

- Ma R, Zhu Y-, Fan X-, Tian C- (2016) Canker disease of willow and poplar caused by Cryptosphaeria pullmanensis recorded in China. Forest Pathology 46 (4): 327-335. https://doi.org/10.1111/epf.12261

- Mehrabi M, Hemmati R, Vasilyeva LN, Trouillas FP (2015) A new species and a new record of Diatrypaceae from Iran. Mycosphere 6 (1): 60-68. https://doi.org/10.5943/mycosphere/6/1/7

- Mehrabi M, Hemmati R, Vasilyeva LN, Trouillas FP (2016) Diatrypella macrospora sp. nov. and new records of diatrypaceous fungi from Iran. Phytotaxa 252 (1). https://doi.org/10.11646/phytotaxa.252.1.4

- Mehrabi M, Asgari B, Hemmati R (2019) Two new species of Eutypella and a new combination in the genus Peroneutypa (Diatrypaceae). Mycological Progress 18: 1057-1069. https://doi.org/10.1007/s11557-019-01503-4

- Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for inference of large phylogenetic trees. New Orleans, Louisiana, November, 1–8. In Gateway Computing Environments Workshop 2010 (GCE). https://doi.org/10.1109/GCE.2010.5676129

- Nitschke TR (1867) Pyrenomycetes Germanici Die Kernpilze Deutschlands Bearbeitet von. [Dr. Th. Nitschke]. Erster Band, Erste Lieferung i-ii. Germany, Bresalau, 156 pp. [In Verlag von Eduard Trewendt].

- Nylander JA (2004) MrModeltest v.2.2. Program distributed by the author: 2. Evolutionary Biology Centre. Uppsala University 1–2.

- O'Donnell K, Cigelnik E (1997) Two divergent intragenomic rDNA ITS2 types within a monophyletic lineage of the fungus Fusarium Are Nonorthologous. Molecular Phylogenetics and Evolution 7 (1): 103-116. https://doi.org/10.1006/mpev.1996.0376

- Paolinelli-Alfonso M, Serrano-Gomez C, Hernandez R (2015) Occurrence of Eutypella microtheca in grapevine cankers in Mexico. Phytopathologia Mediterranea 54: 86-93.

- Pazoutova S, Srukta P, Holusa J, Chudickova M, Kubatova A, Kolarik M (2012) Liberomyces gen. nov. with two new species of endophytic coelomycetes from broadleaf trees. Mycologia 104: 198-210. https://doi.org/10.3852/11-081

- Peršoh D, Melcher M, Graf K, Fournier J, Stadler M, Rmbold G (2009) Molecular and morphological evidence for the delimitation of Xylaria hypoxylon. Mycologia 101: 256-268. https://doi.org/10.3852/08-108

- Phukhamsakda C, McKenzie EHC, Phillips AJL, Gareth Jones EB, Jayarama Bhat D, Stadler M, Bhunjun CS, Wasasinghe DN, Thongbai B, Camporesi E, Ertz D, Jayawardena RS, Perera RH, Ekanayake AH, Tibpromma S, Doilom M, Xu J, Hyde KD (2020) Microfungi associated with Clematis (Ranunculaceae) with an integrated approach to delimiting species boundaries. Fungal Diversity 102 102: 1-203. https://doi.org/10.1007/s13225-020-00448-4

- Rambaut A (2012) Fig.Tree. Tree Fig. Drawing Tool. v. 1.4.0. URL: http://tree.bio.ed.ac.uk/software/fgtree/.
• Rannala B, Yang Z (1996) Probability distribution of molecular evolutionary trees: a new method of phylogenetic inference. Journal of Molecular Evolution 43: 304-311. https://doi.org/10.1007/BF0238839
• Rappaz F (1987) Taxonomie et nomenclature des Diatrypaceae à asques octospores. Mycologia Helvetica 2: 285-648.
• Rolshausen PE, Mahoney NE, Molyneux RJ, Gubler WD (2006) A reassessment of the species concept in *Eutypa lata*, the causal agent of *Eutypa* dieback of grapevine. Phytopathology 96: 369-377. https://doi.org/10.1094/PHYTO-96-0369
• Sales R, Santana CVS, Nogueira DRS, Silva KJP, Guimarães IM, Michereff SJ, Abad-Campos P, García-Jiménez J, Armengol J (2010) First report of *Monosporascus cannonballus* on watermelon in Brazil. Plant Disease 94 (2): 278-278. https://doi.org/10.1094/pdis-94-2-0278b
• Senanayake IC, Maharachchikumbura SN, Hyde KD, Bhat JD, Jones EG, McKenzie EH, Dai DQ, Daranagama DA, Dayarathne MC, Goonasekara ID, Konta S, Li WJ, Shang QJ, Stadler M, Wijayawardene NN, Xiao YP, Norphanphoun C, Liu XY, Bahkali AH, Kang JC, Wang Y, Wen TC, Wendt I, Xu JC, Camporesi E (2015) Towards unraveling relationships in Xylariomycetidae (Sordariomycetes). Fungal Diversity 73: 73-144. https://doi.org/10.1007/s13225-015-0340-y
• Senwanna C (2017) Novel taxa of Diatrypaceae from para rubber (*Hevea brasiliensis*) in northern Thailand; introducing a novel genus *Allocryptovalsa*. Mycosphere 8 (10): 1835-1855. https://doi.org/10.5943/mycosphere/8/10/9
• Shang QJ, Hyde K, Jeewon R, Khan S, Promputtha I, Phookamsak R (2018) Morpho-molecular characterization of *Peroneutypa* (Diatrypaceae, Xylariales) with two novel species from Thailand. Phytotaxa 356 (1). https://doi.org/10.11646/phytotaxa.356.1.1
• Spegazzini C (1898) Fungi Argentini novi v. critici. Anales Museo Nacional Buenos Aires 6: 1-23.
• Stamatakis A (2014) RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30 (9): 1312-1313. https://doi.org/10.1093/bioinformatics/btu033
• Thambugala K, Daranagama D, Phillips AL, Bulgakov T, Bhat D, Camporesi E, Bahkali A, Eungwanichayapant P, Liu Z, Hyde K (2016) Microfungi on *Tamarix*. Fungal Diversity 82 (1): 239-306. https://doi.org/10.1007/s13225-016-0371-z
• Thiagaraja V, Senanayake IC, Wanasinghe DN, Karunarathna SC, Worthy FR, To-Anun C (2019) Phylogenetic and morphological appraisal of *Diatrype lijiangensis* sp. nov. (Diatrypaceae, Xylariales) from China. Asian Journal of Mycology 2: 198-208. https://doi.org/10.5943/ajom/2/1/10
• Trouillas F, Úrbez-Torres J, Gubler W (2010) Diversity of diatrypaceous fungi associated with grapevine canker diseases in California. Mycologia 102 (2): 319-336. https://doi.org/10.3852/08-185
• Trouillas F, Pitt W, Sosnowski M, Huang R, Peduto F, Loschiavo A, Savocchia S, Scott E, Gubler W (2011) Taxonomy and DNA phylogeny of Diatrypaceae associated with *Vitis vinifera* and other woody plants in Australia. Fungal Diversity 49 (1): 203-223. https://doi.org/10.1007/s13225-011-0094-0
• Trouillas FP, Gubler WD (2004) Identification and characterization of *Eutypa leptoplaca*, a new pathogen of grapevine in Northern California. Mycological Research 108: 1195-1204. https://doi.org/10.1017/S0953756204000863
• Trouillas FP, Hand FP, Inderbitzin P, Gubler WD (2015) The genus Cryptosphaeria in the western United States: taxonomy, multilocus phylogeny and a new species, C. multicontinentalis. Mycologia 107 (6): 1304-1313. https://doi.org/10.3852/15-115

• Úrbez-torres JR, Peduto F, Striegler RK, Urrearomero KE, Rupe JC, Cartwright RD, Gubler WD (2012) Characterization of fungal pathogens associated with grapevine trunk diseases in Arkansas and Missouri. Fungal Diversity 52: 169-189. https://doi.org/10.1007/s13225-011-0110-4

• Úrbez-Torres JR, Gubler WD (2009) Pathogenicity of Botryosphaeriaceae species isolated from grapevine cankers in California. Plant Disease 93 (6): 584-592. https://doi.org/10.1094/pdis-93-6-0584

• Vasilyeva LN, Stephenson SL (2005) Pyrenomycetes of the Great Smoky Mountains National Park. II. Cryptovalsa Ces. et De Not. and Diatrypella (Ces. et De Not.) Nitschke. Fungal Diversity 1-12.

• Vieira M, Hughes A, Gil V, Vaz A, Alves T, Zan C, Rosa CA, Rosa LH (2011) Diversity and antimicrobial activities of the fungal endophyte community associated with the traditional Brazilian medicinal plant Solanum cernuum Vell. (Solanaceae). Canadian Journal of Microbiology 58: 54-56. https://doi.org/10.1139/w11-105

• Vieira ML, Hughes AF, Gil VB, Vaz AB, Alves TM, Zani CL, Rosa CA, Rosa LH (2012) Diversity and antimicrobial activities of the fungal endophyte community associated with the traditional Brazilian medicinal plant Solanum cernuum Vell. (Solanaceae). Canadian Journal of Microbiology 58: 54-66. https://doi.org/10.1139/w11-105

• Vu D, Groenewald M, de Vries M, Gehrmann T, Boekhout T, Crous PW, Robert V, Verkley GJ (2019) Large-scale generation and analysis of filamentous fungal DNA barcodes boosts coverage for kingdom fungi and reveals thresholds for fungal species and higher taxon delimitation. Study in Mycology 92: 135-154. https://doi.org/10.1016/j.simyco.2018.05.001

• White TJ, Bruns T, Lee J, Taylor SB (1990) Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In: Innis MA, Gelfand DH, Sninsky JJ, White TJ (Eds.) (Eds.) PCR protocols: a guide to methods and applications. Academic Press, San Diego, California, USA., 315–322 pp. https://doi.org/10.1016/B978-0-12-372180-8.50042-1

• Wijayawardene NN, Hyde KD, Al-Ani LKT, Tedersoo L, Haelewaters D, Rajeshkumar KC, Zhao RL, Aptroot A, Leontyev DV, Saxena RK, Tokarev YS, Dai DQ, Letcher PM, Stephenson SL, Ertz D, Lumbsch HT, Kukwa M, Issi IV, Madrid H, Phillips AJL, Selbmann L, Pflieger WP, Horváth E, Kirk P, Kolaříková Z, Raja HA, Radek R, Papp V, Dima B, Ma J, Malosso E, Takamatsu S, Rambold G, Kannibal PB, Triebel D, Gautam AK, Avasthi S, Suetrong S, Timdal E, Fryar SC, Delgado G, Réblóvá M, Doirom M, Dolatabadi S, Pawłowska J, Humber RA, Kodseub R, Sánchez-Castro I, Goto BT, Silva DKA, De Souza FA, Oehl F, Da Silva GA, Silva IR, Blaszkowski J, Jobim K, Maia LC, Barbosa FR, Fiuza PO, Divakar PK, Shenoy BD, Somrithipol S, Karunarathna SC, Tibromma S, Mortimer PE, Wanasinghe DN, Phookamsak R, Xu J, Wang Y, Fenghua T, Alvarado P, Li DW, Kušan I, Matočec N, Maharachchikumbura SSN, Papizadeh M,
Heredia G, Wartchow F, Bakhshi M, Boehm E, Youssef N, Hustad VP, Lawrey JD, Santiago ALCMA, Souza-Motta CM, Firmino AL, Tian Q, Houbraken J, Hongsanan S, Tanaka K, Dissanayake AJ, Monteiro JS, Grossart HP, Suija A, Weerakoon G, Etayo J, Tsurykau A, Kuhnert E, Vázquez V, Mungai P, Damm U, Li QR, Zhang H, Boonmee S, Lu YZ, Becerra AG, Kendrick B, Brearley FQ, Motiejūnaitė J, Sharma B, Khare R, Gaikwad S, Wijesundara DSA, Tang LZ, He MQ, Flakus A, Rodriguez-Flakus P, Zhurbenko MP, McKenzie EHC, Stadler M, Bhat DJ, Liu JK, Raza M, Jeewon R, Nassonova ES, Prieto M, Jayalal RGU, Yurkov A, Schnittler M, Shchepin ON, Novozhilov YK, Cavender JC, Kang Y, Mohammad S, Zhang LF, Xu RF, Li YM, Dayarathne MC, Ekanayaka AH, Wen TC, Deng CY, Lateef AA, Pereira OL, Navathe S, Hawksworth DL, Fan XL, Dissanayake LS, Erdoðdu M (2020) Outline of fungi and fungus-like taxa. Mycosphere 11: 1060-1456. https://doi.org/10.5943/mycosphere/11/1/8

- Wu ZH, Wang TH, Huang W, Ou YB (2001) A simplified method for chromosome DNA preparation from filamentous fungi. Mycosistema 20: 575-577.

- Zhaxybayeva O, Gogarten JP (2002) Bootstrap, Bayesian probability and maximum likelihood mapping: exploring new tools for comparative genome analyses. MBC genomics 3: 4. https://doi.org/10.1186/1471-2164-3-4