Two new species in a new genus and a critical revision of Brachybasidiaceae (Exobasidiales, Basidiomycota) in honor of Franz Oberwinkler

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Abstract

The Brachybasidiaceae are a family of 22 known species of plant-parasitic microfungi belonging to Exobasidiales, Basidiomycota. Within this family, species of the largest genus *Kordyana* develop balls of basidia on top of stomatal openings. Basidial cells originate from fungal stroma filling substomatal chambers. Species of *Kordyana* typically infect species of Commelinaceae. During fieldwork in the neotropics, fungi morphologically similar to *Kordyana* spp. were found on *Goeppertia* spp. (syn. *Calathea* spp., Marantaceae), namely on *G. panamensis* in Panama and on *G. propinqua* in Bolivia. These specimens are proposed as representatives of a genus new to science, *Marantokordyana*, based on the distinct host family and molecular sequence data of ITS and LSU rDNA regions. The specimens on the two host species represent two species new to science, *M. oberwinkleriana* on *G. panamensis* and *M. boliviana* on *G. propinqua*. They differ by the size and shape of their basidia, molecular sequence data of ITS and LSU rDNA regions, and host plant species. In the past, the understanding of Brachybasidiaceae at order and family level was significantly improved by investigation realized by Franz Oberwinkler and his collaborators at the University of Tübingen, Germany. On species level, however, our knowledge is still very poor due to incomplete species descriptions of several existing names in literature, scarceness of specimens, as well as sequence data lacking for many taxa and for further barcode regions. Especially species of *Kordyana* and species of *Dicellomyces* are in need of revision.

Keywords Bolivia · *Calathea* · *Dicellomyces* · *Kordyana* · Marantaceae · Panama

Introduction

The Exobasidiomycetes (Ustilaginomycotina, Basidiomycota) include orders traditionally considered smut fungi because of the presence of thick-walled probasidia (teliospores), like in Doassansiales, Entylomatales, Georgfischeriales, and Tilletiales. This class also includes fungi without teliospores mainly in Ceraceosorales, Exobasidiales, and Microstromatales (Begerow et al. 2002, 2018). On the basis of different types of soral structures, basidia, and ultrastructural characteristics, Bauer et al. (1998, 2001) distinguished four families within the Exobasidiales, namely Brachybasidiaceae, Cryptobasidiaceae, Exobasidiaceae, and Graphiolaceae. These groups have also been treated as orders in the past as well as recently by Oberwinkler (2012a, b). Distinctive characteristics relevant for the concept of Exobasidiales and morphologically closely related...
orders, for families of Exobasidiales, and for genera of Brachybasidiaceae are summarized in Table 1, following the way of presentation used by de Beer et al. (2006) for Microstromatales. For species included in the genera of Brachybasidiaceae, host species, and known geographic distribution, see Online Resource 1. Selected species of

**Exobasidiomycetes**
- parasitic on plants, mostly on leaves
- mostly with holobasidia, often growing out of thick-walled probasidia (telosporas)
- asexual growth as yeasts or with hyphae and yeast-like conidia
- host-parasite interaction with complex interaction apparatus

| Exobasidiomycetes | Ceraceosorales | Microstromatales |
|-------------------|----------------|------------------|
| **basidia**       | - without telosporas | - without telosporas |
| - protruding through epidermis or stomata | - protruding through epidermis or stomata | - protruding through stomata |
| - with 2-4-8 basidiospores mostly on stigmatum | - forming sorus-like hymenium | - carrying more than 2, mostly 6 basidiospores directly on the basidial cell (gastroid) |
| basidiospores | - septate upon germination | - septate upon germination |
| - germinate with hyphae and conidia | - germinate with hyphae and conidia | - germinate with probasidal sacs |
| ultrastructure | - with interaction rings | - with interaction rings |

**Brachybasidiaceae**
- mostly monocots
- without galls
- hosts: Arecaceae
- basidia: in suprastomatall balls
- without paraphyses
- conidia: unknown

**Crypobasidiaceae**
- dicots, mostly Lauraceae
- with galls of various host organs
- basidia: developing inside host tissue
- carrying 4 or more basidiospores mostly directly on the basidial cell (gastroid)
- no probasidal sacs
- basidiospores: blastospore
- thick-walled
- ornamented
- slightly pigmented

**Kordyana** (7+3 doubtful species)
- Commelinaceae (Bigononiaceae, Bursariaceae, Poaceae)
- basidia: in suprastomatall balls
- sometimes with paraphyses
- some species with probasidal sacs in substomatall chambers
- conidia: fusiform, globose, or subpyriform

**Marantokordyana** (2 species)
- Marantaceae
- basidia: in suprastomatall balls
- with paraphyses
- with probasidal sacs in substomatall chambers
- conidia: fusiform

**Meira** (6 species)
- isolated from various substrates
- saprotrophic
- basidia: absent (asexual stage)
- conidia: fusiform

**Dicellomyces** (1+1 doubtful sp.)
- Poaceae (Arecaceae)
- basidia: in discoid, gelatinous basidioecarps breaking through epidermis
- thin-walled persistent probasidia
- without paraphyses
- conidia: globose

**Dicellomyces” scirpi** (1 sp.)
- Cyperaceae
- basidia: in gelatinous basidioecarps breaking through epidermis
- swollen, not persistent probasidia
- with paraphyses
- conidia: allantoid

**Exobasidiellum** (1 species)
- Poaceae
- basidia: in a layer on the surface of host tissue
- without paraphyses
- with probasidal swellings
- conidia: ovoid

**Proliferobasidioum** (1 species)
- Heliconiaceae
- basidia: in gelatinous pustules breaking through epidermis or stomata
- with thick-walled probasidia
- without paraphyses
- with repeated proliferation from within the probasidal wall
- conidia: globose
Fig. 1 Original line drawings by Franz Oberwinkler of selected species of Brachybasidiaceae (included with the approval by Barbara Oberwinkler).

a  *Brachybasidium pinangae* (comp. Fig. 13 in Begerow et al. 2002). Ball of basidia with probasidia on top of a stomatal opening.

b  *Exobasidiellum graminicola* (comp. Fig. 14 in Begerow et al. 2002). Basidia with thick-walled probasidia forming a layer on host tissue as well as just liberated and germinating basidiospores with conidia.

c  *Dicellomyces gloeosporus*.

d  *Kordyana tradescantiae* (comp. Fig. 15 in Begerow et al. 2002)

For localizations of drawings labeled with non-bold letters c–f see sketch at non-bold a. (a) Schematically drawn longitudinal section of a basidiocarp. (b) Gelatinous basidiocarp on host tissue. (c) Basidia with probasidial thickenings, basidiospores, and conidia. (d) Hyphae in a gelatinous matrix. (e) Parallel hyphae in the base of the basidiocarp. (f) Hyphae with swollen tips at the lower surface of the basidiocarp.
Brachybasidiaceae are illustrated by drawings elaborated by Franz Oberwinkler (Fig. 1).

The taxonomic history of the group of fungi currently classified in Brachybasidiaceae started in 1892 when Patouillard and von Lagerheim (1892) described *Exobasidium tradescantiae* Pat. (now *Kordyana tradescantiae* (Pat.) Racib.) based on a fungus infecting a species of *Tradescantia* (Commelinaceae) from Ecuador, and when Bresadola (in Krieger, Fung. Saxii. Exsicc., Pilze Sachsen’s: no. 664, 1892) presented the species *Exobasidium graminicola* Bres. (now *Exobasidiellum graminicola* (Bres.) Donk) on Poaceae (Online Resource 1). The oldest genus currently placed in Brachybasidiaceae is *Kordyana*, established by Raciborski in 1900 including *Kordyana pinangae* Racib. (now *Brachybasidium pinangae* (Racib.) Gãûm.) and *Kordyana tradescantiae* that was selected as type species by Gãûmán (1922). Gãûmán (1922) created the genus *Brachybasidium* with *B. pinangae* as type species and established the family Brachybasidiaceae in 1926 (Gãûmán 1926). Further genera currently placed in Brachybasidiaceae are *Exobasidiellum* (Donk 1931), *Dicellomyces* (Olive 1945), *Proliferobasidium* (Cunningham et al. 1976), and *Meira* (Boekhout et al. 2003), with the latter including up to now only asexually multiplying yeasts.

The genus *Dicellomyces* with the type species *D. gloeosporus* L.S. Olive on a species of Poaceae has been used to classify four species of plant-parasitic fungi forming gelatinous, pustulate basidiocarps breaking through the host tissue. *Dicellomyces bombacis* B.K. Bakshi infecting a species of Malvaceae, however, was separated from other *Dicellomyces* spp. as *Ceraceosorus bombacis* (B.K. Bakshi) B.K. Bakshi first within the Brachybasidiaceae (Cunningham et al. 1976) and later in the order Ceraceosorales based on molecular sequence data (Begerow et al. 2006; Piątek et al. 2016).

Raciborski (1900) described a further genus called *Lelum* based on *Lelum ustilaginoides* Racib. on *Persea* sp., which is attributed to Brachybasidiaceae by some authors (e.g., Kirk 2019). However, due to the host species belonging to Lauraceae, a light brown spore mass, and basidiospores with longitudinal ridges, we suppose that this species may belong to Cryptobasidiaceae. Though basidia carry two basidiospores each, this characteristic does not fit to the morphological concept of Cryptobasidiaceae. Thus, this species needs further analyses.

The authors contributing to the knowledge of Brachybasidiaceae in the past noticed similarities of the species included in Brachybasidiaceae and Exobasidiaceae, but they also noted important differences, especially the fact that species of Brachybasidiaceae develop basidia with only two sterigmata each, in contrast to basidia of typical species of Exobasidiaceae that develop mostly 4 (2–8) basidiospores. Basidiospores of *Exobasidium* spp. further differ from basidiospores of species of Brachybasidiaceae by apiculi with abaxial orientation. Forked basidia with two basidiospores that develop septa during germination are typical for species of Brachybasidiaceae and for species of Dacrymycetales (Agaricomycotina). The sterigmata of dacrymycetalean basidia, however, are much longer than those of the species of Brachybasidiaceae, and species of Dacrymycetales are saprotrophs in contrast to species of Brachybasidiaceae that are exclusively plant-parasitic during their sexual development (Oberwinkler 1982).

In the working group of Franz Oberwinkler, R. Bauer, D. Begerow, and F. Oberwinkler himself made important contributions to the systematics of Ustilaginomycotina at order and family level. On genus and species level, however, our knowledge of Brachybasidiaceae is still very incomplete, as evident by incomplete morphological descriptions, several taxonomic problems, and molecular sequences lacking for most described species in this group (comp. Begerow et al. 2018; Berndt and Sharma 1998). By the present publication, we contribute new data to this group based on recently collected specimens from Bolivia and Panama in the neotropics.

### Materials and methods

### Specimens

The specimens of Brachybasidiaceae cited in this study were collected during fieldwork for research and teaching. The new species from Panama was first noticed in the context of the Majagua project (M numbers, Piepenbring et al. 2012b). The new species collected in Bolivia was found during the “Curso de hongos” (CH, Course on Fungi), realized at the Universidad Autónoma Gabriel René Moreno in Santa Cruz, Bolivia. The specimens were analyzed, preserved, and deposited in the herbarium of the Universidad Autónoma de Chiriquí, Panama (UCH), the national herbarium of the Universidad de Panamá (PMA), in the herbarium of the Museo de Historia Natural Noël Kempff Mercado at the Universidad Autónoma Gabriel René Moreno, Bolivia (USZ), and/or in the Botanische Staatssammlung München, Germany (M).

### Identification of host plants

The host plant species of the specimen collected in Panama was identified as *Goepertia panamensis* (Rowlee) Borchts. & S. Suárez (syn. *Calathea panamensis* Rowlee, Marantaceae) based on keys and descriptions in the Flora of Panama (Woodson and Schery 1943) and herbarium specimens deposited in PMA.
The host plant of the specimen from Bolivia has been identified with the help of the checklist of plants for Bolivia (Jørgensen et al. 2014), diverse Florae (Costa Rica: Hammel et al. 2003; Panama: Woodson and Schery 1943), molecular sequence data of the ITS region and of the rbcL gene, and labeled photos available on the Internet.

In order to obtain DNA sequence data, the following primers have been used: for the ITS region, ITS5A-F (Stanford et al. 2000) and 241r-R (Michelangeli et al. 2004) and for the rbcLa marker, the primers rbcLa-F (SI-For) and rbcLa-R (SI-Rev) (Kress et al. 2009).

The ITS sequence (GenBank acc. no. MN399972) obtained from the host plant from Bolivia shows identities of 97% and more with several species of Goeppertia (Calathea). Among these species, the only one that is reported for Bolivia and morphologically similar to the specimen collected in Bolivia is Goeppertia propinqua (Poeppl. & Endl.) Borchs. & S. Suárez, that is represented by the sequence JQ341297 (published by Borchsenius et al. 2012) in GenBank and has 97% percent identity with the sequence from the plant from Bolivia. For the rbcL sequence (GenBank acc. no. MN953612), there is no sequence of Goeppertia propinqua available in GenBank for comparison.

Morphological analysis

Fresh and dried specimens of recently collected Brachybasidiaceae were investigated morphologically following standard methods as described by Judith et al. (2015). Measurements represent mean values plus/minus standard deviation and minimum and maximum values are given in parentheses. Hyphae, basidia, basidiospores, and conidia of these fungi are relatively small and thin-walled. They easily collapse when dried. Therefore, preferably fresh specimens were analyzed or sections were made by hand with new razor blades from fresh sori. These sections were preserved on glass slides below cover slips in a mounting medium for semi-permanent preparations with cotton blue.

Molecular analysis of the fungi

DNA extraction, PCR, and sequencing

Genomic DNA was isolated directly from the herbarium specimens. Isolation of fungal material, DNA extraction, amplification, purification of PCR products, and sequencing were provided by the ALVALAB, Oviedo, Spain. For some specimens, this was done with innuPREP Plant DNA Kit (Analytikjena, Jena, Germany) and sequences were obtained from Microsynth Seqlab (Göttingen, Germany). The ITS1-5.8S-ITS2 region of the ribosomal DNA (ITS) was amplified using the primer pair ITS1-F (Gardes and Bruns 1993) and ITS4 (White et al. 1990). The 5′-end of the nuclear large subunit ribosomal DNA (LSU) was amplified using the primer pair LR0R and LR5 (White et al. 1990; O’Donnel 1992, 1993). The DNA sequences determined for this study were deposited in GenBank (GenBank accession numbers are given in Fig. 2 and the taxonomy section).

Phylogenetic analyses

To elucidate the phylogenetic position of the specimens from Bolivia and Panama, their sequences were analyzed within a concatenated ITS + LSU dataset. Since preliminary analyses and the comparison of the sequences to available data in GenBank using BLAST (Altschul et al. 1997) revealed an affinity of the specimens to the Exobasidiales, the dataset was reduced to members of the Exobasidiales and Rhamphospora nymphaeae (Doassansiales) as outgroup. The dataset covered all Exobasidiales genera of which sequences were available in GenBank. If present in GenBank, sequences of the respective type species were used. In addition, sequences of all available species of Brachybasidiaceae and Graphioliaceae were added including sequences from the public catalog of the NITE Biological Resource Center collection (NBRC), Japan. GenBank accession numbers of the sequences used (Beigerow et al. 2002; Boekhout et al. 2003; Cao et al. 2018; Crous et al. 2003; Kottke et al. 2010; Kruse et al. 2017; Macedo et al. 2016; Maier et al. 2006; Matheny et al. 2006; Nasr et al. 2019; Piepenbring et al. 2010, 2012a; Schoch et al. 2014; Sepúlveda et al. 2017; Tanaka et al. 2008; Wang et al. 2015; Yasuda et al. 2005) are cited in Fig. 2.

Sequence alignment was obtained independently for the ITS and LSU part of the ITS + LSU dataset using MAFFT 7.313 (Katoh and Standley 2013) with the L-INS-i option. To obtain reproducible results, manipulation of the alignments by hand as well as manual exclusion of ambiguous sites were avoided as suggested by Gatesy et al. (1993) and Giribet and Wheeler (1999). Instead, highly divergent portions of the alignments were omitted using GBlocks 0.91b (Castresana 2000) with the following options for the ITS part of the ITS + LSU dataset: “Minimum Number of Sequences for a Conserved Position”: 9, “Minimum Number of Sequences for a Flank Position”: 9, “Maximum Number of Contiguous Non-conserved Positions”: 8, “Minimum Length of a Block”: 5, and “Allowed Gap Positions” to ‘With half.’ For the LSU part of the ITS + LSU dataset, these parameters were set to 14/14/11/5/“With half,” respectively. Afterwards, the ITS and LSU sequence alignments were concatenated.

The resulting alignment (new number of positions, 1099 (23% of the original 4783 positions); number of variable sites, 543) was used for phylogenetic analyses applying Bayesian inference (BI) and maximum likelihood (ML). For the BI analysis, a Markov chain Monte Carlo technique was used as implemented in MrBayes 3.2.6 (Ronquist et al. 2012). Two runs over 5,000,000 generations, each consisting of four
chains, were implemented using the general time-reversible model of DNA substitution with gamma-distributed substitution rates and an estimated proportion of invariant sites, random starting trees, and default starting parameters of the DNA substitution model. A 50% majority-rule consensus tree was computed from 75,000 trees that were sampled after the process had become stationary. The topology was rooted with *Rhamphospora nymphaeae* (Doassansiales). Numbers on branches before slashes are estimates for a posteriori probabilities; numbers on branches after slashes are ML bootstrap support values. Branch lengths were averaged over the sampled trees. They are scaled in terms of expected numbers of nucleotide substitutions per site. The applied taxonomical concepts follow Begerow et al. (2014). Species belonging to the new genus are indicated in bold.

25% of trees from each run were discarded (burn-in). The remaining trees were used to compute a 50% majority-rule consensus tree to obtain estimates for the a posteriori probabilities of groups of species. This Bayesian approach to phylogenetic analysis was repeated five times to test the independence of the results from topological priors (Huelsenbeck et al. 2002). ML analysis (Felsenstein 1981) was conducted.
with RAxML 8.2.11 (Stamatakis 2014) invoking the GTRGAMMA and the rapid bootstrap option (Stamatakis et al. 2008) with 1000 replicates, but omitting an estimated proportion of invariant sites following the advice given in the user manual (Stamatakis 2016).

Results

Phylogenetic analyses

For the two specimens of Brachybasidiaceae on Goeppertia panamensis from two localities in Panama, the ITS and LSU sequences were identical, while the sequences obtained from the specimen collected in Bolivia on Goeppertia propinqua differed in 36 bp/6.4% for the ITS region and 32 bp/2.6% for the LSU.

The different runs of the BI and the ML analysis yielded consistent topologies. To illustrate the results, the consensus tree of one run of the BI is presented (Fig. 2). Using Rhamphospora nymphaeae as outgroup, the clades in the phylogenetic tree were congruent to the families as presented by Begerow et al. (2014). In all analyses, the specimens from Bolivia and Panama formed a strongly supported monophylum in a well-supported group including all sequences from Dicellomyces, Kordyana, and Meira specimens. However, relations between these taxa were not resolved. Species of Kordyana formed a monophylum together with Dicellomyces gloeosporus and Dicellomyces as well as Meira were revealed to be polyphyletic.

Taxonomy

Marantokordyana M. Piepenbr., Maike Hartmann, T.A. Hofm. & M. Lutz, gen. nov. (MycoBank number MB 832383)

Type species: Marantokordyana oberwinkleri M. Piepenbr., Maike Hartmann, T.A. Hofm. & M. Lutz, see below

Etymology: The name refers to morphologically similar species of Kordyana and to the host family Marantaceae.

Description:

Plant-parasitic fungi infecting species of Marantaceae. Fungal hyphae in intercellular spaces in leaves, causing yellowish spots without hypertrophic growth, filling substomatal chambers, protruding through stomatal openings. Basidia exposed on top of stomata on the abaxial side of leaf blades in spherical balls, mixed with paraphyses, each basidium with two straight sterigmata carrying one basidiospore each. Basidiospores blastosporic, with conspicuous hilum, without apiculus, cylindrical to slightly allantoid, often liberated in pairs, with one central septum at maturity. Basidiospores germinating with thin hyphae producing tiny, rod-shaped to

Fig. 3 Species of Marantokordyana. a–c M. oberwinkleri on Goeppertia panamensis (MP 5127). a Infected plant in the field. b The upper side and the lower side of leaf blades with leaf spots. c A leaf spot seen from the lower side of the blade by a stereomicroscope. Each white dot is a ball of basidia. Scale bar = 2 mm. d–f M. boliviana on Goeppertia propinqua (CH 11). d Infected plant in the field. e The upper side of a leaf blade with leaf spots. f Leaf spots seen from the lower side of the blade. White dots are balls of basidia
fusiform conidia that multiply by budding yeast cells.

**Marantokordyana oberwinkleriana** M. Piepenbr., Maike Hartmann, T.A. Hofm. & M. Lutz, sp. nov. (MycoBank number MB 832384) (Figs. 3a–c, 4, and 5)

Type on *Goeppertia panamensis* (Rowlee) Borchs. & S. Suárez (syn. *Calathea panamensis* Rowlee; Marantaceae). Panama, Chiriquí province, Dolega district, Los Algarrobos, trail to Río Majagua, 8° 29′ 22″ N, 82° 26′ 1″ W, 110 m asl., 10. 8. 2018, leg. M. Piepenbring and M.U. Schmidt MP 5412 (holotype PMA 0123802, isotypes M 141363, UCH 11709).

GenBank: ITS = MN275897, LSU = MN275900.

**Etymology:** This species is named in honor of Franz Oberwinkler (1939–2018) who made important contributions to the knowledge of heterobasidiomycetes (comp. Piepenbring et al. 2019).

**Description:**

Leaf blades with scattered to numerous spots due to infection. **Infected areas** elongated rectangular and rounded at the ends, laterally delimited by veins of the leaf blade, not swollen, in surface view mostly (3–)4–10(–11) × (1.5–)2–2.5(–3) mm (n = 10), sometimes larger by fusion, adaxially yellow to slightly orange colored, with white balls of basidia evident with a hand lens or a stereomicroscope. All the substomatal chambers of an infected area filled by fungal hyphae in dense stromata, hyphae protruding through the stomatal openings, developing balls of basidia mixed with paraphyses, one ball on the top of each stoma, balls of basidia easily breaking off and being dispersed.

**Substomatal chambers** filled with fungal stroma formed by dense fungal hyphae, more voluminous than substomatal chambers without fungal stroma (Fig. 4), cellular details of fungal stroma difficult to distinguish, in very thin sections three types of fungal hyphae distinguishable, very thin (approx. 1 μm wide) hyphae close to host cells, thin (approx. 1.5 μm wide) hyphae mixed with the thick hyphae, and thick (approx. 3 μm wide) hyphae with swellings of up to 5 μm width at the base.

**Balls of basidia** spherical to globose, composed of basidia in different developmental stages and 1.5 μm wide paraphyses densely packed in the center and loosely exposed at the surface of a ball, (40–)55–80(–90) μm diam. (n = 20), not pigmented.

**Basidia** holobasidia, with basidial cell cylindrical, of variable length of up to at least 70 μm, sometimes shorter by retraction septa formed during sporulation, 3–4(–4.5) μm wide (n = 10), thinner after the liberation of the basidiospores, with two apical, straight or bent, elongated and often slightly swollen sterigmata of (2–)3.5–6.5(–7.5) × 1.5–2 μm (n = 20), with one basidiospore developing at the tip of each sterigma. After liberation of the basidiospores basidial cells empty, sometimes with scattered tiny remnants of cytoplasm, with scars evident at the tips of the sterigmata.

**Basidiospores** liberated singly or in pairs, at the moment of liberation one-celled but soon afterwards two-celled due to a central septum, cylindrical or slightly allantoid and slightly fusoid at the base, with a conspicuous hilum (scar) at the base of each basidiospore, (10–)11–16(–19) × 3–4.5(–6) μm (n =
30), hyaline, smooth, densely filled with oil drops or with homogeneous cytoplasm. The two basidiospores of one basidium conjugating by fusing or individual basidiospores germinating with about 1 μm wide hyphae originating at their tips or laterally and developing conidia on more or less evident sterigma-like outgrowths or cells of basidiospores directly developing conidia usually at their tips.

Conidia mostly one-celled, rarely with septum, rod-shaped to fusiform or slightly allantoid, with scar at the point of detachment, 3–8(−11) × (0.5−)1(−1.5) μm (n = 20), hyaline, smooth, germinating with thin hyphae or budding forming further conidia (yeast cells).

**Further specimens examined:** On *Goepertia panamensis*. Panama, Chiriqui province, locality of the type, 12. 8. 2009, leg. M. Piepenbring and T.A. Hofmann M 409 (M 141366, UCH 11710). Same locality, 31. 8. 2009, leg. M. Piepenbring and T.A. Hofmann M 504 (M 141365, UCH 11711). Almost the same locality, 8° 29′ 18″ N, 82° 26′ 2″ W, 107 m asl., 27. 7. 2012, leg. M. Piepenbring and A. Krohn MP 5127 (M 141364, PMA
Marantokordyana boliviana M. Piepenbr., Maike Hartmann, M. Lutz & T.A. Hofm., sp. nov. (MycoBank number MB 832385) (Figs. 3d–f and 6)

**Type** on Goeppertia propinqua (Poepp. & Endl.) Borchs. & S. Suárez (syn. Calathea propinqua (Poepp. & Endl.) Körn.; Marantaceae). Bolivia, Department of Santa Cruz, Andrés Ibañez Province, Municipio Porongo, Terebinto, Hacienda Privada Arubai, 17° 41′ 10″ S, 63° 25′ 15″ E, 448 m asl., 13. 2. 2019, M. Piepenbring and participants of the Curso de Hongos CH 88 (holotype USZ). GenBank: ITS = MN275898, LSU = MN275901.

**Etymology:** The type of this species was discovered in Bolivia.

**Description:**
Leaf blades with scattered to sometimes very numerous spots due to infection. Infected areas elongated rectangular and often poorly delimited, on the abaxial side laterally delimited by veins of the leaf blade, not swollen, on the abaxial side intercostal areas with balls of basidia mostly (6–)7–

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Fig. 6 Marantokordyana boliviana (CH 88). a Fungal cells isolated from a substomatal chamber, without adjacent host cells. Thick hyphae with probasidial swellings directed towards the stomatal opening (not shown) and thin, indistinguishable hyphae filling the space in between. b Basidia after the liberation of basidiospores. c Liberated basidiospores in pairs, one pair with a conjugation bridge at the base. d Conjugated basidiospores germinating with hyphae. Note the retraction septum in the cell on the left hand side. e One half of a basidiospore forming a conidium at the tip of a hypha. Note the retraction septum. f Yeast-like conidia budding or forming conidia on sterigma-like outgrowths. g An intercalary cell of a hypha with an outgrowth developing a conidium. h A hypha developing conidia. Septa of the hypha are difficult to distinguish. Scale bars = 10 μm
14(−15) × 1.5−2 mm (n = 10), probably older infected areas much larger by fusion of leaf spots, adaxially yellowish, abaxially whitish to slightly yellow to orange colored areas of sporulation surrounded by whitish host tissue, with white balls of basidia evident with a hand lens or a stereomicroscope. All the substomatal chambers of an infected area filled by fungal hyphae in dense stroma, hyphae protruding through the stomatal openings, developing balls of basidia mixed with paraphyses, one ball on the top of each stoma, balls of basidia easily breaking off and being dispersed.

**Substomatal chambers** filled with fungal stroma formed by dense fungal hyphae, cellular details of fungal stroma difficult to distinguish, in the stroma isolated from a substomatal chamber two types of fungal hyphae distinguishable, thick hyphae that penetrate through the stoma and develop basidia outside the host tissue and thin hyphae that fill the space between the thick hyphae. Thick hyphae mostly about 4 μm wide, cylindrical with more or less undulated walls or thinner and sometimes with a basal, probably probasidial swelling of 4–6 μm width at their base. Thin hyphae about 1 μm wide, curled, ramified, very difficult to distinguish by light microscopy.

**Balls of basidia** globose or of irregular shape, composed of basidia of different ages and approx. 1 μm wide paraphyses densely packed in the center and loosely exposed at the surface of a ball, (60−)70−90(−100) μm diam. (n = 10), not pigmented. In young balls of basidia, only thick hyphae are present protruding through the stomatal openings, larger (older) balls contain thick hyphae forming basidia and paraphyses that are probably the prolongations of the thin hyphae filling the substomatal chamber.

**Basidia** holobasidia, cylindrical, delimited at their base by a retraction septum, i.e., the subbasidial cell is empty, without plasm, and collapses, basidia (10−)11−18(−20) × 3−4 μm (n = 10) when sterig mata are present (not including the sterigmata), sometimes shorter by retraction septa formed during sporulation, each basidium with two apical, straight, or slightly bent sterigmata, (1.5−)2−4 μm (n = 10) long, with one basidiospore developing at the tip of each sterigma. After liberation of the basidiospores, basidal cells with scattered tiny remnants of cytoplasm, with scars evident at the tips of the sterigmata.

**Basidiospores** liberated single or in pairs, two-celled due to a central septum, cylindrical or slightly allantoid and slightly fusoid at the base, with a conspicuous hilum (scar) at the base of each basidiospore, (11−)12−16(−18) × 3−4.5(−5) μm (n = 30), hyaline, smooth. The two basidiospores of one basidium conjugating by fusing at their bases or individual basidiospores germinating with about 1 μm wide hyphae and developing conidia on more or less evident sterigma-like outgrowths or directly developing conidia usually at their tips. Hyphae approx. 1 μm wide, septa very difficult to distinguish by light microscopy.

**Conidia** mostly one-celled, rarely with septum, rod-shaped, sometimes slightly allantoid, or filiform, with more or less evident scar at the point of detachment, (2−)3−9(−15) × (0.5−)1(−1.5) μm (n = 20), hyaline, smooth, germinating with thin hyphae or budding forming further conidia (yeast cells).

**Discussion**

**Morphological structures of species of Brachybasidiae**

The presence of balls of basidia formed on the top of stomatal openings by cells originating from fungal stroma in substomatal chambers and protruding through the stomatal openings is a structural feature present in distantly related species of Microstromatales (e.g., Volvocisporium; Begerow et al. 2001; Oberwinkler 2012b; Ritschel et al. 2008), Exobasidiaceae (e.g., Exobasidium oxyiocci, illustrated by F. Oberwinkler in Begerow et al. 2001), and Brachybasidiaceae (Brachybasidium, Kordyana, Marantokordyana). Such structures apparently are the result of convergences: If fungal cells are not able to break through the epidermis, they use stomatal openings (Gäumann 1922). Similar strategies for the liberation of spores (not basidiospores) exposed by hyphae protruding through stomatal openings can also be observed for cercosporoid fungi (Mycosphaerellales, Ascomycota), Entylomella spp. (Entylomatales), and Peronosporales (Oomycota) (Pepenbring 2015). Repeatedly, swellings were documented for thick hyphae in the stroma filling substomatal chambers. They were observed in very thin sections but even in very thin sections, they could not always be distinguished because cells are densely packed and often gelatinized (comp. Gäumann 1922; Berndt and Sharma 1998). In addition, they might be ephemeral because their function may be related to the fact that the fungus needs to force its cells through the stomatal opening or to the development of basidia that need high inner pressure (turgor) for their development. High inner pressure in the substomatal chambers is evident by the fact that the substomatal chambers filled with stroma are swollen (Fig. 4). The swellings may be the place of karyogamy of dikarya and therefore interpreted as probasidia. Gäumann (1922) observed karyogamy and following meiosis in probasidial cells of Brachybasidium pinangae. More or less conspicuous probasidial swellings without observations concerning the nuclei were illustrated repeatedly by F. Oberwinkler (Fig. 1d) and some further authors for Kordyana tradescantiacea, Microstroma juglandis (Begerow et al. 2001), Muribasidiospora hesperidium (Begerow et al. 2001), Volvocisporium grewiae (Ritschel et al. 2008), and Volvocisporium triumfetticola (Begerow et al. 2001). Apparently, it is a widely spread structure that is easily overlooked and therefore not useful to distinguish taxa within Brachybasidiaceae. While thick hyphae forming part of the
stroma filling substomatal chambers are considered basal parts of basidia, thin hyphae of the stroma are supposed to protrude through the stomatal opening and to form paraphyses.

The question whether basidiospores are actively discharged, e.g., ballistosporic, or blastosporic was not addressed by the first authors like Gäumann (1922). Maybe since Cunningham et al. (1976) described “discharged” basidiospores, authors described basidia of Brachybasidiaceae as ballistosporic without critical reflection. Based on own observations and critical interpretation of information in literature, however, basidiospores of Brachybasidiaceae cannot be ballistosporic: (1) The sterigmata of the basidia are straight or stout (e.g., Reid 1976) and not curved as in the case of ballistosporic basidia. (2) According to drawings available in literature and own observations, basidiospores of species of Brachybasidiaceae have broad hila at their basis but no apiculi. There is confusion in the application of the terms “apiculus” and “hylum” (e.g., Reid 1976). The hilum is the area of the attachment of the basidiospore to the basidium, evident as a scar at the basis of the detached basidiospore, while the apiculus is a projection close to the hilum where the Buller’s drop develops and causes active liberation of the basidiospore (Piepenbring 2015). (3) Basidiospores of species of Kordyana and Marantokordyana are often found attached to balls of basidia and in pairs. The diaspores responsible for dispersal of these species are the balls of basidia, basidiospores, and conidia.

As typical for species of Basidiomycota classified in heterobasidiomycetes (comp. Piepenbring 2015), the result of the germination of basidiospores is variable. In the case of species of Brachybasidiaceae, basidiospores can fuse by a conjugation bridge, directly form conidia, or germinate forming thin hyphae that produce conidia. Conidia can be cylindrical, rod-shaped, allantoid, fusiform, or globose. The conidia germinate forming hyphae or they produce further spores, often on short outgrowths. Budding conidia can be considered yeast cells. The shape of conidia may be an important characteristic to distinguish systematic groups and species of Exobasidiales (comp. “Dicellomyces” scirpi).

New genus and species of Marantokordyana

The specimens of Brachybasidiaceae recently collected in Bolivia and Panama are morphologically close to Kordyana spp. by basidia in suprastomatal balls, basidia forming two basidiospores each, and basidiospores germinating after septation with hyphae forming elongate conidia (yeast cells). We classify them in a separate genus new to science because in contrast to species of Kordyana that mostly infect species of Commelinaceae (Commelinales, Commelinidae), the two new species infect species of Marantaceae (Zingiberales, Commelinidae) and because the two new species form a well-supported monophylum in the molecular phylogenetic analyses. Within Brachybasidiaceae, no species infecting Marantaceae are known up to now. In the phylogenetic tree, the monophylum of the new genus forms part of a polytomy including all the species of Brachybasidiaceae represented by molecular sequence data. Within the family, no specific sister relationship can be distinguished for the members of the new genus. Further described genera differ morphologically by fungal stromata breaking through host tissue, absence of paraphyses, proliferating basidia, and/or different shapes of conidia (comp. Table 1). We consider host specificity on family level as an important argument for the definition of systematic groups, because host specificity has been shown to be indicative of systematic relationships for several systematic groups of Exobasidiales, e.g., Graphiola spp. exclusively on Areaceae and Exobasidium spp. on Ericaceae and closely related families.

Both species proposed as new species in Marantokordyana infect species of the same genus (Goeppertia, syn. Calathea) and are morphologically very close to each other. The only evident morphological differences are the shorter size of basidia and straighter and shorter sterigmata in M. boliviana with a length of 2–4 μm versus 3.5–6.5 μm in M. oberwinkleri. Further, but probably less reliable differences are the yellowish to orange color of the lower side of the leaf spots in M. boliviana versus white color in M. oberwinkleri, and slightly larger balls of basidia in M. boliviana. The distinction of the two species is confirmed by differences in sequence data of the ITS and LSU regions of rDNA.

Taxonomy of Kordyana spp.

During the compilation of species known for Brachybasidiaceae (Online Resource 1), several taxonomic problems were noticed for species of Kordyana: Kordyana indica lacks a diagnosis. Kordyana polliae var. microspora lacks a diagnosis and is considered a synonym of Kordyana polliae by Index Fungorum, but according to the description it may be a species distinct from K. polliae. The type species of Kordyana (K. tracyscantia) and most other species described in Kordyana infect species of Commelinaceae, so Kordyana boswelliae on Burseraceae, Kordyana cyphelloidis described from Bignoniaceae, and Kordyana polliae on Poaceae are doubtful. In addition to the different host family, K. cyphelloidis differs from typical Kordyana spp. by basidiocarps on cancers on leaf blades, leaf petioles, or branches, and by basidia developing from regularly septate hyphae in a hymenium surrounded by a peridium formed by setae with ornamented surface (Viégas 1945). As for many species of Kordyana, morphological characterizations are incomplete and because most of the species of Kordyana lack
DNA sequence data, it is not possible to clarify these taxonomic problems without new specimens and DNA sequence data.

“Dicellomyces” scirpi

The species Dicellomyces scirpi is presently classified in the genus Dicellomyces because of morphological similarities with D. gloeosporus, the type species of Dicellomyces, namely by being a plant-parasitic fungus on a grass developing gelatinous stromata breaking through host tissue and by developing bisterigate basidia with probasidial swellings in a hymenium. The two species differ by their host family (Dicellomyces gloeosporus: Poaceae, D. scirpi: Cyperaceae), the shape of the sori (discoid versus elongated), the presence of paraphyses (absent versus present, described as “hyphidia” by Parmasto 1968), more persistent and evident probasidial swellings in D. gloeosporus, and the shape of conidia formed by germinating basidiospores (globose versus allantoid, also described as “crescentiform”) (Ingold 1985; Olive 1945; Parmasto 1968; Reid 1976).

Analyses of LSU sequence data including D. gloeosporus and D. scirpi resulted in conflicting hypotheses considering the congeneric classification. While Tanaka et al. (2008) revealed monophyly of both Dicellomyces species (support: 72/90) being sister taxon of Meira spp. (support: 73/53), Nasr et al. (2019) placed the two Dicellomyces species in different clades with Kordyana spp. as closest relatives of D. gloeosporus (support: 100/99) and Meira spp. as closest relative of D. scirpi (support: 92/62). The results of Nasr et al. (2019) are confirmed by the analyses presented here based on extended data (ITS + LSU) and additional species included (Kordyana brasiliensis, Meira miltonrushii, M. nicotianae). Meira is revealed polyphyletic with Dicellomyces scirpi as closest relative of M. nicotianae.

Dicellomyces scirpi most probably does not belong to the genus Dicellomyces, however, we refrain from establishing a new genus for D. scirpi, because several other generic concepts are available that are not represented by molecular sequences and therefore not included in the molecular analyses. In order to demonstrate this conclusion, we write the genus name of “D.” scirpi between quotation marks.

Conclusion

In the present publication, we draw some taxonomic conclusions for species of Brachybasidiaceae. Many species concepts in this relationship, however, are still incompletely known, because they were only presented by short, incomplete species descriptions in literature. For 12 of now 24 known species of Brachybasidiaceae (Online Resource 1), LSU and for some of them also ITS sequence data are available, only four of seven described genera are included in the phylogenetic analyses. Although species in this group are inconspicuous, without economic importance, and rare, they should be collected, well preserved, and analyzed in order to improve our understanding of this interesting group of fungal plant parasites.

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