

# **Classes and phyla of the kingdom** *Fungi*

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# **Abstract**

Fungi are one of the most diverse groups of organisms with an estimated number of species in the range of 2–3 million. The higher-level ranking of fungi has been discussed in the framework of molecular phylogenetics since Hibbett et al., and the defnition and the higher ranks (e.g., phyla) of the 'true fungi' have been revised in several subsequent publications. Rapid accumulation of novel genomic data and the advancements in phylogenetics now facilitate a robust and precise foundation for the higher-level classifcation within the kingdom. This study provides an updated classifcation of the kingdom *Fungi*, drawing upon a comprehensive phylogenomic analysis of *Holomycota*, with which we outline well-supported nodes of the fungal tree and explore more contentious groupings. We accept 19 phyla of *Fungi,* viz*. Aphelidiomycota*, *Ascomycota*, *Basidiobolomycota*, *Basidiomycota*, *Blastocladiomycota*, *Calcarisporiellomycota*, *Chytridiomycota*, *Entomophthoromycota*, *Entorrhizomycota*, *Glomeromycota*, *Kickxellomycota*, *Monoblepharomycota*, *Mortierellomycota*, *Mucoromycota*, *Neocallimastigomycota*, *Olpidiomycota*, *Rozellomycota*, *Sanchytriomycota,* and *Zoopagomycota*. In the phylogenies, *Caulochytriomycota* resides in *Chytridiomycota*; thus, the former is regarded as a synonym of the latter, while *Caulochytriomycetes* is viewed as a class in *Chytridiomycota*. We provide a description of each phylum followed by its classes. A new subphylum, *Sanchytriomycotina* Karpov is introduced as the only subphylum in *Sanchytriomycota*. The subclass *Pneumocystomycetidae*

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Kirk et al. in *Pneumocystomycetes*, *Ascomycota* is invalid and thus validated. Placements of fossil fungi in phyla and classes are also discussed, providing examples.

**Keywords** *Caulochytriomycota* · Phylogenomic analyses · *Pneumocystomycetidae* · *Sanchytriomycotina*

# **Introduction**

The Kingdom *Fungi* is one of the most diverse groups of living organisms on earth and its members occur across a broad range of ecosystems including extreme environments. Fungi are primarily heterotrophs that occur as uni- to multicellular organisms and are vital in ecosystem functioning (Kendrick [2000;](#page-142-0) Kirk et al. [2008;](#page-142-1) Hyde et al. [2018](#page-140-0)). Over time, understanding diferent aspects of fungi (i.e. taxonomy, diversity, species number, genetics, chemistry) has improved rapidly by incorporating molecular and bioinformatics tools with traditional approaches (Wijayawardene et al. [2021a](#page-160-3)). Many studies have been published based on these polyphasic approaches and these have expanded the magnitude of the kingdom (Hyde [2022](#page-140-1)). Several studies have predicted a higher diversity of diferent taxonomic groups (e.g. freshwater fungi *fde* Calabon et al. [2022](#page-132-0); telemorphic *Ascomycota fde* Senanayake et al. 2022; *Basidiomycota fde* He et al. [2022a](#page-139-2); asexually producing taxa *fde* Wijayawardene et al. [2022b](#page-160-4)). Currently, about 160,000 species of fungi are accepted by Species Fungorum (2024; accession data 12.04.2024) though the estimation of the number of species is 2–3 million (Niskanen et al. [2023\)](#page-148-0). Revealing these unknown or 'missing taxa' is one of the biggest challenges, since many species do not produce any (easily) observable phenotypic characters, i.e. dark taxa (Page [2016](#page-151-0); Wang et al. [2016a,](#page-159-0) [b](#page-159-1)), and it is thus impossible to identify and describe them from conventional techniques (Mitchell and Zuccaro [2006](#page-148-1); Seifert [2009](#page-154-1)). Over three decades, i.e. since White et al. ([1990](#page-159-2)), Sanger sequencing has played a vital role in species recognition, determining the higher-level classifcation and other fundamental taxonomic approaches such as linking sexual and asexual or dimorphic taxa. Nevertheless, Sanger sequencing can only sequence specimens or cultures individually (Shokralla et al. [2012\)](#page-155-0). Margulies et al. ([2005\)](#page-146-0) and Shendure et al. ([2005\)](#page-154-2) developed 'sequencing-by-synthesis technology' to overcome the major drawbacks of Sanger sequencing, which is currently called 'high-throughput sequencing' (HTS) or 'next-generation sequencing'(NGS) (Schuster [2008](#page-154-3); Shendure and Ji [2008](#page-154-4)). At present, HTS/NGS plays an important role in microbial and mycological research, especially in determining dark taxa from environmental samples which could represent novel lineages in the Kingdom *Fungi* (Tedersoo et al. [2018\)](#page-157-0).

# **Higher‑ranking classifcation of fungi and current status**

Higher-level classifcation of diferent organisms has been debated among taxonomists in diferent disciplines. Ruggiero et al. [\(2015\)](#page-152-0) mentioned that 'biological classifcation (taxonomy) aims to simplify and order the immense diversity of life into coherent units called taxa that have widely accepted names and whose members share important properties.' The Linnaean system has played a signifcant role for over 250 years (since mid-eighteenth century) and is broadly adopted by taxonomists (Schuh [2003;](#page-154-0) Ruggiero et al. [2015](#page-152-0)). The molecular techniques introduced in the last three decades have progressed the Linnaean system to 'a system of modern biological classifcation' (Liu et al. [2016\)](#page-145-0).

Nowadays, molecular systematics has become the basis for current classifcation schemes of fungi at various ranks. Hibbett and Donoghue ([1998\)](#page-139-0) discussed the importance of using DNA-based phylogenetic analyses to classify fungi. Hibbett et al. [\(2007\)](#page-139-1) was one of the first significant studies that broadly and comprehensively discussed the higherlevel ranking of fungi. Subsequently, diferent studies have addressed the higher-level ranking of fungi but are restricted to diferent taxa (Table [1\)](#page-2-0).

A milestone in recent fungal classifcation was the study of Tedersoo et al. ([2018\)](#page-157-0) based on both phylogenies and divergence time estimates. In their analyses, they accepted 18 phyla based on their DNA sequence analyses, though the taxa in *Caulochytriomycota* lack DNA sequences. Galindo et al. ([2021](#page-137-0)) introduced a new phylum, *Sanchytriomycota*, in *Holomycota*. Wijayawardene et al. [\(2020a,](#page-160-0) [2022a](#page-160-1)) accepted 20 phyla in the kingdom *Fungi* of which, 17 have been proposed to accommodate early divergent taxa, i.e., excluding *Ascomycota*, *Basidiomycota,* and *Entorrhizomycota* in *Dikarya* (Table [2](#page-3-0))*.* However, recent subsequent studies provided diferent conclusions on the classifcation of Early Divergent Fungi (EDF) (see below). Wijayawardene et al. ([2020b\)](#page-160-2) recommended the importance of maintaining a dynamic database on fungal classifcation.

## **Early divergent fungi (EDF)**

The discussion of 'what delineates true fungi from fungilike protists' has been a controversial topic for decades. The discovery and introduction of novel early lineages of *Holomycota* and the advancements in our knowledge of

Targeted taxon (Class/Phyla/Kingdom)*	Loci	References
Kingdom Fungi	rpb1 and rpb2	Tedersoo et al. $(2018)$
Kingdom Fungi	LSU and SSU	Voigt et al. $(2021)$
Kingdom Fungi	LSU, SSU, rpb1 and rpb2	Wijayawardene et al. (2020a)
Kingdom Fungi	Genome	Li et al. $(2021a, b)$
Kingdom Fungi	Genome	Strassert and Monaghan (2022a, b)
Subkingdom Mucoromyceta	Genome	Zhao et al. $(2023)$
Phylum Basidiomycota	nrLSU, nrSSU, 5.8S, tef1- $\alpha$ , rpb1 and rpb2	Zhao et al. $(2017)$
	Phylogenomic	Mishra et al. $(2018)$
Phylum Glomeromycota	SSU, ITS, and partial LSU	Oehl et al. $(2011a,b)$
Phylum Mucoromycota	<b>ITS, LSU</b>	Zhao et al. $(2023)$
'Zygomycota'	Conserved orthologous proteins	Spatafora et al. $(2016)$
Subphylum Pucciniomycotina	LSU, SSU, tef1- $\alpha$ , rpb1, rpb2, and 5.8 s	He et al. $(2019)$
Subphylum Ustilaginomycotina	LSU, SSU, 5.8 s, tef1- $\alpha$ , rpb1, and rpb2,	He et al. $(2019)$
Class Dothideomycetes	LSU, SSU, tef1- $\alpha$ and rpb2	Hongsanan et al. $(2020a, b)$
Class Leotiomycetes	Genome	Johnston et al. $(2019)$
Class Sordariomycetes	LSU, SSU, tef1- $\alpha$ and rpb2	Hongsanan et al. $(2017)$ and Hyde et al. $(2020)$

<span id="page-2-0"></span>**Table 1** Examples of major studies that have addressed the higher-level classifcation of fungi

\* We only list studies published after Hibbett et al. ([2007\)](#page-139-1) that focus on classes and higher taxa (including intermediate/ auxiliary ranks) as this paper focuses only on them

zoosporic fungi have blurred the lines between fungi and their relatives. However, the molecular-based phylogenetic analyses have resolved many of the deepest nodes of the fungal phylogeny, and now provide a solid foundation to defne the 'true fungi'. The early-diverging lineages of *Holomycota* are represented by phagotrophic nucleariids (*Cristidiscoidea*), rozellids+*Microsporidia* (collectively referred to as *Cryptomycota* or *Rozellomycota*), and aphelids, which branch off in succession from the ancestral lineage leading to the traditional fungal phyla, i.e. the 'true fungi' (Tedersoo et al. [2018](#page-157-0); Wijayawardene et al. [2020a;](#page-160-0) Galindo et al. [2021](#page-137-0); Mikhailov et al. [2022\)](#page-147-0). The aphelids and rozellids comprise parasitic intracellular organisms, which infect the host (algae for *Aphelida* or a diverse range of hosts for *Rozellomycota*) and phagocytize the cell contents before producing zoospores. Many mycologists include rozellids+*Microsporidia* and *Aphelida* inside *Fungi*, despite their diferences in the mode of nutrition (e.g. Tedersoo et al. [2018\)](#page-157-0). The phagotrophic mode of nutrition is absent in true fungi, and the establishment of exclusively osmotrophic nutrition has coincided with the losses of some phagotrophy-related proteins early in their evolution (Mikhailov et al. [2022](#page-147-0)). At the same time, phagotrophic holomycotan lineages lack some of the saprotrophy-linked proteins found in the true fungi (Pozdnyakov et al. [2023\)](#page-150-0). Whether this should be viewed as a basis for separating *Fungi* from the lineages of rozellids+*Microsporidia* and aphelids is a matter of debate. From a taxonomic perspective, expanding *Fungi* and dividing them into

*Eumycota* and other early divergent lineages is acceptable, but it requires a clear characterization of the *Fungi*, which is presently not available. In addition, there is a temptation to expand *Fungi* to *Holomycota*, which was already suggested by Berbee et al. ([2017](#page-132-1)). In the present review, we accept the broader version of *Fungi* by Tedersoo et al. [\(2018](#page-157-0)), but we have to keep in mind an alternative view and try to distinguish the true *Fungi* from their close relatives on the base of morpho-biological and genomic characters.

Recent phylogenomic analyses (Galindo et al. [2021](#page-137-0); Mikhailov et al. [2022;](#page-147-0) Strassert and Monaghan [2022a,](#page-156-0) [b](#page-156-1)) have shown that *Chytridiomycota* formed the earliest branch of 'true fungi'—a position contested previously by *Blastocladiomycota* in some studies (Chang et al. [2015;](#page-133-0) Li et al. [2021b](#page-145-1)). In the newer phylogenies *Blastocladiomycota* formed the next branch of 'true fungi' together with the novel phylum *Sanchytriomycota*. The relatively derived position of *Blastocladiomycota* corresponds to their advanced characters, such as sporic meiosis, hyphal-like apical growing structures (*Allomyces*), narrow sporangia exit tubes (*Catenaria*), and relatively small numbers of carbohydrate metabolism genes (Galindo et al. [2021](#page-137-0)). The *Chytridiomycota*, *Blastocladiomycota*, and *Sanchytriomycota* comprise the zoosporic lineages of fungi along with *Olpidiomycota*, which is the latest lineage to diverge from the fungal stem before the transition of fungi to terrestrial habitats (Chang et al. [2021](#page-133-1)).

Subkingdoms	Phyla	Subphyla	Classes
Aphelidiomyceta	Aphelidiomycota	Aphelidiomycotina	Aphelidiomycetes
Basidiobolomyceta	Basidiobolomycota	Basidiobolomycotina	Basidiobolomycetes
Blastocladiomyceta	Blastocladiomycota	Blastocladiomycotina	Blastocladiomycetes
			Physodermatomycetes
	Sanchytriomycota	Sanchytriomycotina subphylum nov	Sanchytriomycetes
Chytridiomyceta	Chytridiomycota	Chytridiomycotina	Caulochytriomycetes Chytridiomycetes
			Cladochytriomycetes
			Mesochytriomycetes
			Lobulomycetes
			Polychytriomycetes
			Rhizophlyctidomycetes
			Rhizophydiomycetes
			Spizellomycetes
			Synchytriomycetes
	Monoblepharomycota	Monoblepharomycotina	Hyaloraphidiomycetes
			Monoblepharidomycetes
	Neocallimastigomycota	Neocallimastigomycotina	Neocallimastigomycetes
Dikarya	Ascomycota	Pezizomycotina	Arthoniomycetes
			Dothideomycetes
			Eurotiomycetes
			Laboulbeniomycetes
			Lecanoromycetes
			Leotiomycetes
			Lichinomycetes
			<b>Orbiliomycetes</b>
			Pezizomycetes
			Sordariomycetes
			Xylobotryomycetes
		Saccharomycotina	Alloascoideomycetes
			Dipodascomycetes
			Lipomycetes
			Pichiomycetes
			Saccharomycetes
			Sporopachydermiomycetes
			Trigonopsidomycetes
		Taphrinomycotina	Archaeorhizomycetes
			Neolectomycetes
			Novakomycetes
			Pneumocystomycetes
			Schizosaccharomycetes
			Taphrinomycetes
	Basidiomycota	Agaricomycotina	Agaricomycetes
			Bartheletiomycetes
			Dacrymycetes
			Tremellomycetes
		Pucciniomycotina	Agaricostilbomycetes
			Atractiellomycetes
			Classiculomycetes
			Cryptomycocolacomycetes

<span id="page-3-0"></span>**Table 2** List of phyla, subphyla and classes in diferent subkingdoms (based on this study)

#### **Table 2** (continued)



# **Phyla of the Kingdom** *Fungi*

In this compilation, we accept 19 phyla (viz., *Aphelidiomycota*, *Ascomycota*, *Basidiomycota*, *Basidiobolomycota*, *Blastocladiomycota*, *Calcarisporiellomycota*, *Chytridiomycota* (=*Caulochytriomycota*), *Entomophthoromycota*, *Entorrhizomycota*, *Glomeromycota*, *Kickxellomycota*, *Monoblepharomycota*, *Mortierellomycota*, *Mucoromycota*, *Neocallimastigomycota*, *Olpidiomycota*, *Rozellomycota*, *Sanchytriomycota* and *Zoopagomycota*).

### *Aphelidiomycota* Tedersoo etal.

Tedersoo et al. ([2018\)](#page-157-0) introduced the phylum *Aphelidiomycota* to accommodate *Aphelidium* (Zopf) Gromov and three other related genera, *Amoeboaphelidium* Scherf., *Paraphelidium* Karpov et al. and *Pseudaphelidium* Schweikert & Schnepf. Members of *Aphelidiomycota* (i.e. aphelids) are a relatively poorly studied algal parasitic group (Karpov et al. [2014a,](#page-142-2) [2017a,](#page-142-3)[b](#page-142-4)). On the basis of early phylogenies, Karpov et al. ([2014a,](#page-142-2) [2017a](#page-142-3),[b\)](#page-142-4) regarded aphelids as members of a novel superphylum *Opisthosporidia*, which also included *Rozellomycota* (=*Cryptomycota*) along with *Microsporidia*. The proposed *Opisthosporidia* were viewed as a sister group to the fungal clade, but still retained in *Holomycota*. Subsequent analyses found no support for the union of aphelids and *Rozellomycota*, thus rendering the *Opisthosporidia* unsubstantiated. Accordingly, aphelids were proposed to constitute a distinct lineage of fungi at the phylum level (*Aphelidiomycota*) or even subkingdom level (*Aphelidiomyceta*) (Tedersoo et al. [2018\)](#page-157-0). Subsequent outlines by Wijayawardene et al. ([2018a,](#page-160-5) [2020a,](#page-160-0) [2022a](#page-160-1)) followed Tedersoo et al. ([2018\)](#page-157-0) and listed *Aphelidiomycota* as a distinct phylum in the fungal clade.

#### *Ascomycota* Caval.-Sm.

Cavalier-Smith ([1998\)](#page-133-2) introduced the phylum *Ascomycota*, one of the largest phyla in the kingdom. *Ascomycota* is further divided into three subphyla, *Pezizomycotina*, *Saccharomycotina* and *Taphrinomycotina* (James et al. [2006a;](#page-141-2) Liu et al. [2006](#page-145-3); Spatafora et al. [2006](#page-155-2); Sugiyama et al. [2006\)](#page-156-2). Currently, 24 classes have been listed under the phylum which comprises fungi isolated from a broad range of environments including terrestrial and aquatic ecosystems. The subphylum *Pezizomycotina* comprises multicellular, lichenized (e.g. Class *Lecanoromycetes*) and non-lichenized taxa (e.g. Class *Dothideomycetes*). The subphyla *Saccharomycotina* and *Taprinomycotina* comprise the yeasts assigned to *Ascomycota* which are "characterized by budding or fssion as the primary means of asexual reproduction, and have sexual states that are not enclosed in fruiting bodies" (Kurtzman et al. [2011\)](#page-143-0). Until recently *Saccharomycotina* has comprised a single class, *Saccharomycetes* and a single order *Saccharomycetales* (Kurtzman et al. [2011](#page-143-0)). More than 1,200 species were assigned to the single class and order of this subphylum. However, based on relative evolutionary divergence (RED) analyses, Li et al. ([2021a](#page-145-2)) revealed that the subphylum *Saccharomycotina* is seriously under-classifed. While it is characterised by higher levels of genomic diversity and evolutionary rates than *Pezizomycotina* (Shen et al. 2020), it contained only one class as opposed to the 16 classes of *Pezizomycotina*. The other sister subphylum, *Taphrinomycotina*, although it accommodates about ten times fewer species than *Saccharomycotina*, involves six classes. Recent genome-scale phylogenetic analyses divided the subphylum *Saccharomycotina* into 12 major clades (Shen et al. [2016](#page-154-5), [2018\)](#page-154-6). Groenewald et al. ([2023\)](#page-137-1) proposed a new genome-informed higher rank classifcation of *Saccharomycotina*. Based on genome-scale phylogenetic analyses and RED analyses they proposed an updated classifcation of *Saccharomycotina* refecting better the evolutionary divergence of the subphylum. According to the new classifcation *Saccharomycotina* consists of seven classes (*Alloascoideomycetes*, *Dipodascomycetes*, *Lipomycetes, Saccharomycetes*, *Pichiomycetes*, *Sporopachydermiomycetes* and *Trigonopsidomycetes*) and twelve orders. The third subphylum, *Taphrinomycotina*, mainly consists of yeast-like taxa.

#### *Basidiobolomycota* Doweld

Doweld ([2001\)](#page-135-0) introduced this phylum, to accommodate the class *Basidiobolomycetes* which also comprises of one order and one family, i.e. *Basidiobolales* Jacz. & P.A. Jacz. and *Basidiobolaceae* Engl. & E. Gilg. Tedersoo et al. [\(2018\)](#page-157-0) introduced the subkingdom *Basidiobolomyceta* Tedersoo et al. to accommodate *Basidiobolomycota* since it showed a distinct lineage from other subkingdoms. Currently, the phylum consists of one subphylum, *Basidiobolomycotina*.

### *Basidiomycota* R.T. Moore

Moore [\(1980](#page-148-2)) introduced *Basidiomycota*, which is a major phylum of the Kingdom *Fungi*, being second in species numbers to *Ascomycota* (Begerow et al. [2018](#page-131-0); He et al. [2019](#page-139-3)). The phylum comprises four subphyla, namely *Agaricomycotina*, *Pucciniomycotina*, *Ustilaginomycotina* and *Wallemiomycotina* (He et al. [2019,](#page-139-3) [2022a,](#page-139-2) [b](#page-139-4)). Whereas the typical mushroom-forming fungi are restricted to *Agaricomycotina* while *Pucciniomycotina* and *Ustilaginomycotina* are mainly characterised by phytoparasites. The members of *Basidiomycota* are mainly reported as saprobes, pathogens of animals and plants, and ectomycorrhizal species (He et al. [2022a\)](#page-139-2). Traditionally, fungi producing sexually by means of basidia have been assigned to this phylum. However, based on cultural, microscopic, and molecular phylogenetic methods, several genera with asexually reproducing species were also recognised as members of *Basidiomycota* (Hyde et al. [2011](#page-140-5); Wijayawardene et al. [2012,](#page-159-3) [2020a,](#page-160-0) [2021b,](#page-160-6) [2022a](#page-160-1)[,b](#page-160-4)). *Basidiomycetous* yeasts are also an important group in this phylum and occur in *Agaricomycotina*, *Pucciniomycotina*, and *Ustilaginomycotina* (e.g. Wang et al. [2015b;](#page-159-4) Oberwinkler [2017\)](#page-149-2).

#### *Blastocladiomycota* T.Y. James

James et al. [\(2006b](#page-141-3)) introduced the phylum *Blastocladiomycota* which was considered to be a monophyletic group within the phylum *Chytridiomycota* divergent from other zoosporic fungi, clustering among non-zoosporic fungi (Powell [2017\)](#page-150-1). The phylum *Blastocladiomycota* is further divided into the subphylum *Blastocladiomycotina* (Tedersoo et al. [2018](#page-157-0)) and consists of two classes, *Blastocladiomycetes* Doweld., and *Physodermatomycetes* Tedersoo et al. with less than 200 species described. They live in aquatic and terrestrial environments as saprotrophs or parasites of algal groups (Money [2016\)](#page-148-3).

#### *Calcarisporiellomycota* Tedersoo et al.

The phylum *Calcarisporiellomycota* was introduced by Tedersoo et al. [\(2018](#page-157-0)) and accommodates the genera *Calcarisporiella* de Hong., and *Echinochlamydosporium* Jiang et al. The phylum depicts a deep lineage with the phyla *Mucoromycota* (Hirose et al. [2012;](#page-139-5) Yamamoto et al. [2015\)](#page-161-1) or *Mortierellomycota* (Jiang et al. [2011](#page-141-4)). The vegetative structure shows a branched thallus with septate hyphae with uninucleate, ovoid to ellipsoid spores. The genera are also saprotrophic in soil (Jiang et al. [2011](#page-141-4); Hirose et al. [2012](#page-139-5)).

#### *Chytridiomycota* Doweld

#### =*Caulochytriomycota* Doweld

Doweld ([2001\)](#page-135-0) introduced the phylum *Chytridiomycota* and this was phylogenetically confrmed in subsequent studies (e.g. James et al. [2006a,](#page-141-2)[b](#page-141-3); Letcher et al. [2006;](#page-144-0) Mozley-Standridge et al. [2009](#page-148-4)). Tedersoo et al. [\(2018\)](#page-157-0) introduced the subkingdom *Chytridiomyceta* Tedersoo et al. to accommodate *Chytridiomycota*, *Monoblepharomycota* and *Neocallimastigomycota*. Currently, the phylum comprises one subphylum and ten classes. Members of the phylum are reported from both aquatic and terrestrial ecosystems, mainly as saprotrophic, with many facultative and obligate parasites of algae (Powell and Letcher [2014;](#page-150-2) Seto and Degawa [2017\)](#page-154-7). Galindo et al. [\(2021\)](#page-137-0) and Strassert and Monaghan ([2022a,](#page-156-0) [b\)](#page-156-1) regarded *Caulochytriomycota* as a synonym of *Chytridiomycota* and our analyses also agree with it.

#### *Entomophthoromycota* Humber

The phylum *Entomophthoromycota* was introduced by Humber ([2012](#page-140-6)) and is further divided into the subphylum *Entomophthoromycotina* and three classes, *Basidiobolomycetes, Entomophthoromycetes* and *Neozygitomycetes.* The phylum *Entomophthoromycota* consists of hyphal bodies or yeast-like cells with conidiophores. They are usually parasitic on algal groups where some might subsist as saprobes. The class *Basidiobolomycetes* contains unusually large nuclei with uninucleate cells, while *Neozygitomycetes* contains nuclei with vermiform, moderately sized chromosomes, with rod-like hyphal bodies containing 3 to 5 nuclei. The class *Entomophthoromycetes* lacks uniformly uninucleate cells or uniform number of nuclei, yet consists of coenocytic mycelium or rod-like to variably-shaped hyphal bodies (Humber [2012](#page-140-6)).

# *Entorrhizomycota* R. Bauer et al.

The phylum *Entorrhizomycota* was introduced by Bauer et al. [\(2015](#page-131-1)) to accommodate the genus *Entorrhiza* Weber. The phylum consists of one class, *Entorrhizomycetes*. It was further divided into order *Entorrhizales* Bauer et al., and *Talbotiomycetales* Riess et al. (Wijayawardene et al. [2022a](#page-160-1)). This taxon infects roots and its members have regularly septate coiled hypha. It is a small genus of only 14 species. However, with phylogenetic analyses and comparative studies it was found that this genus is a widespread but inconspicuous group of fungi (Bauer et al. [2015\)](#page-131-1).

#### *Glomeromycota* C. Walker & A. Schüßler

Schüssler et al. (2001) introduced *Glomeromycota* to accommodate arbuscular mycorrhizal fungi and Geosiphon pyriforme (Kütz.) F.Wettst. *Geosiphon* is the only member of the monophyletic group known to form symbioses with *Nostoc* (cyanobacteria) and not with plants (Schüssler et al. 2001). The placement of *Glomeromycota* in the fungal tree was accepted by several subsequent studies (e.g. James et al. [2006a;](#page-141-2) White et al. [2006](#page-159-5)). Nevertheless, Spatafora et al. ([2016\)](#page-155-1) who regarded *Glomeromycota* as 'phylogenetic enigma because it lacks any known form of sexual reproduction', diminished the Phylum *Glomeromycota* to Subphylum *Glomeromycotina* in *Mucoromycota*, along with *Mucoromycotina*, and *Mortierellomycotina*. However, Tedersoo et al. [\(2018](#page-157-0)) accepted the Phylum *Glomeromycota* in the Subkingdom *Mucoromyceta*. The fndings of Tedersoo et al. ([2018\)](#page-157-0) were followed by subsequent studies (Wijayawardene et al. [2018b](#page-160-7), [2020a](#page-160-0), [2022a;](#page-160-1) Voigt et al. [2021\)](#page-158-0).

#### *Kickxellomycota* Tedersoo et al.

Tedersoo et al. ([2018\)](#page-157-0) introduced the phylum *Kickxellomycota* in the subkingdom *Zoopagomyceta*, along with *Entomophthoromycota*, and *Zoopagomycota*. The phylum comprises one subphylum, *Kickxellomycotina* Benny (Hibbett et al. [2007](#page-139-1)). Currently, six classes are included in *Kickxellomycota* viz*., Asellariomycetes*, *Barbatosporomycetes*, *Dimargaritomycetes*, *Harpellomycetes*, *Kickxellomycetes*, and *Ramicandelaberomycetes* (Tedersoo et al. [2018\)](#page-157-0)*.* Members of *Kickxellomycota* have been reported as insect parasites, coprophilous, and soil-inhabiting taxa.

#### *Monoblepharomycota* Doweld

Doweld [\(2001\)](#page-135-0) introduced this phylum which is currently accommodated in subkingdom *Chytridiomyceta* (Tedersoo et al. [2018](#page-157-0)). Tedersoo et al. ([2018\)](#page-157-0) also introduced subphylum *Monoblepharomycotina* Tedersoo et al., the only subphylum in *Monoblepharomycota*. Currently, *Monoblepharomycota* comprises two classes, *Monoblepharidomycetes* J.H. Schafn. and *Hyaloraphidiomycetes* Doweld. Members of *Hyaloraphidiomycetes* have been reported from both aquatic and terrestrial environments as saprobes.

### *Mortierellomycota* Tedersoo et al.

*Mortierellomycota* was introduced by Tedersoo et al. ([2018\)](#page-157-0) and is regarded as a phylum in the subkingdom *Mucoromyceta*, along with *Calcarisporiellomycota*, *Glomeromycota*, and *Mucoromycota.* Previous studies based on phylogeny showed that his phylum has a distinct lineage (James et al. [2006a](#page-141-2); Sekimoto et al. [2011;](#page-154-8) Spatafora et al. [2016;](#page-155-1) Tedersoo et al. [2018](#page-157-0)).

#### *Mucoromycota* Doweld

The phylum *Mucoromycota* was introduced by Doweld ([2001](#page-135-0)). However, White et al. ([2006\)](#page-159-5) regarded *Mucorales*, the type order of *Mucoromycota*, as an order of *Zygomycota* while James et al. ([2006a](#page-141-2)) and Sekimoto et al. ([2011\)](#page-154-8) accepted subphylum *Mucoromycotina* in *Zygomycota*. Spatafora et al. [\(2016](#page-155-1)) accepted *Mucoromycota* and showed three subphyla within the phylum, i.e.

*Mucoromycotina*, *Mortierellomycotina*, and *Glomeromycotina*. However, *Glomeromycota* has been accepted as a well-defned phylum by several authors (e.g. Walker et al. [2007;](#page-158-1) Oehl et al. [2011a](#page-149-0), [b;](#page-149-1) Tedersoo et al. [2018;](#page-157-0) Wijayawardene et al. [2020a](#page-160-0); Voigt et al. [2021\)](#page-158-0). The subphylum *Mortierellomycotina* was upgraded to a phylum by Tedersoo et al. ([2018](#page-157-0)) thus *Mucoromycota* has one subphylum i.e. *Mucoromycotina.*

#### *Neocallimastigomycota* M.J. Powell

James et al. ([2006a,](#page-141-2) [b](#page-141-3)) recognized *Neocallimastigales* J.L. Li et al. (type: *Neocallimastix fde* Li et al. [1993](#page-145-4)) which formed a distinct clade in *Chytridiomycota.* Hibbett et al. ([2007](#page-139-1)) introduced the class *Neocallimastigomycetes* and the phylum *Neocallimastigomycota* to accommodate *Neocallimastigales.* Tedersoo et al. [\(2018\)](#page-157-0) introduced *Neocallimastigomycotina* Tedersoo et al., the only subphylum of *Neocallimastigomycota*.

#### *Olpidiomycota* Doweld

*Olpidiomycota* Doweld was established to accommodate *Olpidiales* (Doweld [2013a](#page-135-1)), which was earlier accepted in *Chytridiomycetes* Caval-Sm. as *Spizellomycetales* D.J.S. Barr (Barr [1980](#page-131-2)). The members of *Olpidiomycota* are cosmopolitan in distribution as parasites of economically important crop plants, algae and rotifers (Kirk et al. [2008](#page-142-1); Powell and Letcher [2014](#page-150-2); Lay et al. [2018](#page-144-1); Wijayawardene et al. [2018a,](#page-160-5) [2022a](#page-160-1), Naranjo‐Ortiz and Gabaldón [2019](#page-148-5)). The recent revision of *Fungi* and fungus-like taxa (Wijayawardene et al. [2022a](#page-160-1)) accepted one class, one order, one family and four genera in *Olpidiomycota*.

#### *Rozellomycota* Doweld

The placement of *Rozellomycota* and *Microsporidia* in the kingdom *Fungi* is a controversial topic in the higher-level classifcation of fungi and several major studies recognized them as early divergent groups of fungi (e.g. James et al. [2006a;](#page-141-2) Hibbett et al. [2007](#page-139-1); Corsaro et al. [2014](#page-134-0)). Nevertheless, some studies showed that *Aphelida* (*Aphelidiomycota*) and *Rozellomycota* as members of Opisthosporidia represent the sister groups to *Fungi* (Karpov et al. [2013](#page-142-5), [2014a](#page-142-2), [2017b](#page-142-4); Letcher et al. [2017\)](#page-145-5). In this study, we follow the treatment of Tedersoo et al. ([2018](#page-157-0)) and Voigt et al. [\(2021](#page-158-0)) and accept *Rozellomycota* in the kingdom *Fungi*.

#### *Sanchytriomycota* Galindo et al.

Galindo et al. ([2021\)](#page-137-0) introduce this phylum to accommodate two genera *Sanchytrium* Karpov & Aleoshin and *Amoeboradix* Karpov, López-García, Mamkaeva et Moreira (Karpov et al. [2018\)](#page-142-6), and emended the diagnoses of the class *Sanchytriomycetes* (Tedersoo et al.) Galindo et al. and the order *Sanchytriales* (Tedersoo et al.) Galindo et al. Voigt et al. [\(2021](#page-158-0)) accepted this phylum as a distinct phylum with a single class, order, family, and two genera.

#### *Zoopagomycota* Gryganskyi et al.

Spatafora et al. ([2016](#page-155-1)) introduced this phylum and recognized three subphyla, *Entomophthoromycotina* Humber, *Kickxellomycotina* Benny and *Zoopagomycotina* Benny. However, Tedersoo et al. ([2018](#page-157-0)) upgraded *Entomophthoromycotina* and *Kickxellomycotina* to phyla, thus currently *Zoopagomycota* comprises only one subphylum, *Zoopagomycotina.* Tedersoo et al. [\(2018](#page-157-0)) confrmed the placement of *Zoopagomycota* in the Subkingdom *Zoopagomyceta* along with *Entomophthoromycota* and *Kickxellomycota*.

### **Aim and the content of the compilation**

This study aims to provide a broad concept of the kingdom *Fungi* and present all fundamental concepts of its taxonomy and classifcation. Furthermore, we provide the phylogenomic tree of phyla of the Kingdom *Fungi*. Each phylum and its classes are meticulously described, accompanied by illustrations, life modes, distribution, and notes. We have also included controversial opinions prevalent among the research communities, thereby providing a comprehensive and up-to-date view of the current scientifc discourse.

# **Materials and methodology**

#### **Extraction of data and arrangement of the article**

This manuscript focuses only on classes and phyla of the kingdom *Fungi*. We follow the higher ranks of early divergent fungi (EDF) proposed by Tedersoo et al. ([2018](#page-157-0)) that have 16 phyla (*Aphelidiomycota*, *Basidiobolomycota*, *Blastocladiomycota*, *Calcarisporiellomycota*, *Caulochytriomycota*, *Chytridiomycota*, *Entomophthoromycota*, *Glomeromycota*, *Kickxellomycota*, *Monoblepharomycota*, *Mortierellomycota*, *Mucoromycota*, *Neocallimastigomycota*, *Olpidiomycota*, *Rozellomycota* and *Zoopagomycota*). Subsequently, Galindo et al. [\(2021](#page-137-0)) introduced *Sanchytriomycota*. However, several studies have suggested diferent arrangements for the higher ranks of EDF (Table [3](#page-8-0)). Besides, *Caulochytriomycota* was not accepted as a phylum by Voigt et al. ([2021\)](#page-158-0) and Strassert and Monaghan ([2022a,](#page-156-0) [b\)](#page-156-1).

Placements of the phyla in *Dikarya* (i.e. *Ascomycota*, *Basidiomycota* and *Entorrhizomycota*) have been confrmed in several studies (e.g. Tedersoo et al. [2018;](#page-157-0) Wijayawardene et al. [2020a\)](#page-160-0).

The nomenclature of fungi and the authors of fungal names follow Index Fungorum ([2024\)](#page-141-5), except for the genus originally described with *Rhizophagus populinus* P.A. Dang



<span id="page-8-0"></span>**Table 3** Other prevailing classes and phyla classifcations of early divergent fungi Ŕ l,  $\ddot{ }$  $\overline{a}$ ن<br>ب l, ÷ l, Ŕ ÷  $\frac{1}{2}$ Table 3 Oth

as type species of *Rhizophagus*, later replaced by *Rhizophagus intraradices*. We have adopted the arguments presented by Sieverding et al. ([2015\)](#page-155-4) and have opted to use the generic name *Rhizoglomus* Sieverd., G.A. Silva & Oehl instead of *Rhizophagus*. Nevertheless, this arrangement needs to be approved by the International Commission of Taxonomy of Fungi.

#### **Phylogenomic analysis**

The dataset for phylogenomic analysis was prepared using a set of 300 orthologous groups, constructed for phylogeny inferences in an earlier study (Mikhailov et al. [2022\)](#page-147-0). We have updated the original dataset, which featured 45 species of Holomycota, to include 40 more members of the fungal lineage, with emphasis on improving the sampling of poorly represented phyla. The genomic and transcriptomic data were collected from the NCBI GenBank database ([https://](https://www.ncbi.nlm.nih.gov/genbank/) [www.ncbi.nlm.nih.gov/genbank/](https://www.ncbi.nlm.nih.gov/genbank/)) and the JGI GenomePortal ([https://genome.jgi.doe.gov/portal/\)](https://genome.jgi.doe.gov/portal/). Prefltered transcriptomes of *Aphelidium* species were obtained from a public repository [\(https://fgshare.com/projects/Aphelida\\_](https://figshare.com/projects/Aphelida_Extended_Data/111539) [Extended\\_Data/111539\)](https://figshare.com/projects/Aphelida_Extended_Data/111539) (Galindo et al. [2022\)](#page-137-2). Orthologous sequences were identifed in the assembled collection of proteomes using a previously developed dataset expansion pipeline (Tikhonenkov et al. [2020\)](#page-157-1), involving HMMER searches (Eddy [2011\)](#page-136-0) with the alignment profles, and reciprocal BLAST searches (Altschul et al. [1997](#page-130-0)). The sets of identifed orthologous sequences were processed using an automated quality fltering procedure of PREQUAL (Whelan et al. [2018](#page-159-6)) with all-against-all sequence comparisons and a posterior probability fltering threshold of 0.95. The fltered sets of sequences were aligned with MAFFT using the localpair (L-INS-i) algorithm (Katoh and Standley [2013](#page-142-7)), followed by manual inspection of alignments using BioEdit (Hall [1999](#page-138-0)). Phylogenetic trees were reconstructed for each of the 300 orthologous groups using IQ-TREE (Nguyen et al. [2015](#page-148-6)) to identify and eliminate contamination detected in the sequence sets, specifcally extraneous sequences found in the *Coelomomyces lativittatus* transcriptomic data. The cleaned alignments were fnally processed using a custom mask, designed to exclude ambiguously aligned and lowcomplexity regions. The trimmed alignments were concatenated using SCaFoS (Roure et al. [2007\)](#page-152-1) into a matrix with 113 K amino acid sites.

Bayesian inference with the concatenated alignment was performed with PhyloBayes MPI (Lartillot et al. [2013](#page-144-2)) using the site-heterogeneous CAT-GTR model with four discrete Gamma-distributed rate categories. Four independent chains were run for 10 K cycles and monitored for convergence. The consensus Bayesian tree was reconstructed using the four chains with a 25% burn-in. Trees were also reconstructed for each of the analysis chains independently to confrm convergence to a unique tree topology within each chain. The mean posterior relative exchangeabilities and site-specifc equilibrium frequencies were extracted from the chain fles with a 25% burn-in using the PhyloBayes readpb utility, and converted for downstream analyses using the available scripts (Szantho et al. [2023](#page-156-3)).

Maximum likelihood analysis with the concatenated alignment was performed with IQ-TREE (Nguyen et al. [2015](#page-148-6)). Tree reconstruction employed the  $LG + C60 + F + G4$ profle mixture model, and node support was evaluated using the ultrafast bootstrap approximation (Hoang et al. [2018\)](#page-140-7) with 1000 replicates. Additionally, tree inferences were conducted using the posterior mean site frequency model (Wang et al. [2018\)](#page-159-7), employing the exchangeabilities and site-specifc frequencies derived from the PhyloBayes analysis, as introduced by the CAT-PMSF approach (Szantho et al. [2023\)](#page-156-3). The CAT-PMSF inference utilized PhyloBayes-optimized parameters obtained without a fxed tree topology constraint, although the sampled portions of the analysis chains displayed strict preference for a unique tree topology. Accordingly, tree reconstruction with the CAT-PMSF approach was also carried out independently with the parameters obtained from each of the four PhyloBayes chains. Node support for the CAT-PMSF tree inference was evaluated similarly with ultrafast bootstrap approximation using 1000 replicates.

Approximately unbiased tests (Shimodaira [2002](#page-154-9)) with alternative tree topologies were performed by IQ-TREE with 100,000 resamplings using the RELL method. The tree topologies, representing the conceivable phylogenetic relationships around the contentious nodes of the fungal phylogeny, were constructed using the MEGA software (Kumar et al. [2016\)](#page-143-1). Site-wise likelihoods were calculated under the  $LG + C60 + F + G4$  evolutionary model using the original ML tree for parameter estimation, and under the CAT-PMSF model in four variations, corresponding to the parameters derived from the four PhyloBayes analysis chains.

# **Results and discussion**

#### **Phylogenomic tree of** *Holomycota*

The phylogeny reconstructions were performed using maximum likelihood (ML) and Bayesian methods with a 300 gene dataset and a taxonomic selection balanced toward early-diverging fungal lineages (the dataset is available at Mendeley Data, <https://doi.org/10.17632/9jchyjrxmy.1>). For both reconstruction methods, we employed site-heterogeneous models of sequence evolution, as they provide a better approximation of natural processes (Lartillot and Philippe [2004](#page-144-3); Quang le et al. [2008\)](#page-151-1) and are generally more robust to reconstruction artefacts than site-homogenous models (Philippe et al. [2011](#page-150-3)). The phylogenies produced by both methods show agreement across most of tree nodes, yet similarly to earlier analyses (Galindo et al. [2021;](#page-137-0) Li et al. [2021a](#page-145-2); Strassert and Monaghan [2022a](#page-156-0), [b;](#page-156-1) Mikhailov et al. [2022](#page-147-0)), some of the key nodes of fungal phylogeny are unstable or show discordance between the analyses (Fig. [1](#page-10-0)).

Both ML and Bayesian trees fully support the early split of *Rozellomycota* from the fungal stem lineage, and the



<span id="page-10-0"></span>**Fig. 1** Phylogeny of *Holomycota* reconstructed with a 300-gene dataset. The consensus tree obtained using four CAT-PMSF inferences with parameters from each of the four chains of PhyloBayes, featuring support values from diferent reconstruction methods: ML inference with the  $LG + C60 + F + G4$  profile mixture model, Bayesian inference with the CAT-GTR model of PhyloBayes, and ML inference under CAT-PMSF; tree nodes where the reconstruction methods fail to achieve maximal support, or are in disagreement, are highlighted in red, and the corresponding support values are shown; the

support values in the ML analyses correspond to the ultrafast bootstrap approximation; the support values for PhyloBayes inference are posterior probabilities; for the Bayesian inference and the CAT-PMSF reconstructions the support values are depicted for each of the four analysis chains in the four-leaf clover arrangements, with the red-coloured clover leaves highlighting instances where the best tree for the chain/analysis deviated from the consensus shown here; the branch of *Sanchytriomycota* was shortened by one-third for the illustration

subsequent separation of *Aphelidiomycota* from the ancestral lineage of true fungi. This result is in agreement with the previous large-scale phylogenies (Torruella et al. [2018](#page-157-2); Galindo et al. [2022\)](#page-137-2), which similarly found no support for the superphylum *Opisthosporidia*, hypothesized to house *Aphelidiomycota* and *Rozellomycota* (Karpov et al. [2014a\)](#page-142-2). The trees also agree on the sister position of *Neocallimastigomycota* and *Monoblepharomycota* to the *Chytridiomycota*, and the placement of the whole lineage as the earliest branch of true fungi.

The sole sequenced representative of the proposed phylum *Caulochytriomycota* (Doweld [2014a](#page-135-2)), *Caulochytrium protostelioides*, is found nested within *Chytridiomycota*, joining support for its deranking to a member of *Chytridiomycota* (Ahrendt et al. [2018](#page-130-1); Strassert and Monaghan [2022a,](#page-156-0) [b](#page-156-1)). Yet, the reconstructions fail to fnd the exact placement of *Caulochytrium* relative to the chytrid orders—its position within chytrids varies in both ML and Bayesian analyses (Supplementary Figs. 1 & 2). The newly proposed phylum *Sanchytriomycota* (Galindo et al. [2021](#page-137-0)), which includes two species, *Amoeboradix gromovii* and *Sanchytrium tribonematis*, with highly divergent sequences, is fully supported as a sister group to *Blastocladiomycota*, together forming the second earliest branch of true fungi. The last lineage to diverge from the fungal stem before the emergence of terrestrial non-zoosporic fungi is a chytrid-like fungus *Olpidium* (Chang et al. [2021](#page-133-1)). The Bayesian reconstruction fully supported the sister position of *Olpidium* to the non-zoosporic fungi. In the ML analysis, however, the support wavered, with 7% of bootstrap replicates instead placing *Olpidium* further into the clade of non-zoosporic fungi, sister to the group uniting *Mucoromycota*, *Mortierellomycota*, *Calcarisporiellomycota*, *Glomeromycota*, and *Dikarya*.

Remarkably, most internal tree nodes, where the analyses failed to reach a consensus, involve groups of terrestrial fungi rather than the deeper branching zoosporic fungal phyla. The most prominent confict is observed in the relative branching of *Mucoromycota*, *Mortierellomycota*, *Glomeromycota*, and *Dikarya*. While all reconstructions agree on the sister position of *Calcarisporiellomycota* to *Mucoromycota*, confrming earlier rDNA trees (Hirose et al. [2012;](#page-139-5) Tedersoo et al. [2018](#page-157-0)), the exact phylogenetic context of the *Mucoromycota*+*Calcarisporiellomycota* group varies in the analyses. In the ML tree the *Mucoromycota* (incl. *Calcarisporiellomycota*), *Mortierellomycota*, and *Glomeromycota* form a fully-supported monophyletic group sister to *Dikarya*, with *Glomeromycota* being more closely related to *Mucoromycota* than *Mortierellomycota*, at 89% bootstrap support (Supplementary Fig. 1). In the Bayesian inference, which was conducted using four independent analysis chains, we observed three diferent arrangements for these phyla; one variant repeating the ML tree branching order, and the other variants placing *Glomeromycota* sister to *Dikarya*, with *Mortierellomycota* appearing either as sisters to *Mucoromycota*+*Calcarisporiellomycota* or as sisters to the whole lineage comprising *Mucoromycota, Glomeromycota,* and *Dikarya* (Supplementary Fig. 2). Lineages of former zygomycete fungi that were collectively referred to as *Zoopagomycota* in prior taxonomic revisions (Spatafora et al. [2016\)](#page-155-1) formed a monophyletic group sister to other terrestrial fungi in all analyses, but the relationship between the constituent phyla—*Basidiobolomycota*, *Entomophthoromycota*, *Kickxellomycota*, *Zoopagomycota* sensu stricto—difered between the analyses. The Bayesian inference fully supported the early branching of *Basidiobolus* within the group and the sister position of *Zoopagomycota* to the monophyletic group uniting *Entomophthoromycota* and *Kickxellomycota* (Supplementary Fig. 2). The ML tree placed *Basidiobolus* sister to *Zoopagomycota*, but neither the *Basidiobolomycota*+*Zoopagomycota* nor the *Entomophthoromycota*+*Kickxellomycota* were highly supported (Supplementary Fig. 1).

The conflicting results of the ML and Bayesian trees around the position of *Glomeromycota* were noted previously in phylogenomic analyses (Galindo et al. [2021;](#page-137-0) Mikhailov et al. [2022;](#page-147-0) Strassert and Monaghan [2022a](#page-156-0), [b](#page-156-1)), and were shown to stem from the inherent limitations of the site-heterogeneous profle mixture model implemented in the ML treebuilding software (Strassert and Monaghan [2022a](#page-156-0), [b\)](#page-156-1). Thus, the disagreement between the ML and Bayesian trees on the relative branching of *Mucoromycota*, *Mortierellomycota*, *Glomeromycota*, and *Dikarya* was anticipated for the updated dataset. Unexpectedly, however, our Bayesian inference itself also failed to converge on a single branching arrangement for these groups. Past 1.5 K cycles and up to 10 K cycles, when the run was halted, each independent chain of the inference strictly adhered to its own tree topology, resulting in an ambiguous relationship between these groups in the consensus of the four chains. In an attempt to resolve this ambiguity, we have applied the CAT-PMSF method (Szantho et al. [2023](#page-156-3)), reconstructing additional ML trees using the evolutionary model parameters obtained from the Bayesian analysis chains. Reconstructions under the CAT-PMSF managed to reconcile the position of *Glomeromycota*, placing the group sister to *Dikarya* with nearly full support for all four chains, despite the conficting result of the Bayesian inference (Fig. [1\)](#page-10-0). The CAT-PMSF reconstruction also switched toward supporting the sister position of *Caulochytrium* to the *Spizellomycetales* and *Rhizophydiales* in *Chytridiomycota* for the one chain of Bayesian inference, where an alternative placement of *Caulochytrium* sister to *Cladochytriales* and *Chytridiales* was observed. In other instances where the individual Bayesian trees were incongruent, the CAT-PMSF trees also reproduced the disagreement. These include the placement of *Mortierellomycota*, the positions of *Pneumocystis* and *Schizosaccharomyces* in *Taphrinomycotina*, the relative branching of *Mucor*, *Absidia*, and *Phycomyces* in *Mucoromycota*, and the position of *Polychytrium* in *Chytridiomycota*. Notably, CAT-PMSF analysis also lowered support for several tree nodes where the Bayesian trees were resolute and even fully supported by the original ML analysis with the site-heterogeneous profle mixture model. The latter include the positions of *Saitoella* and *Neolecta*, and the monophyly of *Taphrinomycotina* (Fig. [1\)](#page-10-0).

To characterize the spectrum of ambiguous tree topologies systematically, we carried out hypothesis testing with an exhaustive set of alternatives around the conficting nodes of fungal phylogeny. The hypotheses were tested using the approximately unbiased (AU) test under the site-heterogeneous profle mixture model and the CAT-PMSF models derived from the Bayesian inference. For the tree nodes involving the relative branching of *Mucoromycota*, *Mortierellomycota*, *Glomeromycota*, and *Dikarya*, the profle mixture model accepts only two variants, both of which include the union of *Mucoromycota*, *Mortierellomycota*, and *Glomeromycota* (Supplementary Table 1). In contrast, tests under the CAT-PMSF reject any such variant at the 5% significance level. CAT-PMSF did not reject only variants with *Glomeromycota* sister to *Dikarya*. These include a variant with *Mortierellomycota* sister to the whole lineage of *Mucoromycota*, *Glomeromycota*, and *Dikarya* or a variant with a more commonly seen sister relationship between *Mortierellomycota* and *Mucoromycota* (incl. *Calcarisporiellomycota*). The latter group, however, was rejected in one of the three chains tested with CAT-PMSF. In *Chytridiomycota* the profle mixture model fails to reject 6 out of 15 possible arrangements for the lineages of *Caulochytrium*, *Polychytrium*, *Spizellomycetales*+*Rhizophydiales*, and *Cladochytriales*+*Chytridiales*. With CAT-PMSF there are only two accepted topologies for chytrids, which include a sister relationship between *Caulochytrium* and *Spizellomycetales*+*Rhizophydiales*, with *Polychytrium* emerging either as a sister to *Cladochytriales*+*Chytridiales* or at the base of the whole group. In one of the chains, an additional topology with *Caulochytrium* sister to *Cladochytriales*+*Chytridiales* also appears as a likely variant. Tests for the relative branching of *Basidiobolomycota*, *Entomophthoromycota*, *Kickxellomycota*, and *Zoopagomycota* fail to reject 5 out of 15 arrangements with the profle mixture model. With CAT-PMSF only two topologies for these groups are accepted, which include a monophyletic group of *Entomophthoromycota*+*Kickxellomycota* with either *Basidiobolus* or *Zoopagomycota* as its sister lineages. Tests with CAT-PMSF reject an alternative placement of *Olpidium* within non-zoosporic fungi, which was observed in a small percentage of bootstrap replicates of the original ML tree and was accordingly not rejected by the test with the profle mixture model. For members of *Taphrinomycotina,* all but one of the tree topologies were rejected by the test with the profle mixture model. CAT-PMSF, on the other hand, failed to reject four topologies, including those where *Saitoella* is sister to *Taphrinomycotina* or where *Taphrinomycotina* are

polyphyletic and *Saitoella*+*Neolecta* emerge as the earliest branch of *Ascomycota* or *Saitoella* alone moves to the *Pezizomycotina*+*Saccharomycotina* branch.

#### **Taxonomy**

In this section, all the phyla are provided with important taxonomic data, history, distribution, life cycle data, and life or nutritional modes. The classes of each phylum (Fig. [2\)](#page-13-0) are listed according to the placements in the respective subclasses (if more than one subclass).

*APHELIDIOMYCOTA* Tedersoo, Sanchez-Ramirez, Kõljalg, Bahram, M. Döring, Schigel, T.W. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 147 (2018)

Syn. *Aphelida* Karpov, Aleoshin & Mikhailov Index Fungorum Registration Identifer 553,990

Amoeboid endobiotic parasites of algae. Dispersal amoeboid zoospores attach to a new host cell and encyst (either sessile on the substrate or producing a stalk; "apophyse"; Gromov [2000](#page-137-3)). Amoeboid body penetrates the host's cell through a cyst stalk. The intracellular amoeba engulfs the contents of the host's cell, forming food vacuoles that transport food into the central digestive vacuole. An excretory body is formed in the digestive vacuole. The amoeboid trophont grows into a plasmodium, which totally replaces the cytoplasm of a host cell; the multinuclear plasmodium develops into an unwalled sporangium and divides into uninucleate amoeboid zoospores. Intracellular resting spores are present (Karpov et al. [2014a](#page-142-2)).

#### **Life modes and distribution:** Parasites of algae

**Notes:** The aphelids are a small group of intracellular parasites of algae, including three freshwater genera, *Aphelidium* (Zopf) Gromov (Fig. [3\)](#page-14-0), *Amoeboaphelidium* Scherf., *Paraphelidium* Karpov et al. and a marine genus, *Pseudaphelidium* Schweikert & Schnepf (Schweikert and Schnepf [1996](#page-154-10), [1997](#page-154-11); Gromov [2000](#page-137-3); Karpov et al. [2017a\)](#page-142-3).

Only the asexual part of the life cycle is known for aphelids: amoeboid zoospore attaches to the host algae, and involves a fagellum and encysts. A cyst germinates and penetrates the host cell wall with an infection tube and injects the cyst contents into the host. The parasite becomes the intracellular phagotrophic amoeba which engulfs the host cytoplasm with pseudopodia and forms a central digestive vacuole containing a red residual body. The parasite grows forming an endobiotic multinucleate plasmodium with the residual body in a large central vacuole while it consumes the whole cytoplasm of the host cell. The mature plasmodium then divides into a number of uninucleated cells. After maturation, the amoeboid zoospores are released from the empty host cell through the hole made earlier by the infection tube and infect other algae.



<span id="page-13-0"></span>**Fig. 2** Diagrammatic representation of phyla, subphyla and classes of the kingdom *Fungi*; **a** EDF. **b** *Dikarya*

All known aphelids can produce multiple infections, thus, the trophonts from diferent strains/populations may fuse to each other during the development of plasmodium which can possess more than one sort of nuclei (Tcvetkova et al. [2023](#page-156-4)).

The phylogenetic position of the aphelids became clear after five genes sequencing of *Amoeboaphelidium protococcorum* (Karpov et al. [2013\)](#page-142-8). Subsequently, Letcher et al. [\(2013](#page-145-6)) confrmed this result on the base of SSU gene sequence of *Am. occidentale*. The aphelids grouped with *Rozella* and microsporidia forming a cluster sister to *Fungi*, which was called the super phylum *Opisthosporidia* a year later (Karpov et al. [2014a](#page-142-2)). The rank of the aphelids has been raised from the class *Aphelidae* (Gromov [2000](#page-137-3)) to the phylum *Aphelida* (Karpov et al. [2014a](#page-142-2), [b\)](#page-142-9). The position of the aphelids on ribosomal trees was not stable. It varied from sister group to *Fungi*+*(Rozella*+*Microsporidia*) (Corsaro et al. [2014,](#page-134-0) [2019](#page-134-1)), to *Rozella*+*Microsporidia* (in most published trees), and to the Fungi (Tedersoo et al. [2018](#page-157-0)). Many mycologists included aphelids in the *Fungi* based predominantly on their life cycle, which superficially resembles those of zoosporic fungi, and on the monophyly of the cluster including *Fungi* and *Opisthosporidia* (James et al. 2013). The subsequent outline by Wijayawardene et al. [\(2020a\)](#page-160-0) followed Tedersoo et al. ([2018](#page-157-0)) and listed *Aphelidiomycota* as a distinct phylum in the fungal clade.

Intensive study of the aphelids during last decade gave more knowledge based on the light microscopic and ultrastructural observations, and molecular phylogeny including genomics. A new genus *Paraphelidium* with two species was described (Karpov et al. [2017a](#page-142-3),[b\)](#page-142-4), several new species in *Aphelidium* (Letcher et al. [2017;](#page-145-5) Tcvetkova et al. [2019](#page-156-5); Seto et al. [2020,](#page-154-12) [2022](#page-154-13)) and *Amoeboaphelidium* (Letcher et al. [2015\)](#page-145-7) were added based on morphological and molecular data. Genomes of three (Mikhailov et al. [2022;](#page-147-0) Pozdnyakov et al. [2023](#page-150-0)) and transcriptomes of four aphelid species (Torruella et al. [2018](#page-157-2); Galindo et al. [2022](#page-137-2)) were assembled, and a multigene phylogeny unambiguously showed a sister position of the *Aphelida* and fungi.

Karpov et al. [\(2014a,](#page-142-2) [b](#page-142-10)) believe that phylum *Aphelidiomycota* (=*Aphelida*) having a phagotrophic cellular mode of nutrition does not belong to the kingdom *Fungi* with the totally saprotrophic cellular mode of nutrition. From this view, the original phylum name *Aphelida* (Karpov et al. [2014a,](#page-142-2) [b\)](#page-142-10) has to be retained. We temporarily accept the layout of the present paper dealing with *Fungi* sensu lato*,* which also includes here the phagotrophic rozellids and follows the taxonomy by Tedersoo et al. ([2018](#page-157-0)).

Currently, the phylum consists of one subphylum (*Aphelidiomycotina*) and one class (*Aphelidiomycetes*).

*Aphelidiomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 147 (2018); Fig. [3](#page-14-0)

Index Fungorum Registration Identifer 553991

The class description is similar to the description of the phylum

**Type order:** *Aphelidiales* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov 2018



<span id="page-14-0"></span>**Fig. 3** Light microscopic images of three stages of life cycle of *Aphelidium* spp. life cycle. Modifed after: Karpov et al. [\(2014a\)](#page-142-2) (**e**, **f**, **g**), 2020 (**a**, **b**, **h**), Tcvetkova et al. ([2019\)](#page-156-5) (**c**, **d**). **a**–**e** – zoospores of *Aph. insulamus* (**a**, **b**), *Aph. arduennense* (**c**, **d**) and *Aph. melosirae* (**e**); **a**, **c**, **e** Swimming zoospores. **b**, **d** Crawling zoospores with sim-

**Type family**: *Aphelidiaceae* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 147 (2018)

**Type genus:** *Aphelidium* Zopf, Encyklop. Naturwiss. (Breslae): 30 (1885)

**Type species:** *Aphelidium deformans* Zopf, Encyklop. Naturwiss. (Breslae): 30 (1885)

# *ASCOMYCOTA* Caval.-Sm., Biol. Rev. 73: 247 (1998)

Index Fungorum Registration Identifer 90031

Sexual morph: Reproducing by formation of an ascoma or rarely naked, without an organised ascoma. Ascomata when present apothecial, perithecial, pseudothecial or cleistothecial. Ascomatal cavity when present with a centrum or hamathecium composed of cellular or flamentous, thread-like paraphyses or pseudoparaphyses. Asci unitunicate—operculate, unitunicate-inoperculate, prototunicate, bitunicate. Ascospores with varied shapes, colour, septation, size, with or without surface ornamentation and

ple flopodia (**f**) and branching flopodia (bf). **f** Plasmodium of *Aph. melosirae* in the host cell with central digestive vacuole (cv) containing residual body (rb). **g** Rounded resting spore (rs) of *Aph. melosirae*. **h** elongated resting spore (rs) of *Aph. insulamus*. Scale bars:  $a-e=3 \mu m$ ,  $f-h=10 \mu m$ 

appendages. Asexual morph: coelomycetous—producing pycnidia or acervuli or intermediate forms. Hyphomycetous—producing sporodochia, synnemata or individual conidiophores; conidia produced by diferent modes—arthric, blastic (holoblastic and enteroblastic), phialidic, annelidic, basauxic; conidia with diferent shapes, sizes, dry or wet, with or without surface appendages. Yeasts—budding or fission as the primary means of asexual reproduction, and have sexual morph that are not enclosed in fruiting bodies.

**Life modes and habitats:** Saprobes, pathogens of plants and animals, or lichen-forming, lichenicolous, fungicolous, endophytes, yeast-like or flamentous.

**Notes:** Whittaker ([1959\)](#page-159-8) introduced the term Ascomycota which was invalid. Cavalier-Smith [\(1998\)](#page-133-2) introduced the legitimate term, and subsequent studies accepted it (e.g. Hibbett et al. [2007](#page-139-1)). *Ascomycota* is the largest phylum in the kingdom of *Fungi*. The higher-level classifcation of the phylum was

frst compiled as the 'Outline of *Ascomycota'* by Eriksson and Hawksworth ([1986\)](#page-136-1). Subsequent updates and changes were published by Eriksson and Winka ([1997](#page-136-2)), and Lumbsch and Huhndorf [\(2007](#page-146-1), [2010\)](#page-146-2). Important studies of the taxonomy and classifcation of *Ascomycota* and its classes since Lumbsch and Huhndorf [\(2010](#page-146-2)) are listed in Table [4](#page-15-0).

<span id="page-15-0"></span>**Table 4** Important studies (on taxonomy and classifcation) published since Lumbsch and Huhndorf ([2010\)](#page-146-2) on *Ascomycota* and its classes

Article	Rank	Content
Rosling et al. (2011)	Archaeorhizomycetes	Taxonomy/classification/phylogeny
Gazis et al. (2012)	Xylonomycetes	Taxonomy/classification
Hansen et al. (2013)	Pezizomycetes	Phylogeny/classification
Hustad et al. $(2013)$	Geoglossomycetes	Phylogeny/classification
Hyde et al. (2013)	Dothideomycetes	Taxonomy/classification
Prieto et al. (2013)	Candelariomycetes	Phylogeny/classification
Beimforde et al. (2014)	Ascomycota	Estimating the Phanerozoic history
Ertz et al. (2014)	Arthoniomycetes	Phylogeny/classification
Gueidan et al. (2014)	Eurotiomycetes	Phylogeny/classification
Johnston et al. (2014)	Leotiomycetes	Taxonomy/classification
Menkis et al. (2014)	Archaeorhizomycetes	Phylogeny/classification
Miądlikowska et al. (2014)	Lecanoromycetes	Phylogeny/classification
Haelewaters et al. (2015)	Laboulbeniomycetes	Phylogeny/classification
Maharachchikumbura et al. (2015)	Sordariomycetes	Taxonomy/classification
Shen et al. (2016)	Saccharomycotina	Taxonomy/phylogeny
Baral et al. (2018)	Xylonomycetes	Phylogeny
Ekanayaka et al. (2017)	Neolectomycetes	Taxonomy/classification
Hongsanan et al. (2017)	Sordariomycetes	Phylogeny/classification based on molecular clock evidence
Liu et al. (2017)	Dothideomycetes	Ranking higher taxa using divergence times
Lücking et al. (2017)	Arthoniomycetes	Taxonomy/classification
Lücking et al. (2017)	Lichinomycetes	Taxonomy/classification
Wijayawardene et al. (2017a)	Ascomycota	Notes for the genera in Ascomycota
Baral et al. (2017)	<i>Orbiliomycetes</i>	Taxonomy/classification
Goldmann and Weir (2018)	Laboulbeniomycetes	Phylogeny/classification
Hibbett et al. (2018)	Ascomycota	Phylogenetic taxon definition
Rikkinen et al. (2018)	Coniocybomycetes	Fossil study
Tedersoo et al. $(2018)$	Collemopsidiomycetes	Phylogeny
Wijayawardene et al. (2018a)	Ascomycota	Outline/classification
Ametrano et al. (2019)	Dothideomycetes	Phylogeny
Haelewaters et al. (2019)	Laboulbeniomycetes	Phylogeny/classification
Johnston et al. (2019)	Leotiomycetes	Phylogeny/classification
Samarakoon et al. (2019)	Ascomycota	Divergence time calibrations for ancient lineages
Voglmayr et al. (2019)	Candelariomycetes	Phylogeny/classification
Voglmayr et al. (2019)	Xylobotryomycetes	Phylogeny/classification
Blackwell et al. (2020)	Laboulbeniomycetes	Evolution
Haelewaters et al. (2020)	Laboulbeniomycetes	Phylogeny/taxonomy/classification
Hyde et al. (2020)	Sordariomycetes	Taxonomy/classification
Shen et al. (2020)	Ascomycota	Phylogeny
Hongsanan et al. (2020a,b)	Dothideomycetes	Taxonomy/classification
Čadež et al. (2021)	Novakomycetes	Phylogeny/classification
Haelewaters et al. (2021a)	Laboulbeniomycetes	Phylogeny/classification
Haelewaters et al. (2021c)	Leotiomycetes	Phylogeny/taxonomy/classification
Hashimoto et al. (2021)	Xylonomycetes	Phylogeny/classification
Latinne et al. (2021)	Pneumocystidomycetes	Phylogeny/host specificity
Díaz-Escandón et al. (2022)	Lichinomycetes	Phylogeny
Brysch-Herzberg et al. (2022, 2023)	Schizosaccharomycetes	Taxonomy/phylogeny

<span id="page-16-0"></span>

<b>Table 5</b> Subphyla and classes of Ascomycota	Phylum	<b>ASCOMYCOTA</b>		
	Subphyla	Pezizomycotina	Saccharomycotina	Taphrinomycotina
	Classes	Arthoniomycetes Dothideomycetes Eurotiomycetes Laboulbeniomycetes Lecanoromycetes Leotiomycetes Lichinomycetes <i>Orbiliomycetes</i> Pezizomycetes Sordariomycetes	Alloascoideomycetes Dipodascomycetes Lipomycetes Pichiomycetes Saccharomycetes Sporopachydermiomycetes Trigonopsidomycetes	Archaeorhizomycetes Neolectomycetes Novakomycetes Pneumocystomycetes Schizosaccharomycetes Taphrinomycetes
		Xylobotryomycetes		

Currently, *Ascomycota* comprises three subphyla and 24 classes (Table [5](#page-16-0)).

### **Pleomorphism and one fungus one name**

Saccardo ([1904\)](#page-152-4) suggested implementing the Dual Nomenclature system for pleomorphic fungi. Many ascomycetous taxa have been named with two names when they are reported with two diferent morphs. However, on 30 July 2011, the dual nomenclature system was abandoned, thus it was encouraged to link diferent morphs and propose one name for one fungus. Several important publications have been published during the last decade, and these are summarized in Table [6.](#page-16-1)

#### **Evolutionary studies of** *Ascomycota*

In recent years, *Ascomycota* have been the focus of in-depth molecular dating, paleoecological reconstruction, and morphological character evolution studies. Taylor and Berbee [\(2006\)](#page-156-6) reviewed the molecular-based divergence time estimation for fungi and provided estimated divergence times for the major lineages of the Fungal Tree of Life. A remarkable fossil, *Paleopyrenomycites devonicus*, discovered in the cortex just beneath the epidermis of aerial stems and rhizomes of the vascular plant *Asteroxylon* (Taylor et al. [2005](#page-156-7)), shaped the subsequent evolutionary studies in *Ascomycota* significantly. Based on different calibrations, including signifcant ascomycetous fossil data, the divergence of the *Ascomycota* may have occurred during the 512–588 MYA (Gueidan et al. [2011](#page-138-8); Prieto and Wedin [2013](#page-150-5); Beimforde et al. [2014;](#page-131-3) Pérez-Ortega et al. [2016](#page-149-3)). A recent genome-scale evolutionary study shows the origin of the *Ascomycota* has taken place at 563 MYA (631–495 MYA), while the origin of the *Pezizomycotina* is at 407.7 MYA (631–405 MYA), the *Saccharomycotina* at 438.4 MYA (590–304 MYA), and the *Taphrinomycotina* at 530.5 MYA (620–417 MYA) (Shen et al. [2020](#page-154-14)). Table [7](#page-17-0) summarizes the signifcant publications related to the evolution of *Ascomycota* published after Taylor and Berbee [\(2006\)](#page-156-6).

<span id="page-16-1"></span>**Table 6** Important studies focused on one fungus, one name in *Ascomycota*



**Subphylum** *Pezizomycotina* O.E. Erikss. & Winka, Myconet 1(1): 9 (1997)

Index Fungorum Registration Identifer 501468

**Notes:** Eriksson and Winka ([1997\)](#page-136-2) introduced the subphylum *Pezizomycotina* to accommodate *Arthoniomycetes*, *Dothideomycetes*, *Eurotiomycetes*, *Lecanoromycetes, Leotiomycetes*, *Pezizomycetes* and *Sordariomycetes.* Hibbett et al. ([2007](#page-139-1)) listed *Laboulbeniomycetes* under *Pezizomycotina*. Subsequent studies by Eriksson et al. [\(2003](#page-136-5)), Reeb et al. ([2004](#page-151-3)) and Voglmayr et al. ([2019\)](#page-158-2) introduced *Lichinomycetes*, *Orbiliomycetes*, and *Xylobotryomycetes* respectively. Currently, *Pezizomycotina* has eleven classes (Table [5\)](#page-16-0).

# *Arthoniomycetes* O.E. Erikss. & Winka, Myconet 1(1): 4 (1997)

Index Fungorum Registration Identifer 501475; Fig. [4](#page-18-0)

Thallus lichenized with chlorococcoid or trentepohlialean algae or non-lichenized, usually well-developed, crustose, leprarioid, byssoid, placodioid or fruticose, or in form of black yeasts (*Phaeococcomyces* de Hoog) or stromatic; vegetative lichenized diaspores (isidia, soredia or pseudoisidia)



<span id="page-17-0"></span>**Table 7** Important studies (evolution and ancestral character reconstruction related) published since Taylor and Berbee [\(2006](#page-156-6)) on *Ascomycota* and its classes and subclasses

present in some species; ascomata usually present, stromatic (e.g., *Etayoa* Diederich & Ertz, *Lichenostigma* Hafellner), apothecioid, rounded to stellate, lirelliform or not welldefned and globose asci loosely scattered in the thallus (*Crypthothecia* Stirt., *Myriostigma* Kremp.); hamathecium present in form of paraphysoids (between asci or enclosing the asci) or absent; asci bitunicate, globose to clavate, with a strongly thickened tholus, with  $K$ ,  $I + blue$  ring in the

<span id="page-18-0"></span>**Fig. 4** Morphology of selected members of *Arthoniomycetes*. **a** *Alyxoria varia* (Poland, UGDA L-41544). **b** *Arthonia radiata* – type species (Poland, UGDA L-60149). **c** *Coniocarpon cinnabarinum* (Poland, UGDA L-20564). **d** *Herpothallon rubrocinctum* with pseudoisidia (Bolivia, UGDA L-19608). **e** *Lecanactis abietina* with pycnidia (Poland, UGDA L-25338). **f** *Lichenostigma chlaroterae*, stromata on *Lecanora pulicaris* (Poland, UGDA L-21140). **g** *Myriostigma napoense* with not well-defned ascomata (Bolivia, UGDA L-19564). **h** *Roccella tinctoria* (Canary Islands, UGDA L-20824). Scale bars: **a**, **b**, **c**, **e**, **f**=500 µm; **d**,  $g=1000 \mu m; h=10 \text{ mm}$ 



lower part of the apical dome (near the ocular chamber), sometimes also ascus tholus amyloid; ascospores hyaline to brown, simple, transversely septate or muriform. *Conidia* simple to muriform, produced by budding or in conidiomata (pycnidia or sporodochia). Lichenized species often produce secondary lichen metabolites. In general taxa can be included in *Arthoniomycetes* based on unique to the class asci with thickened tholus, with  $KI + blue$  ring in the lower part of the apical dome, and/or on the molecular data, especially in case of sterile, peculiar species (Tehler et al. [2010](#page-157-3); Ertz and Tehler [2011](#page-136-6); Ertz et al. [2011](#page-136-7), [2014](#page-136-3), [2015](#page-136-8); Diederich et al. [2012](#page-135-4); Frisch et al. [2014,](#page-137-6) [2015](#page-137-7); Thiyagaraja et al. [2020](#page-157-4)).

**Life modes and distribution**: Lichenized, saprotrophic or lichenicolous (parasitic or commensals), on bark of trees, wood, rocks, leaf litter, and other lichens (apothecia or thallus), epiphyllous, cosmopolitan, members of several cryptogamous communities, but do not play important role in building main biomes (Lutzoni et al. [2004;](#page-146-8) Spatafora et al. [2006](#page-155-2); Ertz et al. [2009,](#page-136-9) [2011,](#page-136-7) [2014,](#page-136-3) [2015;](#page-136-8) Tehler et al. [2010](#page-157-3); Ertz and Tehler [2011;](#page-136-6) Diederich et al. [2012;](#page-135-4) Frisch et al. [2014,](#page-137-6) [2015;](#page-137-7) Thiyagaraja et al. [2020\)](#page-157-4), cosmopolitan, but common in temperate to tropical regions, in coastal, desert, mountainous, or forest habitats.

**Notes**: At frst, the species now placed in *Arthoniales* were accepted in two orders, *Arthoniales* and *Opegraphales* M. Choisy ex D. Hawksw. & O.E. Erikss., by Hawksworth and Eriksson ([1986](#page-139-9)). The similarity and close relationships of *Arthoniales* and *Opegraphales* were pointed out by Hafellner ([1988](#page-138-11)) and later included in the class *Arthoniomycetes* by Eriksson and Winka ([1997\)](#page-136-2) within the newly established superclass *Leotiomyceta* O.E. Erikss. & Winka. Lutzoni et al. ([2004\)](#page-146-8) suggested a sister relationship with *Dothideomycetes* O.E. Erikss. & Winka, but with low support, and Spatafora et al. ([2006\)](#page-155-2) confrmed this relationship. At present, *Arthoniomycetes* includes two orders, *Arthoniales* (syn. *Opegraphales*) and *Lichenostigmatales*, the latter with one family (*Phaeococcomycetaceae* McGinnis & Schell) and three genera. The majority of taxa are placed in *Arthoniales*, within seven families (*Andreiomycetaceae* B.P. Hodk. & Lendemer, *Arthoniaceae*, *Chrysotrichaceae* Zahlbr., *Lecanographaceae* Ertz, Tehler, G. Thor & Frisch, *Opegraphaceae* Körb. ex Stizenb., *Roccellaceae* Chevall., *Roccellographaceae* Ertz & Tehler), however, 21 genera are considered *incertae sedis* and perhaps more families can be established when more species are included in the phylogeny of the class (Wijayawardene et al. [2022a](#page-160-1)).

**Type order:** *Arthoniales* Henssen ex D. Hawksw. & O.E. Erikss., Syst. Ascom. 5(1): 177 (1986).

**Type family:** *Arthoniaceae* Reichenb. ex Reichenb., Deut. Bot. Herb.-Buch: 13 (1841).

**Type genus:** *Arthonia* Ach., Neues J. Bot. 1(3. Stück): 3 (1806).

**Type species:** *Arthonia radiata* (Pers.) Ach., K. Vetensk-Acad. Nya Handl. 29: 131 (1808).

*Dothideomycetes* O.E. Erikss. & Winka, Myconet 1(1): 5 (1997).

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**Sexual morph:** *Ascomata* stromatic, uni- or multi-loculate, pseudothecial, thyriothecial, cleistothecial, hysterothecial or apothecial, solitary to gregarious, globose to subglobose, pyriform to depressed, crustose or stellate, discoid to cupulate, immersed to superficial, coriaceous, with or without ostioles, glabrous to setose, periphysate, brown to black. *Peridium* single or multi-layered, composed of angular to globose or fliform, black to hyaline cells towards the inside lighter. *Hamathecium* comprising or lacking pseudoparaphyses. *Asci* cylindrical to clavate or ellipsoidal to subglobose, bitunicate, fssitunicate, with or without an ocular chamber. *Ascospores* fliform or ellipsoidal to fusiform, aseptate or septate, hyaline or colored, smooth- or rough-walled, with or without guttules, thin- or thick-walled, with or without various types of appendages and/or sheaths. **Asexual morph:** coelomycetous or hyphomycetous. **Coelomycetes:** *Conidiomata* acervular or pycnidial, solitary or gregarious, immersed to erumpent, unior multi-loculate, with or without an ostiole. *Conidiomatal wall* composed of angular to globose or fliform, hyaline to brown cells. *Conidiophores* cylindrical to ampulliform or fliform, septate, branched to unbranched, sometimes reduced to conidiogenous cells. *Conidiogenous cells* holoblastic, enteroblastic, phialidic, annellidic, arthric or tretic, integrated to discrete, doliiform, lageniform or cylindrical, sometimes with sympodial or percurrent proliferations, hyaline to brown. *Conidia* globose to oblong, cylindrical to fliform, hyaline to dark brown, aseptate or septate, smooth, striate, spinulose or verruculose, thin- or thick-walled, with or without appendages or gelatinous sheaths. **Hyphomycetes:** *Conidiomata* sporodochial or synnematal, solitary to gregarious, with or without setae. *Conidiophores* macronematous to micronematous, mononematous or synnematous, straight or fexuous, aseptate or septate, unbranched or branched, hyaline to brown. *Conidiogenous cells* blastic, phialidic, annellidic or tretic, integrated or discrete, determinate or proliferating sympodially or percurrently, cylindrical to oblong. *Conidia* solitary or catenate, acrogenous, pleurogenous or acropleurogenous, globose to ellipsoidal, cylindrical, clavate to obclavate, fusiform, fliform or reniform to sickle-shaped, straight to curved or helicoid, rounded or obtuse to truncate at apex, truncate at base, aseptate or septate, hyaline to dark brown, smooth or verruculose.

**Life modes and distribution:** The members of *Dothideomycetes* are distributed in a wide range of ecosystems around the world and have diverse life modes. Saprobic fungi are found in all environments and are important for decomposition and nutrient cycles in ecosystems (e.g. *Hysterium*, *Lindgomyces*). Phytopathogenic fungi are responsible for causing leaf, stem, root, and fruit diseases (e.g. *Alternaria*, *Botryosphaeria*, *Cercospora*, *Mycosphaerella*), while some have been reported as pathogenic to arthropods and mammals (e.g. *Hortaea*, *Medicopsis*, *Myriangium*, *Piedraia*). Lichens (e.g. *Monoblastia*, *Strigula*, *Trypethelium*) are found in the tropics and subtropics (Hongsanan et al. [2020b](#page-140-3)). Lichenicolous fungi (e.g. the *Phoma* complex) are also reported (Lawrey

<span id="page-20-0"></span>**Fig. 5** Morphology in species of *Dothideomycetes*. **a–d** Symptoms (**a** *Pseudocercospora cladrastidis* on leaves of *Maackia amurensis*. **b** *Alternaria alternata* on apple leaf. **c** *Venturia inaequalis* on apple fruit **d** *Shiraia bambusicola* on bamboo). **e–g** Ascomata on host surface (**e** *Dothidea puccinioides*. **f** *Caryospora minima*. **g** *Hysterium angustatum*). **h–j** Ascomata in vertical sections (**h** *D. puccinioides*. **i** *Keissleriella quadriseptata*. **j** *Minutisphaera japonica*). **k** pseudoparaphyses (*Lineolata rhizophorae*). **l** Ascus apex (*Jahnula* sp.). **m–p** Asci (**m** *Asterina aucubae*, **n** *Hadrospora fallax*. **o** *Massarina eburnea.* **p** *Neoheleiosa* sp.). **q** Fissitunicate ascus with extending endoascus (*Lindgomyces cinctosporus*). **r** Fissitunicate ascus and discharged ascospores with an elongated sheath (*Tingoldiago graminicola* (mounted in India ink)). **s–v** Ascospores (**s** *Phyllosticta* sp. (stained with blue-black ink). **t** *Rebentischia abietis*. **u** *Patellaria* sp. **v** *Stemphylium* sp.). **w** Conidioma in vertical section (*Versicolorisporium triseptatum*). **x** Macronematous conidiophores (*Cladosporium* sp.). **y** Synnema (*Phragmocephala atra*). **z-f**<sub>1</sub> conidia (**z** *Botryosphaeria dothidea*. **a**<sub>1</sub> *Periconia macrospinosa* (bleached using NaClO). **b**<sub>1</sub> *Muyocopron* sp. **c1** *Berkleasmium concinnum*. **d1** *Prosthemium intermedium*. **e1** *Quadricrura meridionalis*. **f1** *Tubeufa* sp.). **g1** Propagule (*Collapsimycopappus styracis*). Scale bars: **e**–**g**=500 µm, **h**–**j**,  $\mathbf{w} - \mathbf{y} = 100 \, \mu \text{m}$ ,  $\mathbf{m} - \mathbf{r} = 50 \, \mu \text{m}$ ,  $\mathbf{k}$ , **l**, **s**–**v**, **z**–**g**<sub>1</sub> $=$ 20  $\mu$ m



et al. [2012;](#page-144-5) Valenzuela-Lopez et al. [2018\)](#page-158-4). Some members are endophytic (e.g. *Alfoldia*, *Aquilomyces*, *Camarosporidiella*, *Neohendersonia*, *Ramichloridium*), mycorrhizal (e.g. *Cenococcum*), fungicolous (e.g. *Ampelomyces*, *Trichothyrium*) and coprophilous (e.g. *Delitschia*, *Preussia*). There are some rock-inhabiting fungi (e.g. *Coniosporium*, *Cryomyces*) and their species diversity is becoming clearer. The ancestors of *Dothideomycetes* are presumed to have been saprobic and then evolved into other lifemodes such as phytopathogens and lichens (Schoch et al. [2009a;](#page-153-0) Haridas et al. [2020\)](#page-138-12).

**Notes:** *Dothideomycetes*, the largest and most phylogenetically diverse class of *Ascomycota*, were introduced by Eriksson and Winka [\(1997](#page-136-2)). This class encompasses the majority

<span id="page-21-1"></span><span id="page-21-0"></span>

<span id="page-21-2"></span>of species previously classifed under *Loculoascomycetes* (Luttrell [1955\)](#page-146-9) and is characterized by ascolocular development and fssitunicate asci. The classifcation of sexually typifed genera within *Dothideomycetes* was provided by Lumbsch and Huhndorf [\(2007](#page-146-1), [2010](#page-146-2)) alongside the Phylum *Ascomycota*. Similarly, asexual-typifed genera in this class were incorporated into the natural classifcation systems by Wijayawardene et al. ([2014](#page-159-10)), where molecular data are available. The most recent monographs of families of *Dothideomycetes* were presented by Hongsanan et al. ([2020a](#page-140-2), [b](#page-140-3)).

*Dothideomycetes* comprising 49 orders, 222 families and 1748 genera (Wijayawardene et al. [2022a](#page-160-1)) are divided into two subclasses: *Dothideomycetidae* and *Pleosporomycetidae*. Most orders and families are treated as *incertae sedis* within this class (Schoch et al. [2006\)](#page-153-1). Within the subclass *Dothideomycetidae*, there are ten accepted orders, namely: *Aureoconidiellales*, *Capnodiales*, *Cladosporiales*, *Comminutisporales*, *Dothideales*, *Mycosphaerellales*, *Myriangiales*, *Neophaeothecales*, *Phaeothecales*, and *Racodiales* (Wijayawardene et al. [2022a](#page-160-1)). In the *Pleosporomycetidae* subclass, *Gloniales*, *Hysteriales*, *Mytilinidiales*, and *Pleosporales* are accepted (Wijayawardene et al. [2022a](#page-160-1)), whereas 35 orders are categorized as *Dothideomycetes* orders *incertae sedis*.

One fungus one name: In recent years, DNA sequencing and molecular phylogenetic analysis have made it possible to confrm sexual and asexual relationships and to propose one name for a single species or genus (Crous et al. [2009](#page-134-3); Bensch et al. [2012;](#page-131-4) Manamgoda et al. [2012](#page-146-10); Phillips et al. [2013;](#page-150-6) Hyde et al. [2013,](#page-140-9) [2014](#page-140-11)). Thus, it was proposed to adopt one name (between two generic names) when the links between the morphs are confrmed.

The members of *Dothideomycetes* are well-known as pathogens of both plants and animals, and industrially important taxa. Some studies focusing on economically, clinically and industrially important taxa are summarized in Tables [8](#page-21-0), [9](#page-21-1) and [10](#page-21-2) respectively (examples that were published in the last 5 years).

**Type order:** *Dothideales* Lindau, in Engler & Prantl, Nat. Pfanzenfam., Teil. I (Leipzig) 1(1): 373 (1897).

**Type family:** *Dothideaceae* Chevall. [as 'Dothideae'], Fl. gén. env. Paris (Paris) 1: 446 (1826).

**Type genus:** *Dothidea* Fr., Observ. mycol. (Havniae) 2: 347 (1818).

**Type species:** *Dothidea sambuci* (Pers.) Fr., Syst. mycol. (Lundae) 2(2): 551 (1823).

*Eurotiomycetes* O.E. Erikss. & Winka, Myconet 1(1): 6 (1997).

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**Thallus:** Yeast-like to flamentous. **Sexual morph**: *Stromata* occasionally present, reduced, usually pulvinate or (sub)globose, sometimes stipitate, white, brown or black. *Ascomata* apothecial, ascostromatal, cleistothecial, gymnothecial, perithecioid, or mazaediate, mostly (sub)globose, pyriform or variously shaped, sometimes setose or with dictinct peridial appendages, light-coloured to black, solitary to aggregated. *Hamathecium* absent or including paraphyses, pseudoparaphyses, periphysoids and periphyses in diferent combinations. *Asci* usually 8-spored, sometimes 2-spored or polysporous, mostly (sub)globose, clavate or obpyriform, unitunicate or bitunicate, with fssitunicate to evanescent dehiscence. Ascospores usually allantoid, bivalved, ellipsoidal, globose, oblate, (ob)ovoid, reniform or star-shaped, aseptate, transversely septate or dictyoseptate, hyaline, lightcoloured or brown, smooth or showing an equatorial furrow, rim or crests, a reticulum, ridges, spines, warts or other ornamentations, sometimes surrounded by a sheath. **Asexual morph**: coelomycetous or hyphomycetous. **Coelomycetes:** *Conidiomata* pycnidial or pycnothyrial, with or without a central ostiole, commonly fat to globose, brown, immersed to erumpent. *Conidiomatal wall* often of *textura angularis* or *textura epidermoidea* at the outer portion, inner wall covered with hyaline to dark brown, ampulliform, cylindrical, ellipsoidal, fusiform, oval or lens-shaped phialidic conidiogenous cells, or with pale brown toruloid conidiophores producing cheiroid conidia. *Conidia* aseptate, mostly cylindrical to ellipsoid, hyaline or cheiroid, consisting of a central globose, medium brown, verruculose cell giving rise to up to 6 radiating arms of subglobose to ellipsoidal cells, each forming lateral branches. **Hyphomycetes:** *Conidiomata* present or absent, scattered or aggregated, sporodochia or synnemata. *Setae* sometimes present. *Conidiophores* micronematous to macronematous, hyaline to brown, solitary to fasciculate, aseptate or septate, simple or branched, smooth-walled to verrucose, *Conidial ontogeny* enteroblastic, holoblastic, holothallic or thallic-arthric. *Conidiogenous cells* intercalary, lateral or terminal, sometimes extending percurrently or sympodially, sometimes denticulate or with dark or refractive scars. *Conidia* solitary or produced in mucilaginous masses or chains, aseptate, transversely septate or dictyoseptate, highly diverse in shape, usually (sub)cylindrical, (ob) clavate, doliiform, ellipsoid, (sub)globose, falcate, fusiform or guttuliform, smooth-walled or ornamented, usually with more or less developed spines or warts, with or without a basal frill, secession rhexolytic or schizolytic.

**Life modes and distribution:** Members of *Eurotiomycetes* are abundant both in terrestrial and aquatic ecosystems worldwide, occurring in an extremely broad range of habitats. While most of these fungi are degraders of decaying material of plant or animal origin, many species cause diseases in animals (e.g. *Exophiala cancerae*, an agent of Lethargic Crab Disease in the mangrove crab *Ucides cordatus fde* de Hoog et al. [2011](#page-135-5)) (Table [11](#page-25-0)), plants (e.g. *Dolabra nepheliae* on rambutan and lychee *fde* Rossman et al. [2010\)](#page-152-10), or other fungi (e.g. *Cladophialophora pucciniophila* on *Puccinia polygoni-amphibii fde* Park and Shin [2011\)](#page-149-4). This order includes a remarkably high number of clinically relevant species causing diverse pathologies in humans and other animals, including agents of life-threatening systemic mycoses, such as blastomycosis, coccidioidomycosis, histoplasmosis, and paracoccidioidomycosis, as well as fungi causing chromoblastomycosis, dermatophytosis, hyalohyphomycosis, mycetoma, and phaeohyphomycosis (de Hoog et al. [2020](#page-135-6)). Certain lineages in *Eurotiomycetes* contain lichenized fungi, e.g. *Pyrenulales*, *Verrucariales*, and some *Celotheliaceae* (Gueidan et al. [2014](#page-138-2); Pykälä et al. [2020](#page-150-7); Miranda-González et al. [2022\)](#page-147-4) or lichenicolous species (e.g. *Knufa peltigerae* and *Sclerococcum* spp. *fde* Untereiner et al. [2011;](#page-157-5) Diederich et al. [2018\)](#page-135-7).

**Notes**: The class *Eurotiomycetes* was erected by Eriksson and Winka ([1997](#page-136-2)) based on phylogenetic analysis of the 18S rRNA gene (SSU). The original circumscription of this class included most fungi referred to as "Plectomycetes", which were roughly characterized by thin-walled prototunicate asci (unitunicate asci with walls which disintegrate at maturity to release the ascospores passively within the ascoma), ascomata lacking a distinct hymenial layer and with asci scattered within the ascomatal cavity, aseptate ascospores, ascomata gymnothecial to cleistothecial, the latter sometimes produced within a stroma, and morphologically diverse blastic or thallic asexual morphs (Alexopoulos et al. [1996](#page-130-6); Geiser et al. [2006\)](#page-137-9). Eriksson and Winka ([1997\)](#page-136-2) included two orders in this class, i.e. *Eurotiales* and *Onygenales*. The former included the families *Elaphomycetaceae*, *Monascaceae* and *Trichocomaceae*, and the latter included *Arthrodermataceae*, *Ascosphaeraceae*, *Eremascaceae*, *Gymnoascaceae* and *Onygenaceae*. The monophyly of this class was confrmed subsequently in a SSU-based phylogenetic analysis by Geiser and LoBuglio [\(2001\)](#page-137-10). Unexpectedly, early phylogenetic studies (Spatafora et al. [1995](#page-155-5); Berbee [1996](#page-131-5); Silva-Hanlin and Hanlin [1999\)](#page-155-6) revealed that the "monophyletic Plectomycetes" shared a common ancestry with members of the order *Chaetothyriales*, which produce perithecioid ascomata and bitunicate asci with an active, fssitunicate spore release mechanism (Müller et al. [1987](#page-148-9); Barr [1991\)](#page-131-6). The order *Coryneliales* has been considered a "missing link" between the "monophyletic Plectomycetes" and chaetothyrialean genera and related fungi. *Coryneliales* produce ascolocular ascomata and asci which are initially bitunicate, but whose outer layer breaks away early in their development, so that asci appear unitunicate at maturity, releasing spores passively by degradation of the inner ascal wall (Johnston and Minter [1989\)](#page-141-8). Multi-locus phylogenetic studies placed this order basal to the clade comprising *Eurotiales* and *Onygenales* (Geiser et al. [2006\)](#page-137-9), supporting the view that the ascomata of Plectomycetes evolved by loss of the bitunicate ascus and its active spore release mode (Eriksson [1982\)](#page-136-11). The boundaries of *Eurotiomycetes* have been broadened signifcantly in recent years and, at present, it includes six ecologically, morphologically, and phylogenetically diverse subclasses, i.e. *Chaetothyriomycetidae*, *Coryneliomycetidae*, *Cryptocaliciomycetidae*, *Eurotiomycetidae*, *Mycocaliciomycetidae*, and *Sclerococcomycetidae*



(Prieto et al. [2021](#page-150-8)). The class *Eurotiomycetes* currently encompasses twelve orders, 34 families and ca. 291 genera.

Numerous species of *Eurotiomycetes* are pleomorphic, producing sexual and asexual morphs which were traditionally named separately. Recent phylogenetic studies (Réblová et al. [2013;](#page-151-5) Samson et al. [2014](#page-152-11); Yilmaz et al. [2014](#page-161-7); Diederich et al. [2018;](#page-135-7) Sun et al. [2019a](#page-156-9), [b](#page-156-10); Houbraken et al. [2020](#page-140-12);

Kandemir et al. [2022](#page-142-11)) have applied unitary nomenclature in diferent genera in this class, in the pursuit of taxonomic stability.

**Type order:** *Eurotiales* G.W. Martin ex Benny & Kimbr., Mycotaxon 12(1): 23 (1980).

<span id="page-24-0"></span>**Fig. 6** Macro and micro morphological characteristics of eurotiomycetous taxa. **a**, **g**–**k** ascomata (**a** *Agonimia tristicula* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*; **g** *Pyrgillus americanus* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*); **h** *Chaetothyrium guaraniticum* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*); **i**: *Dolabra nepheliae* (*Celotheliaceae*, *Phaeomoniellales*, *Chaetothyriomycetidae*); **j** *Coccodinium bartschii* (*Coccodiniaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*); **k** *Pyrenowilmsia ferruginosa* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **b** Synnema of *Penicilliopsis clavariiformis* (*Aspergillaceae*, *Eurotiales*, *Eurotiomycetidae*). **c** Ascostroma of *Corynelia tripos* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **d** Stipewith volva of *Pseudotulostoma volvatum* (*Elaphomycetaceae*, *Eurotiales*, *Eurotiomycetidae*). **e** Apothecia of *Mycocalicium parietinum* (*Mycocaliciaceae*, *Mycocaliciales*, *Mycocaliciomycetidae*). **f** Stromata *Dendrosphaera eberhardtii* (*Aspergillaceae*, *Eurotiales*, *Eurotiomycetidae*). **h** Ascomata with black setae of *Chaetothyrium guaraniticum* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). l Squash mount of ascoma of *Actinocymbe separatosetosa* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). **m**, **n**, **p** Vertical section of ascoma (**m** *Dermatocarpon miniatum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*); **n** *Dermatocarpella yoshimurae* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **o** Vertical section of apothecia of *Phaeocalicium praecedens* (*Mycocaliciaceae*, *Mycocaliciales*, *Mycocaliciomycetidae*). **q**–**ai** Asci with ascospores. (**q** *Phaeosaccardinula diospyricola* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). **r** *Samarospora potamogetonis* (*Eurotiomycetidae incertae sedis*). **s** *Dermatocarpon miniatum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **t** *Corynelia clavata* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **u** *Caliciopsis pinea* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **v** *Fitzpatrickella operculata* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **w** *Sporodictyon schaererianum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **x** *Actinocymbe separato-setosa* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). **y** *Muellerella polyspora* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **z** *Clypeopyrenis microsperma* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **ab** *Melanotheca coccorum* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **ac** *Pyrenowilmsia ferruginosa* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **ad** *Heuferidium pentagastricum* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **ae** *Dolabra nepheliae* (*Celotheliaceae*, *Phaeomoniellales*, *Chaetothyriomycetidae*). **af** *Parathelium polysemum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **ag** *Phaeocalicium praecedens* (*Mycocaliciaceae*, *Mycocaliciales*, *Mycocaliciomycetidae*). **ah** *Chaenothecopsis rubescens* (*Mycocaliciaceae*, *Mycocaliciales*, *Mycocaliciomycetidae*). **ai** *Moriola descensa* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **aj-at** Ascospores. (**aj** *Verrucaria tartarina* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **ak** *Rhaphidicyrtis trichosporella* (*Incertae sedis*, Pyrenulales, *Chaetothyriomycetidae*). **al** *Tripospora tripos* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **am** *Samarospora potamogetonis* (*Eurotiomycetidae* genera *incertae sedis*). **an** *Lagenulopsis bispora* (*Coryneliaceae*, *Coryneliales*, *Coryneliomycetidae*). **ao** *Heterocarpon ochroleucum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **ap** *Parapyrenis aurora* (*Pyrenulaceae*, *Pyrenulales*, *Chaetothyriomycetidae*). **aq** *Phaeospora rimosicola* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **ar** *Ceratocarpia cactorum* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). **as** *Bicornispora exophiala* (*Rutstroemiaceae*, *Helotiales*, *Leotiomycetidae*). **at** *Sporodictyon schaererianum* (*Verrucariaceae*, *Verrucariales*, *Chaetothyriomycetidae*). **au**, **av** Spores (**au** *Coccodinium bartschii* (*Coccodiniaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). **av** *Chaetothyrium spinigerum* (*Chaetothyriaceae*, *Chaetothyriales*, *Chaetothyriomycetidae*). Scale bars: **a**, **b**=2000 µm, **c**, **f**=1000 µm, **e**, **h**, **i**, **k–o**=100 µm, **g**=500 µm, **p**=50 µm, **q**, **w**, **ad**, **ai**, **av**=20 µm, **r**–**v**, **x**–**ac**, **ae**–**ah**, **ak**, **al**,  $\text{as}-\text{au}=10 \, \mu \text{m}$ ,  $\text{aj}, \text{am}-\text{ar}=5 \, \mu \text{m}$  (photo credit to Qing Tian)

◂ **Type family:** *Aspergillaceae* Link, Abh. dt. Akad. Wiss. Berlin 1824: 165. 1826.

=*Eurotiaceae* Clem. & Shear, Gen. fung., Edn 2 (Minneapolis): 50 (1931).

**Type genus:** *Aspergillus* P. Micheli ex Haller, Hist. stirp. Helv. (Bernae) 3: 113 (1768).

**Type species:** *Aspergillus glaucus* (L.) Link 1809 (Designated by Clements & Shear, Gen. fung., Edn 2 (Minneapolis): 386. 1931).

=*Eurotium herbariorum* (F.H. Wigg.) Link ex Nees, Syst. Pilze (Würzburg): 96 (1816) [1816–17].

*Laboulbeniomycetes* Engler & Prantl, Nat. Pfanzenfam*.*, Teil. I (Leipzig) 1: vi (1897).

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*Laboulbeniales* **and** *Herpomycetales*: No production of hyphae; three-dimensional thallus solitary or multiple, determinate growth, formed by divisions of two-celled ascospore, composed of up to several thousand cells; perithecia composed of two-layered walls, systematically to rarely haustorial (rhizoid-like), haustoria simple or branched, single or multiple per thallus, no asexual reproduction; host specifc, habitat specific, and position specific, on single arthropodonly hosts, ascospores transmitted primarily through direct contact between hosts. *Laboulbeniales***:** Thalli averaging 200–300 µm, microscopic and multicellular, perithecial wall generally arises from perithecial basal cells, with inner cell wall starting at diferent level outer cell walls, rows of outer wall cells growing upwards around and after carpogonial extension, melanin-rich cell walls, unculturable (or if growth, no more than a few cells), rarely haustorial (rhizoid-like), single per thallus; from four to eight two-celled ascospores per ascus, septum positioned near the lower end separating a smaller (directed downward) and larger cell (directed upwards), asci generated from a single series of ascogenic cells, sticky ascospores transmitted directly from infected to uninfected hosts during mating or other contacts; known from all continents (except Antarctica), wide variety of hosts including cockroaches, termites, beetles, fies, true bugs, ants, crickets and allies, lice, and thrips, moderately to highly host-specifc. *Herpomycetales***:** Dioecious; four-celled primary axis of thallus developing directly from ascospore; suprabasal cell in female thallus giving rise to secondary axis (or axes), producing perithecia and connecting directly with integument of the host; perithecia multi-tiered, outer wall rows consisting of many cells equal in height, develop before carpogonial upgrowth which extends between the outer wall cells, bearing single to three apex projections; haustorial, multiple per thallus; ascospores 8 per ascus with median septum resulting in two equal cells; asci produced sequentially from two series of ascogenic

Species name	Human/ animal	References
Aspergillus fumigatus	Main agent of aspergillosis worldwide in human and various warm-blooded animals	Tsang et al. $(2018)$ , Houbraken et al. $(2020)$ , Ortiz et al. (2022)
<b>Blastomyces dermatitidis</b>	Causative agent of blastomycosis in humans and other mammals	Dukik et al. $(2017)$ , Jiang et al. $(2018)$ , Linder & Kauff- man $(2020)$
Cladophialophora carrionii	Main agent of human chromoblastomycosis in arid regions	Deng et al. (2015), de Hoog et al. (2020)
Cladophialophora bantiana	Agent of phaeohyphomycosis in humans and other mammals, usually affecting the brain	Kantarcioglu et al. $(2017)$ , de Hoog et al. $(2020)$
Coccidioides immitis	Agent of human coccidioidomycosis	de Hoog et al. $(2020)$ , Kandemir et al. $(2022)$ , Kirkland et al. $(2022)$
Exophiala dermatitidis	Agent of various forms of phaeohyphomycosis in humans	Song et al. (2017), Kirchhoff et al. (2019)
Fonsecaea pedrosoi	Main agent of human chromoblastomycosis in humid regions	Schneider et al. (2019)
Histoplasma capsulatum	Agent of human histoplasmosis	Sepúlveda et al. (2017), Valdez et al. (2022)
Paracoccidioides brasiliensis	Agent of paracoccidioidomycosis in humans and other mammals	Pinheiro et al. $(2020)$ , Hahn et al. $(2022)$ , Rodrigues et al. $(2023)$ , Vilela et al. $(2023)$
Talaromyces marneffei	Agent of talaromycosis in humans and other mammals	Cao et al. $(2019)$ , Pruksaphon et al. $(2022)$ , Wang et al. (2023a)
Trichophyton rubrum	Human, agent of dermatophytosis	de Hoog et al. $(2017)$ , Chanyachailert et al. $(2023)$

<span id="page-25-0"></span>**Table 11** Medicinally/clinical signifcance, examples:

cells, host-specificity restricted to cockroaches. *Pyxidiophorales***:** Three-morph life-cycle with hyphal growth, including an independent sexual (meiosporic origin) state and two asexual (mitosporic origin) states associated with arthropods for dispersal, some mycoparasitic forming twocelled ascospores dividing into asexual state (*Thaxteriola* Speg.); perithecia produced from a mycelium, composed of single-layered cell walls, reduced number of ascospores per ascus; heteroecious, with fungal host (which is parasitized) and arthropod host (for dispersal), sometimes associated on ephemeral substrates such as herbivore dung.

**Life modes and distribution:** Perithecial fungi obligatorily associated with arthropod integuments, either as ectobionts or for dispersal, on a wide range of terrestrial and aquatic hosts; forming teleomorphic thalli on specifc or unspecifc positions on the host integument, sometimes involved in hyperparasitic associations; or mycoparasitic and producing anamorphic morphs dispersed by arthropods (Fig. [8](#page-27-0)). **Notes**: *Laboulbeniomycetes* is a class of microscopic fungi within *Ascomycota* exhibiting an obligate association with arthropods. Three named (*Laboulbeniales*, *Herpomycetales* and *Pyxidiophorales*) and two unnamed (*Chantransiopsis* and *Laboulbeniopsis*) clades are currently accepted (Haelewaters et al. [2021a\)](#page-138-6). However, the circumscription of these clades has been impeded by numerous taxonomic and phylogenetic misunderstandings and misinterpretations, resulting in a *turbulent* taxonomic history (*fde* Haelewaters et al. [2021b](#page-138-13)). Initially mistaken for parasitic worms or red algal relatives, there was no consensus among biologists and mycologists regarding the fungal nature of these organisms during the twentieth century, mainly due to the lack of "typical" fungal characters. Charles P. Robin (1821–1885) was the frst one to recognize *Laboulbeniomycetes* as fungi, alongside Roland Thaxter (1891–1931) who later considered them to be ascomycetes (Blackwell et al. [2020](#page-132-2)). Thaxter's eforts led to the description of over half of the currently known diversity of *Laboulbeniomycetes*, inevitably based on morphological characters (e.g. perithecium morphology or asci development).

Since Engler ([1898](#page-136-12)) erected the class *Laboulbeniomycetes*, the accumulation of empirical data and the development of molecular-based tools considerably improved our understanding of life history traits and phylogenetic relationships within this group. The class was for the frst time confrmed by molecular data by Weir and Blackwell [\(2001](#page-159-13)) and while molecular-based studies show clear advantages over feld-based observations (Cazabonne et al. [2022\)](#page-133-8), numerous phylogenetic relationships within *Laboulbeniomycetes* remain to be elucidated. Goldmann and Weir ([2018](#page-137-5)) demonstrated that *Laboulbeniomycetes* is a monophyletic group within *Ascomycota*, closely related to *Sordariomycetes*, confrming the previous placement of the *Laboulbeniales*-*Pyxidiophora* clade (Weir and Blackwell [2001](#page-159-13)). The uncertain placement of the genus *Herpomyces* was resolved by Haelewaters et al. [\(2019](#page-138-4)) by the introduction of the order *Herpomycetales* based on molecular phylogenetic, morphological, developmental, and host usage data. Despite those recently undertaken efforts, deeper nodes remain unresolved and the sister group of *Laboulbeniomycetes* is yet to be discovered

<span id="page-26-0"></span>**Fig. 7** *Laboulbeniomycetes*, including (**a**–**l**) *Laboulbeniales*, (**m**–**n**) *Herpomycetales*, and (**o**–**p**) *Pyxidiophorales*. **a** *Stigmatomyces majewskii*. **b** *Teratomyces actobii*. **c** *Rhachomyces anophthalmi*. **d** *Laboulbenia notiophili*. **e** *Dimeromyces rugosus*. **f** Laboulbenia anisodactyli. **g** *Arthrorhynchus eucampsipodae*. **h** *Gloeandromuces cusucoensis*. **i** *Laboulbenia oioveliicola*. **j** *Haplomyces texanus*. **k** *Hesperomyces harmoniae*. **l** *Gloeandromyces plesiosaurus* f. *plesiosaurus*. **m** *Herpomyces stylopygae*. **n** *Herpomyces shelfordellae*. **o** *Thaxteriola* sp. **p** *Pyxidiophora bialowiezensis*. Scale bars: **a**, **c**, **d**, **h**, **l**–**o**=50 µm; **b**, **e**, **g**, **j**, **k**,  $p = 100 \text{ µm}$ ;  $f = 10 \text{ µm}$ ; **i**=20 µm. Photos: Danny Haelewaters; drawings: Jingyu Liu



(Blackwell et al. [2020\)](#page-132-2). Fossil records for *Laboulbeniomycetes* are limited to *Columnomyces electri* from Dominican amber (Perreau et al. [2021](#page-151-6)) and *Stigmatomyces succini* from Bitterfeld amber (Rossi et al. [2005\)](#page-152-13).

**Type order:** *Laboulbeniales* Lindau, in Engler & Prantl, Nat. Pfanzenfam., Teil. I (Leipzig) 1(1): 47 (1897). **Type family:** *Laboulbeniaceae* G. Winter [as 'Laboulbenieae'], Rabenh. Krypt.-Fl., Edn 2 (Leipzig) 1.2: 918 (1886). **Type genus:** *Laboulbenia* Mont. & C.P. Robin, in Robin, Histoire naturelle des Vegetaux Parasites (Paris): 622 (1853). **Type species:** *Laboulbenia rougetii* Mont. & C.P. Robin, in Robin, Histoire naturelle des Vegetaux Parasites (Paris): 622 (1853).

*Lecanoromycetes* O.E. Erikss. & Winka, Myconet 1(1): 7 (1997).

Index Fungorum Registration Identifer 501486; Fig. [9](#page-28-0)



<span id="page-27-0"></span>**Fig. 8** Hosts of *Laboulbeniomycetes* microfungi. **a** *Harmonia quadripunctata* (Coleoptera, Coccinellidae) with *Hesperomyces harmoniae*. **b** *Bembidion femoratum* (Coleoptera, Carabidae) with *Laboulbenia vulgaris*. **c** *Bradycellus harpalinus* (Coleoptera, Carabidae) with *Laboulbenia eubradycelli*. **d** A dried, pinned specimen of *Crepi-*

*dogaster bioculata* (Coleoptera, Carabidae) with *Laboulbenia* sp. **e** *Myrmica sabuleti* (Hymenoptera, Formicidae) with *Rickia wasmannii*. **f** *Penicillidia conspicua* (Diptera, Nycteribiidae) with *Arthrorhynchus nycteribiae*. Photos: a: Gilles San Martin; b, c: Bart Horvers; d: Ondřej Koukol; f: Walter P. Pfiegler; E. Theodoor Heijerman

Sexual morph: Thallus usually superficial on the substratum, more rarely immersed, poikilohydric, usually longlived (on average perennial, up to thousands of years), composed of hyphae and green algae and/or cyanobacteria in defned tissues. *Ascomata* apothecia, often present but unknown in many species, solitary or aggregated, round

to linear, variously coloured, disc usually without algae or cyanobacteria, margin often containing algae or cyanobacteria. *Hamathecium* composed of paraphyses. *Asci* often eight-spored, but varying between 1-spored and multi (>100)-spored, cylindrical to fusiform or clavate, incompletely bitunicate, often with J+staining tholus. *Ascospores*



**Fig. 9** *Lecanora saepiphila.* **a** habitat on wood. **b** section through apothecium, mounted in water. **c** section through apothecium, mounted in 10% KOH. Pictures by L.A. dos Santos

<span id="page-28-0"></span>flamentous, globose, ellipsoidal or fusiform, aseptate or septate, hyaline or brown, often surrounded by a gelatinous sheath. **Asexual morph:** coelomycetous. *Conidiomata* frequently present, pycnidial, developing on the same thalli as the apothecia, usually immersed, globose, coriaceous, with a single locule, dark brown to black, ostiolate. *Conidiomata wall* carbonized. *Conidiophores* lining the inner cavity or reduced, septate, sometimes branched. *Conidiogenous cells* phialidic, discrete, lageniform, hyaline to pale yellow, smooth, producing microconidia and sometimes mesooconidia. *Conidia* hyaline, aseptate to multiseptate, without appendages.

**Life modes and distribution:** Lichen-forming with green algae and/or cyanobacteria, or more rarely parasitic on lichens or saprobic; on bark, wood, rock, soil, detritus, other lichens and/or living leaves. Showing a worldwide distribution, but most abundant in boreal to arctic regions.

**Notes:** The class *Lecanoromycetes*, one of the largest classes of phylum *Ascomycota*, was introduced by Eriksson and Winka [\(1997\)](#page-136-2). It comprises the vast majority of the lichenized fungi. Apart from this, it accommodates some species that are parasitic on lichens, either with their own separate algae, or without separate algae and using the algae of the host; there are also some groups with a saprobic life style. Rambold and Triebel ([1992\)](#page-151-7) provided a discussion and list of such species.

Based on Lücking et al. ([2017](#page-146-4)) and Wijayawardene et al. ([2022a](#page-160-1)), the class comprises 21 orders, 82 families, around 775 genera and around 15,500 species. The saprobic life style is largely restricted to a small part of the *Ostropales*.

The members of the *Lecanoromycetes* are distributed over a broad range of ecosystems. They are most conspicuous in boreal, (ant) arctic and alpine ecosystems, where they cover vast areas of soil and rock. In other ecosystems they are present on tree bark and living leaves. As with almost every group, they are most diverse in the tropics. They are themselves often host to other, so-called intralichenic, flamentous fungi and/or yeasts (Spribille et al. [2016](#page-155-8); Grimm et al. [2021](#page-137-11)).

The main use of *Lecanoromycetes* is for the biomonitoring of air pollution (Giordani [2019](#page-137-12)). Many species are sensitive to sulphur dioxide, while other species respond positively to ammonia pollution.



**Type order:** *Lecanorales* Nannf., Nova Acta R. Soc. Scient. upsal., Ser. 4 8(no. 2): 68 (1932).

**Type family:** *Lecanoraceae* Körb. [as 'Lecanoreae'], Syst. lich. germ. (Breslau): 104 (1855).

<span id="page-30-0"></span>**Fig. 10** *Leotiomycetes* fungi. **a** *Lachnellula willkommii*, rehydrated ◂ apothecia showing dense hairs on the outside, on branch of *Larix decidua*, Sampeyre, Italy (*Lachnaceae*, *Helotiales*). **b** *Phialocephala* sp., Calf Island, Boston Harbor Islands, Massachusetts USA (*Mollisiaceae*, *Helotiales*). **c** *Chlorosplenium cusucoense* (as *Chlorosplenium* sp. 1 in Haelewaters et al. 2021), Boquete, Panama (*Chlorospleniaceae*, *Helotiales*). **d** *Sclerencoelia fraxinicola*, erumpentapothecia on branch of *Fraxinus excelsior*, Tübingen, Germany (*Cenangiaceae*, *Helotiales*). **e** *Propolis farinosa*, erumpent apothecia on branch of Populus sp., Peddocks Island, Boston Harbor Islands, Massachusets, USA (*Marthamycetaceae*, *Rhytismatales*). **f** *Leotia* sp. nov., Koussoucoingou, Benin (*Leotiaceae*, *Leotiales*). **g** *Pezoloma* sp., apothecia on dead leaf of *Acer saccharinum*, New Durham, New Hampshire, USA (*Discinellaceae*, *Helotiales*). **h** *Hymenoscyphus* serotinus, on *Fagus sylvatica* twig, Kaiserslautern, Germany (*Helotiaceae*, *Helotiales*). **i** *Pezoloma* sp., transverse section of 2–4-mm wide apothecium (*Discinellaceae*, *Helotiales*). **j** *Hymenoscyphus* sp. nov., ascus p in Melzer's reagent, Caribou, Colorado. **k** *Ascocoryne* sp., a sci i n M elzer's reagent, World's End Peninsula, Boston Harbor Islands, Massachusets, USA (*Gelatinodiscaceae*, *Helotiales*). Scale bars: j, k  $= 10 \mu m$ . Photos: a, d. Hans-Oto Baral; b, c, k. Danny Haelewaters; e. Alden C. Dirks; f. Meike Piepenbring; g, i. Jason M . Karakehian; h. Petra Behrens; j. Candice C. Perrota

**Type genus**: *Lecanora* Ach., in Luyken, Tentam. Histor. Lichen. Gen. Prim. Lin. Distrib. nov., Göttingae: 90 (1809). **Type species**: *Lecanora argentata* (Ach.) Röhl., Deutschl. Fl. (Frankfurt) 3(2): 82 (1813).

*Leotiomycetes* O.E. Erikss. & Winka, Myconet 1(1): 7 (1997).

Index Fungorum Registration Identifer 501,487; Fig. [10](#page-30-0) *Ascomata* typically small (<2 cm) in size; often apothecial (cup-shaped), of various sizes and forms including cups to earth tongues to perithecioid apothecia, rarely naked asci developing from hyphae due to lack of ascomata (e.g. in *Thelebolales*, genus *Antarctomyces*); frequently open apothecium, closed until maturity (e.g., in *Lahmiales*, *Rhytismatales*) or permanently closed (cleistothecia) (e.g., in *Amorphothecaceae*, *Erysiphaceae*, *Myxotrichaceae*, *Pleuroascaceae*, *Rutstroemiaceae*, *Thelebolaceae*); single or multicelular ascomata when closed. Hymenium exposed or not during ascomata maturation. *asci* inoperculate, with apical ring amyloid, inamyloid, or hemiamyloid, frequently not surrounded by tissue (open apothecia); rarely splitting open at maturity. *Ascospores* frequently without ornamentation, actively (e.g. open apothecia) or passively discharged through wall disintegration (e.g. cleistothecia). *Paraphyses* produced in open apothecia and absent in cleistothecia; some lineages amenable to growth in axenic culture; common in environmental DNA samples; asexual reproduction frequent, certain taxa only known from asexual morphs; yeast forms rare.

**Life modes and distribution:** *Leotiomycetes* have a global distribution and are found on all continents, although historically they have been unevenly sampled (Quandt and Haele-waters [2021](#page-150-11)). There are numerous extremophiles, including psychrophiles (in *Thelebolales*; de Menezes et al. [2017](#page-135-10); Villanueva et al. [2021](#page-158-6)). Environmental DNA studies have shown that *Leotiomycetes* are dominant in many diferent habitats, including arctic tundras, southern temperate forests, and peat bogs (Tedersoo et al. [2014;](#page-157-8) Lamit et al. [2017](#page-144-6)). *Helotiales* is a speciose and a diverse order with numerous ecologies including saprotrophs, plant pathogens, mycorrhizae, and endophytes (Baral et al. [2015](#page-130-7); Ekanayaka et al. [2019;](#page-136-14) Johnston et al. [2019](#page-141-0)). *Chaetomellales*, *Lahmiales*, *Marthamycetales*, *Micraspidales*, and *Rhytismatales* are dominated by taxa growing within and fruiting on living or dead plant tissues (Quandt and Haelewaters [2021\)](#page-150-11). *Leotiales* is ecologically diverse, with saprotrophic, lichenized, plantpathogenic, and potentially mycorrhizal representatives. *Thelebolales* comprises saprotrophs, coprophiles, psychrophiles, and mammal associates including pathogens (Landvik et al. [1998;](#page-144-7) de Hoog et al. [2005;](#page-134-4) Minnis and Lindner [2013](#page-147-5); Haelewaters et al. [2021d\)](#page-138-15). *Phacidiales* includes saprotrophs and parasites of plants, fungi, and lichens (Quandt and Haelewaters [2021](#page-150-11)). *Lauriomycetales* and *Umbellidiales* are only known from asexual morphs found in leaf litter (Hernández-Restrepo et al. [2017](#page-139-10); Barreto et al. [2023\)](#page-131-7). *Medeolariales* only includes one described species that is a plant pathogen (LoBuglio and Pfister [2010\)](#page-145-10).

**Notes:** Eriksson and Winka [\(1997\)](#page-136-2) erected the class *Leotiomycetes* to classify ascomycetes with inoperculate asci, which include the morphological group of the inoperculate discomycetes. Eriksson ([2005](#page-136-15)) excluded *Geoglossaceae* and added *Pseudeurotiaceae* to the class. Later, Wang et al. ([2006\)](#page-159-15) recognized the following higher taxa: *Cyttariales*, *Erysiphales*, *Helotiales*, *Rhytismatales*, *Thelebolales*, *Myxotrichaceae*, and *Pseudeurotiaceae*. Rossman et al. ([2004](#page-152-14)) determined a close relationship between *Chaetomella* and *Pilidium* within *Leotiomycetes*. *Chaetomellaceae* was erected to accommodate the two genera *Chaetomella* and *Pilidium* (Baral et al. [2015](#page-130-7)) and later recognized as an ordinal-level clade by Crous et al. ([2017](#page-134-5)). Using genomic data, Johnston et al. ([2019](#page-141-0)) recognized *Chaetomellales*, *Helotiales*, *Leotiales*, *Marthamycetales*, *Phacidiales*, *Rhytismatales*, and *Thelebolales*. Recent ordinal-level additions include: *Lauriomycetales* (Hernández-Restrepo et al. [2017](#page-139-10); Haelewaters et al. [2021c](#page-138-7)), *Lichinodiales* (Prieto et al. [2019](#page-150-12)), *Micraspidales* (Quijada et al. [2020\)](#page-151-8), and *Umbellidiales* (Barreto et al. [2023](#page-131-7)). Haelewaters et al. ([2021c\)](#page-138-7) formally recognized *Cyttariales* and *Erysiphales* as family-level clades in *Helotiales*, and *Lichinodiales* as a family-level clade in *Leotiales*.

Over time, the following trends can be observed in the class: (1) in general, some genera have been studied signifcantly more than others. For example, the type genus *Leotia* has not received any taxonomic additions in over forty years (Otani [1982\)](#page-149-6). (2) Many species and genera have frequently changed placement both among diferent higher taxa and between taxonomic levels (in both directions). A consequence of this has been an expansive defnition of the now massive order *Helotiales*. (3) The class has been plagued by systematic inconsistencies among diferent studies (Quandt and Haelewaters [2021\)](#page-150-11). Currently, the following orders are recognized: *Chaetomellales*, *Helotiales*, *Lahmiales*, *Lauriomycetales*, *Leotiales*, *Marthamycetales*, *Medeolariales*, *Micraspidales*, *Phacidiales*, *Rhytismatales*, *Thelebolales*, and *Umbellidiales*.

# **Phylogenetic relationships within** *Leotiomycetes*

The classifcation system and taxonomy of *Leotiomycetes* were traditionally based on morphology, in particular sexual and asexual diagnostic characteristics. However, morphologically defned taxa grouped in erected genera, families, and orders are often not supported by molecular phylogenies (e.g., Quijada et al. [2018](#page-151-9); Ekanayaka et al. [2019](#page-136-14); Johnston et al. [2019;](#page-141-0) Haelewaters et al. [2021c;](#page-138-7) Quandt and Haelewaters [2021\)](#page-150-11). In addition, even after the use of DNA characters, molecular phylogenies are often inconsistent and unresolved. The frst molecular phylogenies of helotialean taxa were based on SSU and LSU (Platt and Spatafora [2000](#page-150-13); Gernandt et al. [2001\)](#page-137-13). Note that for some of the currently recognized orders, sequence data does not currently exist (*Lahmiales fde* Quandt and Haelewaters [2021\)](#page-150-11) or has only been generated for ribosomal loci (*Medeolariales fde* LoBuglio and Pfister [2010\)](#page-145-10). In general, limited DNA data are available for most members of the class, notably regarding type specimens.

Based on the most recent studies, current classifcations recognize the following classifcations and associations. The frst diverging order of the class is *Lauriomycetales*, followed by *Umbellidiales* (Barreto et al. [2023](#page-131-7)). Phylogenies disagree on the relationships of other early diverging orders in the class: *Chaetomalleles*, *Leotiales*, *Marthamycetales*, *Micraspidales*, *Phacidiales*, *Rhytismatales*, and *Thelebolales*, and their respective relationships to *Helotiales* (Ekanayaka et al. [2019;](#page-136-14) Johnston et al. [2019;](#page-141-0) Haelewaters et al. [2021d](#page-138-15); Quijada et al. [2022](#page-151-10); Barreto et al. [2023\)](#page-131-7). Historically there has been confusion about how these non-helotialean *Leotiomycetes* taxa are related to one another (Gernandt et al. [2001;](#page-137-13) Lantz et al. [2011](#page-144-8); Quijada et al. [2018](#page-151-9); Karakehian et al. [2019](#page-142-14)). Within *Helotiales*, 53 families were recognized by Quandt and Haelewaters ([2021\)](#page-150-11), and Johnston et al. ([2019\)](#page-141-0) recognized the discinelloid, erysiphoid, helotioid, hyaloscyphoid, mollisioid, pezizelloid, and sclerotinioid clades.

The total number of species in the class difers depending on the classifcation system. According to Wijayawardene et al. ([2020a](#page-160-0)), *Leotiomycetes* comprises 13 orders and 52

families, for a total of 626 accepted genera and 6,453 species, with a major part of *incertae sedis* genera and species distributed in *Helotiales*. Quandt and Haelewaters [\(2021](#page-150-11)) synthesized the numbers of genera and species per family in the then-current classifcation of *Leotiomycetes* and presented 13 orders, with 43 families, 630 genera, and a total of 6,440 species. The most recent compilation of known species and genera of *Leotiomycetes* was presented by Wijayawardene et al. ([2022a](#page-160-1)) and resulted in eleven orders, 55 families, 633 genera, and 6,451 species.

**Type order:** *Leotiales* Korf & Lizoň, Czech Mycol. 52(4): 256 (2001).

**Type family:** *Leotiaceae* Corda, Icon. fung. (Prague) 5: 37 (1842).

**Type genus:** *Leotia* Pers., Neues Mag. Bot.: 31 (1794).

**Type species:** *Leotia lubrica* (Scop.) Pers., Neues Mag. Bot.: 31 (1794).

*Lichinomycetes* V. Reeb, Lutzoni & Cl. Roux, Mol. Phylogen. Evol. 32(3): 1055 (2004).

=*Candelariomycetes* Voglmayr & Jaklitsch, in Voglmayr, Fournier & Jaklitsch, Persoonia 42: 41 (2018).

= *Coniocybomycetes* M. Prieto & Wedin, Cladistics 39(3): 305 (2013).

=*Geoglossomycetes* Zheng Wang, C.L. Schoch & Spatafora, Persoonia 22: 131 ([2009\)](#page-153-0).

=*Sareomycetes* Beimforde, A.R. Schmidt, Rikkinen & J.K. Mitch., Fungal Systematics and Evolution 6: 33 (2020).

= *Xylonomycetes* Gazis & P. Chaverri, in Gazis, Miądlikowska, Lutzoni, Arnold & Chaverri, Mol. Phylogen. Evol. 65(1): 301 (2012).

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**Sexual morph:** *Thallus* when lichen-forming superficial on the substratum, which is usually rock or soil (rarely bark), when not lichen-forming immersed in soil, plants or insects, composed of hyphae or yeast cells. *Ascomata* apothecia, often present, solitary, round and immersed in or superfcial on thallus, or on substrate surface and often stalked. *Hamathecium* composed of paraphyses. *Asci* often eight-spored, but up to multi  $(>100)$ -spored, cylindrical to fusiform or clavate, incompletely bitunicate, often with J+staining tholus. *Ascospores* flamentous, globose, ellipsoidal or fusiform, aseptate or septate, hyaline. **Asexual morph:** coelomycetous. *Conidiomata* infrequently present, pycnidial, developing on the same thalli as the apothecia, usually immersed, globose, with a single locule, pale, ostiolate. *Conidiomata wall* not carbonized. *Conidiophores* lining the inner cavity or reduced, septate, sometimes branched. *Conidiogenous cells* phialidic, discrete, lageniform, hyaline, smooth, producing microconidia. *Conidia* hyaline, aseptate, without appendages.

<span id="page-32-0"></span>**Fig. 11** *Digitothyrea polyglossa*. **a**, **b** habitus on rock. **c**–**f** sections through the thallus. Pictures by F. Schumm

![](_page_32_Figure_3.jpeg)

**Life modes and distribution:** Lichen-forming with cyanobacteria or green algae, or more rarely putative mycorrhizae, endophytes, and yeast-like insect symbionts.

**Notes:** The class *Lichinomycetes* was until recently regarded as one of the smaller classes of the Phylum *Ascomycota,* including only some ascomycetes that are lichenized with cyanobacteria (Díaz-Escandón et al. [2022](#page-135-3)). The study by Díaz-Escandón et al. ([2022](#page-135-3)) however showed that several other fungal classes with widely varying life modes cluster in the clade of *Lichinomycetes*. It regards the (former) classes *Candelariomycetes, Coniocybomycetes, Geoglossomycetes, Sareomycetes,* and *Xylonomycetes*, all of which have to be subsumed into the *Lichinomycetes* following this study.

Based on Lücking et al. ([2017\)](#page-146-4), Díaz-Escandón et al. [\(2022](#page-135-3)), and Wijayawardene et al. ([2022a](#page-160-1)), the class comprises 7 orders, 13 families, around 75 genera, and around 600 species.

The members of the *Lichinomycetes* are distributed over a broad range of ecosystems. They are most conspicuous in desert ecosystems, where they cover vast areas of rock and soil. In all other ecosystems, they are present but less

conspicuously. As with almost every group, they are most diverse in the tropics (Schumm and Aptroot [2023](#page-154-16)).

The main use of *Lichinomycetes* is that they are often a component of soil crusts in desert environments.

**Type order:** *Lichinales* Henssen & Büdel, in Eriksson & Hawksworth, Syst. Ascom. 5(1): 138 (1986).

**Type family:** *Lichinaceae* Nyl., Mém. Soc. Sci. nat. Cherbourg 2: 8 (1854).

**Type genus:** *Lichina* C. Agardh, Syn. Alg. Scand.: xii, 9 (1817).

**Type species:** *Lichina pygmaea* (Lightf.) C. Agardh, Syn. Alg. Scand.: xii, 9 (1817).

*Orbiliomycetes* O.E. Erikss. & Baral, in Eriksson, Baral, Currah & Hansen, Myconet 9: 96 (2003).

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**Sexual morph:** *Ascomata* solitary or aggregated, cupulate to fat or convex, sessile or sometimes distinctly stalked, erumpent or superfcial, usually non-gelatinous, hyaline to bright-coloured (yellow, red, pink etc.), rarely black, exterior

![](_page_33_Picture_2.jpeg)

<span id="page-34-0"></span>**Fig. 12** Macro- and micromorphological characteristics of *Orbili-*◂ *omycetes*. **a‒j** Apothecia; a *Hyalorbilia infatula*. **b** *Orbilia clavuliformis.* **c** *Lilapila oculisporella.* **d** *Orbilia mirabilis.* **e** *O. patellarioides*. **f** *O. orientalis*. **g** *O. rubrovacuolata*. **h** *O. xanthostigma*. **i** *O. fmbriata*. **j** *O. crenatomarginata*; **k** Asci; **k1** *Lecophagus ellipsoideus.* **k**<sub>2</sub> *O. multiaustraliensis*; **l** Paraphyses; **l**<sub>1</sub> *O. sarothamni.* **l**<sub>2</sub> *O. carpoboloides.* **l<sub>3</sub>** *O. eucalypti.* **l<sub>4</sub>** *O. amarilla*; **m.** Ascospores; **m**<sub>1</sub> Lilapila oculispora.  $m_2$  *O.* subsphaerospora.  $m_3$  *Amphosoma resinicola*. **m4** *Hyalorbilia fusispora*. **m5** *O. favida*. **m6** *O. cercidicola*. **m7** *O. xanthoguttulata.*  $m_8$  *O. menageshae.*  $m_9$  *O. carpoboloides.*  $m_{10}$  *O. bicknellensis*. **m11** *O. myriomuscula*. **m12** *O. subaristata*; **n** Conidia;  $n_1$  *O. fissilis.*  $n_2$  *O. aprilis.*  $n_3$  *O. auricolor.*  $n_4$  *O. subvinosa.*  $n_5$  *O. eucalypti*. **n6** *O. patellarioides*. **n7** *Hyalorbilia infatula*

smooth or with short to long, thin-to thick-walled hairs or glassy processes that are frequently agglutinated as teeth. *Ectal excipulum* composed of hyaline isodiametric to prismatic cells. *Medullary excipulum* of hyaline intricate to isodiametric texture. *Hamathecium* composed of paraphyses that are septate, apically uninfated to infated (knob-shaped to spathulate or lanceolate), often covered by exudate, with anastomoses at the base, branched below or also above. *Vacuolar bodies* and *cytoplasmic bodies* typical of various species. *Asci* arising from croziers or usually from simple septa with often bifurcate base, 8–128-spored with often some of the spores inversely oriented,  $\pm$  cylindrical in the region of pars sporifera, unitunicate, forcibly ejecting ascospores as a single compact entity, inamyloid, apex of dead asci hemispherical to truncate (saddle-shaped), thin- or thick-walled, without apical ring. *Ascospores* variously shaped, mostly aseptate, hyaline, smooth-walled, thin-walled, mostly with one or sometimes several globose to lens-shaped or tearshaped to fliform spore bodies which are often associated with some lipid bodies. **Asexual morph:** hyphomycetous; *Conidiomata* solitary, sometimes sporodochial or synnematal. *Setae* lacking. *Conidiophores* macronematous to micronematous, mononematous, scattered or gregarious, hyaline, straight, septate, unbranched or branched, smoothwalled. *Conidiogenous cells* monoblastic, terminal, proliferating sympodially, rarely synchronously, conidial secession schizolytic. *Conidia* solitary, acrogenous, hyaline, thin-walled, smooth, aseptate to multiseptate, unbranched (obovoid, fusiform, cylindrical, falcate, sometimes with protuberances) or variously branched, truncate at the base, eguttulate or guttulate. *Trapping organs* present in *Lecophagus*, *Hyalorbilia*, and *Orbilia* section *Arthrobotrys*: adhesive pegs capturing rotifers, peg-like knobs capturing rhizopods, adhesive knobs, columns or networks, constricting and nonconstricting rings capturing nematodes or rarely arthropods.

**Life modes and distribution:** Members of *Orbiliomycetes* are distributed worldwide in a very broad range of ecosystems with very diferent climatic conditions. Numerous members are drought-tolerant and show their maximum diversity in semihumid to semiarid regions, growing on dead xeric parts of woody plants, while only a smaller part is adapted to moist to semi-aquatic habitats. Various taxa include a parasitic phase: members of *Lecophagus* are adapted to attract and exploit rotifers and some of *Hyalorbilia* prey on rhizopods, while a few closely related species of *Orbilia* section *Arthrobotrys* (*O. alba*, "*Dactylellina*" *copepodii,* "*D.*" *entomopaga*, "*D.*" *ferox*) trap and exploit diferent small arthropods, such as copepods, collemboles, and insects. Many members of section *Arthrobotrys* prey on nematodes and are potential agents in controlling root damage in agriculture. Very few taxa have been exclusively reported as fungicolous, e.g., *Hyalorbilia orbiliicola* growing parasitic on *Orbilia* spp. (Baral et al. [2020\)](#page-131-8).

**Notes**: The class *Orbiliomycetes* was introduced by Eriksson and Baral in Eriksson et al. ([2003\)](#page-136-5) based on extraordinary morphologies and molecular data to include the family *Orbiliaceae* which has extremely increased in the past decades to around 500 species with a known teleomorph (Baral et al. [2020](#page-131-8)). The extraordinary morphology of the bilateral-symmetrical ascus apex and the living ascospores with their spore bodies were first addressed by Baral [\(1994](#page-130-8)). Important comprehensive works on the classifcation of the group were those by Boudier (1904–1910), Velenovský ([1934\)](#page-158-7), Svrček ([1954\)](#page-156-11) and Spooner [\(1987](#page-155-9)) for the teleomorphs. A large number of authors have worked on the predacious anamorphs of *Orbiliomycetes*, one of the most important being Drechsler (e.g., Drechsler [1937\)](#page-136-16). A survey of nematode-trapping taxa was provided by Rubner ([1996\)](#page-152-15) and Yu et al. ([2014\)](#page-161-8).

Pfster [\(1997\)](#page-150-14) and Hagedorn and Scholler ([1999](#page-138-16)) were the frst to apply DNA sequence-based methods to the family to connect sexually and asexually typifed taxa and to suggest relationships for taxa with similar trapping devices by replacing previous concepts that were based on the diferent types of conidiophores and conidia.

According to Baral et al. ([2020\)](#page-131-8), the class currently comprises one order, one family, and ten genera. This was accepted by Wijayawardene et al. ([2020a,](#page-160-0) [2022a](#page-160-1)).

**One fungus one name:** A large number of members of *Orbiliomycetes* have been reported as pleomorphic taxa, and numerous genera have been recognized to accommodate them based on the diferent types of conidiophores and conidia (Baral et al. [2018](#page-131-9)). About 15 of these genera have been accepted prior to the new rules of adopting only one name for one taxon. Diferent opinions exist about the circumscription of a genus. Baral et al. ([2020\)](#page-131-8) proposed a rather wide concept by recognizing a very large genus *Orbilia*.

**Type order:** *Orbiliales* Baral, O.E. Erikss., G. Marson & E. Weber, in Eriksson, Baral, Currah & Hansen, Myconet 9: 96 (2003).

**Type family:** *Orbiliaceae* Nannf., Nova Acta R. Soc. Scient. upsal., Ser. 4 8(no. 2): 250 (1932).

**Type genus:** *Orbilia* Fr., Fl. Scan.: 343 (1836).

**Type species:** *Orbilia xanthostigma* (Fr.) Fr. 1849 (designated by Bachman, Ann. rep. Ohio State Acad. Sci. 5(no. 2): 57. 1909); *fde* Index Fungorum (2023).

*Pezizomycetes* O.E. Erikss. & Winka, Myconet, 1(1): 8 (1997).

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**Sexual morph:** *Ascomata* epigeous or hypogeous, of apothecia or cleistothecia types, more rarely of perithecia type, exceptionally without a complex structure, i.e. asci produced on a simple hyphal structure. Apothecia of various shape, e.g. discoid, cupuliform, pulvinulate, cerebriform, saddleshaped, etc., stipitate or not. Cleisthotecia usually more or less spherical or potato-like. Flesh of apothecia compound with an exposed hymenium and an excipulum, the latter made of a single structure or more complex (multi-layered). Flesh of cleistothecia (also called gleba) has a diferentiate hymenium or not, accompanied with sterile tissues, surrounded by a peridium. *Asci* with a single wall (unitunicate), inamyloid or amyloid (more or less difusely in the whole length, or more strongly at the apex), operculate with an operculum placed at the top or eccentrically, or sometimes without diferentiated apical structure, breaking down before maturity (prototunicate), more rarely evanescent, mainly 8-spored, sometimes 2-, 4-, 16-, 32-spored or more, with or without croziers at the base. *Ascospores* always aseptate, mainly symmetrical, smooth or variously ornamented, with or without oil drops, mainly containing 1 to 4 nuclei, hyaline, more rarely coloured. *Paraphyses* present, septate, simple or branched, sometimes anastomosing, hyaline or containing various type of granular, vacuolar or cristalloid pigments (e.g. carotenoids), sometimes also with external pigments that may form an epithecium. **Asexual morph:** Hyphomycetous, of various types, e.g. chromelosporiumlike and oedocephalum-like in *Pezizaceae*, dichobotrys-like in *Pyronemataceae* and *Pyropyxidaceae*, mollardiomyceslike in *Sarcoscyphaceae*, conoplea-like in *Sarcosomataceae*, verticicladium-like in *Chorioactidaceae*, costantinella-like in *Morchellaceae*, etc. The conidia are formed blastically. Some genera are only known to be anamorphic, e.g. *Glaziella*, *Heydenia*, *Phymatotrichopsis*.

**Life modes and distribution:** *Pezizomycetes* are present all around the world, in a wide range of habitats, although they seem more represented in temperate and boreal zones, as well as in Mediterranean basin. They are mainly saprobic and terrestrial but can be found on other substrates such as decaying wood, dead plants or dung. Some genera are known to have species that form ectomycorrhizae (Hobbie et al. [2001](#page-140-13); Tedersoo et al. [2006,](#page-156-12) [2010\)](#page-156-13), including hypogeous members of these genera. There are also some parasitic genera or species, e.g. *Octospora* Hedw. and *Lamprospora* De Not. are bryoparasites (Döbbeler [1979](#page-135-11); Benkert [1993\)](#page-131-10), *Rhizina undulata* Fr. parasites the roots of pine trees, especially those damaged by fre (Sierota [1998;](#page-155-10) Lee et al. [2005](#page-144-9)), the asexual morph of *Caloscypha fulgens* (Pers.) Boud. — previously known under the name *Geniculodendron pyriforme* — is a seed pathogen of conifers (Salt [1974;](#page-152-16) Schröder et al. [2002](#page-153-3)), the anamorph *Phymatotrichopsis omnivora* (Shear) Hennebert is a root pathogen of dicotyledonous plants (Uppalapati et al. [2010](#page-157-9)). Some species are also known to be strictly pyrophilous, growing in postfre environments (Petersen [1970\)](#page-150-15). Many of them are known to be endophytes or endolichenic fungi (Raudabaugh et al. [2020](#page-151-11); Healy et al. [2021](#page-139-11)).

**Notes:** *Pezizomycetes* was introduced by Eriksson and Winka ([1997](#page-136-2)). This class includes only one order, *Pezizales*, the so-called "operculate discomycetes", a group of cup-fungi distinguished from the other feshy ascomycetes by Boudier ([1885](#page-132-6)) based on the dehiscence mode of asci, i.e. presence of an operculum. This systematics was widely used during the twentieth century, until the works of Berthet [\(1964](#page-132-7)), Eckblad ([1968](#page-136-17)) and Korf ([1972](#page-143-2)) who proposed some changes based on morphological characters and nomenclatural aspects. Main changes were introduced in the molecular era with the frst phylogenetic works by Gargas and Taylor ([1995](#page-137-14)), Landvik et al. ([1997\)](#page-144-10), Norman and Egger ([1999](#page-149-7)), Hansen and Pfister ([2006\)](#page-138-17) and Perry et al. ([2007\)](#page-150-16). Kirk et al. [\(2008](#page-142-1)) listed 16 families, 200 genera and about 1684 species. Jaklitsch et al. [\(2016](#page-141-10)) also listed 16 families, but 172 genera and about 1608 species, while Ekanayaka et al. [\(2018](#page-136-18)) listed 21 families and 218 accepted genera. During the last years, three new families were added (Kraisitudomsook et al. [2020](#page-143-3); Van Vooren [2021;](#page-158-8) Zeng et al. [2022](#page-161-9)). There are about 235 accepted genera based on our last count.

Beyond their roles in natural ecosystems, some *Pezizomycetes* have an important economic impact on the food industry, i.e. truffles culture (Oliach et al.  $2021$ ) and wild mushrooms collecting like morels (Malone et al. [2022](#page-146-12)).

For a more detailed view of *Pezizomycetes*, see Pfister and Healy ([2021](#page-150-17)).

**Type order:** *Pezizales* J. Schröt., *in* Engler & Prantl, Nat. Pfanzenfam., I: 173 (1897) [as "*Pezizineae*"].

**Type family:** *Pezizaceae* Dumort. Analyse des familles des plantes: 72 (1829).

**Type genus:** *Peziza* Dill. ex Fr., Syst. mycol. (Lundae) 2(1): 40 (1822).

**Type species:** *Peziza vesiculosa* Bull., Hist. Champ. Fr., 1(2): 270 (1790), *nom. sanct.*


**Fig. 13** Macro morphologies of taxa in *Pezizomycetes.* **a** *Peziza vesiculosa*. **b** *Daleomyces brunneoviolaceus*. **c** *Cheilymenia crucipila*. **d** *Helvella monachella*. **e** *Caloscypha fulgens*. **f** *Morchella deliciosa.* **g** *Neournula pouchetii*. **h** *Ascobolus lignatilis*. **i** *Discina perlata*.

**j** *Tuber melanosporum*. **k** *Scutellinia nigrohirtula*. **l** *Wilcoxina gregaria*. **m** *Genea verrucosa.* **n** *Pyronema domesticum*. **o** *Orbicula parietina.* **p** *Lamprospora leptodictya*. (All photos by N. Van Vooren, except *Orbicula parietina* with the courtesy of M. Poulain)



<span id="page-38-0"></span>**Fig. 14** Morphological features of diferent taxa in *Sordariomycetes.* ◂ **a** Stromata of *Xylaria* sp. **b** stromata of *Collodiscula* sp. **c** Stromata of *Daldinia* sp. **d** Stromata of *Xylaria bambusicola.* **e** Fruiting bodies of *Nectria pseudotrichia.* **f** Fruiting bodies of *Eutypa linearis.* **g** Stromata of *Xylaria* sp. **h** Fruiting bodies of *Astrocystis mirabilis.* **i** Longitudinal section of the fruiting bodies of *Daldinia bambusicola.* **j, k** Longitudinal section of *Eutypa linearis.* **l, m** Longitudinal section of *Arthrinium longistromum.* **n** Longitudinal section of *Leptosporella bambusae.* **o** Longitudinal section of perithecium of *Anthostomella* sp. **p** Longitudinal section of *Daldinia bambusicola*. **q** Longitudinal section of perithecium of *Lopadostoma* sp*.* **r** Longitudinal section of perithecium of *Fasciatispora* sp. **s** Longitudinal section of perithecium of *Nigropunctata* sp*.* **t** Conidia arrangement of *Pleurophragmium bambusinum.* **u** Ascus of *Reticulosphaeria* sp*.* **v** Ascus of *Xylaria* sp*.* **w** Ascus of *Fasciatispora* sp*.* **x** Ascus of *Pemphidium* sp*.* **y** Ascus of *Nigropunctata* sp*.* **z** Ascus of *Collodiscula* sp*.* **aa** Ascus of *Arecophila* sp. **bb** Ascus of *Eutypella* sp. **cc** Ascus of *Arthrinium rasikravindrae*. **dd** Ascus of *Leptosporella bambusae.* **ee** Ascospore of *Collodiscula* sp*.* **f** Ascospore *Spirodecospora* sp*.* **gg** Ascospore *Anthostomella* sp*.* **hh** Ascospore *Reticulosphaeria* sp*.* **ii** Ascospore *Durotheca* sp. **jj** Ascospore *Pseudomassaria* sp*.* **kk** Ascospore of *Pleurophragmium bambusinum.* **ll** Ascospore of *Cytospora* sp. **mm** Ascospore of order *Diaporthales* sp. **nn** Ascospore of *Arthrinium* sp. **oo** Conidia of *Arthrinium yunnanum.***pp** Ascospore of *Pustulomyces bambusicola*

*Sordariomycetes* O.E. Erikss. & Winka, Myconet 1(1): 10 (1997).

Index Fungorum Registration Identifer 90,350; Fig. [14](#page-38-0)

**Sexual morph:** *Stromata* immersed to erumpent, composed of orange, brown or black parenchymatous tissues, ascomata embedded in stromatic tissues. *Ascomata* solitary or aggregated, subglobose to cylindrical or ovoid to ellipsoidal, immersed to superficial, ostiolate, papillate or epapillate, periphysate, brown to black. *Hamathecium* composed of paraphyses or internal structures absent; centrum initially flled with a hyaline pseudoparenchyma, dissolving at maturity. *Asci* four-spored or eight-spored or multi, cylindrical to fusiform or clavate to ellipsoidal, unitunicate, deliquescent, with J-, apical ring or indistinct. *Ascospores* flamentous or ellipsoidal to fusiform, aseptate or septate, hyaline or brown or becoming dark at maturity, allantoid, clavate, cylindrical to ellipsoid, 0–3-septate, smooth-walled, with or without guttules, thick- or thin-walled, with or without apical chambers. **Asexual morph:** coelomycetous or hyphomycetous. **Coelomycetes:** *Conidiomata* acervular or pycnidial or intermediate, scattered or aggregated, immersed, globose, coriaceous, with single or multi locules, yellow or orange or dark brown to black, ostiolate or aostiolate. *Conidiomata wall* comprising thin-walled, almost hyaline to brown cells of *textura angularis*. *Conidiophores* lining the inner cavity or reduced to basal layer sometimes reduced to conidiogenous cells, occasionally, septate, branched. *Conidiogenous cells* holoblastic, enteroblastic to phialidic, discrete, lageniform, subcylindrical to cylindrical, hyaline to pale yellow, smooth, producing macroconidia and sometimes microconidia from same conidiogenous cell. *Conidia* hyaline when young, hyaline or brown to dark brown at maturity, aseptate to multiseptate, thick-walled or smooth-walled, guttulate or egutulate, with appendage or not. **Hyphomycetes:** *Conidiomata* solitary, sporodochia or synnemata. *Setae* lacking, or present occasionally. *Conidiophores* macronematous to micronematous, mononematous, scattered or gregarious, brown, straight, aseptate or septate, unbranched or branched, smooth-walled. *Conidiogenous cells* monoblastic, terminal, integrated or discrete, percurrent or proliferating sympodially. *Conidia* solitary, acrogenous, brown to dark brown, aseptate to septate, obclavate, fusiform, cylindrical, pyriform to obovoid, straight, subtruncate to obtuse at the apex, truncate or swollen at the base, with or without basal frill eguttulate or guttulate.

**Life modes and distribution:** The members of *Sordariomycetes* are distributed in a broad range of ecosystems (worldwide) and are reported as diferent life modes. Phytopathogenic members of the class are responsible for causing leaf, stem, root, and fruit diseases (e.g. *Cytospora chrysosperma*, *Coniella vitis*, *Diaporthe eres*), while some are reported as pathogenic to arthropods and mammals (e.g. *Sporothrix*, *Fusarium*). Some members are endophytes (e.g. *Pestalotiopsis*) and some are saprobic taxa that are important in decomposition and nutrient cycling (e.g. *Chaetomium*, *Neurospora*). Some taxa have been exclusively reported to be fungicolous.

**Notes:** The class *Sordariomycetes*, one of the largest classes of the phylum *Ascomycota* was introduced by Eriksson and Winka ([1997](#page-136-0)). Nevertheless, previous exceptional studies by Barr ([1983](#page-131-0), [1987,](#page-131-1) [1990](#page-131-2)) and Eriksson and Hawksworth ([1986,](#page-136-1) [1993\)](#page-136-2) provided a comprehensive background for the classifcation of these unitunicate fungi. Classifcation of sexually typifed genera in *Sordariomycetes* was provided along with the Phylum *Ascomycota* by Lumbsch and Huhndorf ([2007](#page-146-0), [2010](#page-146-1)).

Hyde et al.  $(2011)$  $(2011)$  made the first attempt at compiling asexually typifed genera of the kingdom *Fungi* and listed the genera and morphologically doubtful genera (like genera) according to their 'natural classifcation'. After completion, Hyde et al. [\(2011\)](#page-140-0) provided the asexually typifed genera in *Sordariomycetes,* and subsequent studies by Wijayawardene et al. [\(2012,](#page-159-0) [2017b,](#page-160-0) [2022b\)](#page-160-1) updated the lists with the notes, and Wijayawardene et al. [\(2021b\)](#page-160-2) provided the entries for all pleomorphic genera in *Sordariomycetes*. Maharachchikumbura et al. [\(2015\)](#page-146-2) provide the frst attempts at assembling both sexually and asexually typifed genera, along with morphology and DNA sequence-based phylogenies. Hongsanan et al. ([2017\)](#page-140-1) provided the divergence time estimates for the various orders and families in the class. Hyde et al. ([2020\)](#page-141-0) is the latest study that provides a comprehensive background

on the taxonomy of the genera and the higher-level classifcation of *Sordariomycetes*.

*Sordariomycetes* have been recognized with six subclasses i.e. *Hypocreomycetidae*, *Sordariomycetidae* and *Xylariomycetidae* (Eriksson and Winka [1997](#page-136-0)), *Diaporthomycetidae*, *Lulworthiomycetidae* (Maharachchikumbura et al. 2016), and *Savoryellomycetidae* (Hongsanan et al. [2017](#page-140-1)).

Many members of *Sordariomycetes* have been reported as pleomorphic taxa (Kirk et al. [2008](#page-142-0)). Thus, it was proposed to adopt one name (between two generic names) when the links between the morphs are confrmed. Rossman et al. ([2015b\)](#page-152-0) and Réblová et al. [\(2016\)](#page-151-0) are important publications that deal with the recent nomenclatural changes.

**Type order:** *Sordariales* Chadef. ex D. Hawksw. & O.E. Erikss., Syst. Ascom. 5(1): 182 (1986).

**Type family**: *Sordariaceae* G. Winter [as 'Sordarieae'], Rabenh. Krypt.-Fl., Edn 2 (Leipzig) 1.2: 162 (1885).

**Type genus:** *Sordaria* Ces. & De Not., Comm. Soc. crittog. Ital. 1(fasc. 4): 225 (1863).

**Type species:** *Sordaria fmicola* (Roberge ex Desm.) Ces. & De Not. 1863.

*Xylobotryomycetes* Voglmayr & Jaklitsch, Persoonia 42: 41 (2018).

Index Fungorum Registration Identifer 826,791; Fig. [15](#page-40-0)

**Sexual morph:** *Stromata* superficial ascomata bearing branched or unbranched. *Ascomata* perithecioid, perithecioid, subglobose to ellipsoid. *Ostiolar* canal periphysate. *Hamathecium* fliform, septate, hyaline, thin-walled, embedded in a gelatinous matrix, paraphyses. *Asci* 4–8-spored, bitunicate, fssitunicate, apex with an inamyloid apical ring. *Ascospores* ellipsoid to fusiform, 2-celled, brown, septate, with longitudinal germ slits (Voglmayr et al. [2019\)](#page-158-0). **Asexual morph: Coelomycetous:** pycnidial, cylindrical, unilocular. *Conidiogenous cells* meristem, producing pigmented, septate arthroconidia. *Conidia* pigmented, septate (Voglmayr et al. [2019](#page-158-0)).

**Life modes and distribution:** The members of *Xylobotryomycetes* are distributed in a broad range of ecosystems viz*.,* tropical, subtropical to warm temperate, humid areas and are reported as diferent life modes (Réblová and Seifert [2012](#page-151-1); Voglmayr et al. [2019\)](#page-158-0). Most of the members are saprobic.

**Notes:** Voglmayr et al. [\(2019](#page-158-0)) introduced *Xylobotryomycetes* with *Xylobotryales* based on unique morphological features and phylogeny analyses. This order contains two families, *Cirrosporiaceae* and *Xylobotryaceae* (type family) and each family has a single genus *Cirrosporium* and *Xylobotryum*, respectively. The morphology of the type genus of *Xylobotryaceae*, *Xylobotryum*, is very similar to *Xylaria* species, as they produce dark carbonaceous stromata, but *Xylobotryum* produces naked perithecia and twocelled ascospores (Trierveiler-Pereira et al. [2008](#page-157-0)). The position of *Xylobotryum* within *Ascomycota* was uncertain until Voglmayr et al. [\(2019](#page-158-0)) used molecular data to determine its placement. Classifcation of the type genus of *Cirrosporiaceae* (*Cirrosporium*) within *Eurotiomycetes*, which is close to *Mycocaliciales,* has been suggested by Réblová and Seifert ([2012\)](#page-151-1), which can be a new order. An extensive analysis of *Ascomycota* by Schoch et al. [\(2009](#page-153-0)) resolved the ancestor of the *Eurotiomycetes* and Voglmayr et al. ([2019](#page-158-0)) showed that *Cirrosporium* is basal to the *Eurotiomycetes*.

**Type Order:** *Xylobotryales* Voglmayr & Jaklitsch, in Voglmayr, Fournier & Jaklitsch, Persoonia 42: 41 (2018). **Type Family:** *Xylobotryaceae* Voglmayr & Jaklitsch, in Voglmayr, Fournier & Jaklitsch, Persoonia 42: 42 (2018). **Type Genus:** *Xylobotryum* Pat., in Patouillard & Lagerheim, Bull. Herb. Boissier 3(1): 69 (1895). **Type Species:** *Xylobotryum andinum* Pat., in Patouillard & Lagerheim, Bull. Herb. Boissier 3(1): 69 (1895).

**Subphylum** *Saccharomycotina* O.E. Erikss. & Winka, Myconet 1(1): 10 (1997).

Index Fungorum Registration Identifer 501,470.

**Notes:** Eriksson and Winka [\(1997](#page-136-0)) introduced this subphylum to accommodate a single class, *Saccharomycetes*. Groenewald et al. ([2023\)](#page-137-0) comprehensively addressed *Saccharomycotina* and introduced *Alloascoideomycetes*, *Dipodascomycetes*, *Lipomycetes*, *Pichiomycetes*, *Sporopachydermiomycetes* and *Trigonopsidomycetes*. Hence, the subphylum consists of seven classes.

*Alloascoideomycetes* M. Groenew., Hittinger, Opulente & A. Rokas Stud. Mycol. 105: 7 (2023).

Index Fungorum Registration Identifer 847278; Fig. [16](#page-42-0)a, b-d

**Sexual reproduction:** Firm walled *asci* are ellipsoidal to elongate and form laterally or terminally on hyphae, singly or in small whorls. Greater than eight and up to 400 *ascospores* can form in an *ascus* and released by apical deterioration. *Ascospores* are ellipsoidal but can also appear hat-shaped due to a unilateral, mucilaginous brim. **Asexual reproduction:** Colonies are cream-colored and can be dry or slimy. Budding cells, *pseudohyphae* and *true hyphae* are present or absent as species are often dimorphic, with colonies being restricted and yeast-like or expanding and hyphal. Cell division is by multilateral budding and formation of *pseudohyphae* and septate *hyphae*. Clavate *blastoconidia* of variable size form on <span id="page-40-0"></span>**Fig. 15** *Xylobotryum andinum* (WU 39969, epitype). **a** Heads of branched stroma showing ascomata (MJF 07074). **b** Ascomata in vertical section. **c** Peridium. **d** Immature and mature asci. **e–g** Ascospores. Scale bars:  $\mathbf{a} = 1$  mm,  $\mathbf{b} = 0.2$  mm, **c**, **d** = 20 μm, **e**–**g** = 5 μm. Redrawn from Voglmayr et al. ([2019\)](#page-158-0)



wide *hyphae* and may be sessile or arise from denticles and occur singly or in short, branched chains. **Physiology/biochemistry:** Fermentation is absent and relatively few carbon compounds are assimilated. Urease activity is absent. The diazonium blue B reaction is negative or, rarely, weak. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein families OG0009556 and OG0024318 (Groenewald et al. [2023\)](#page-137-0) as well as phylogenetic analyses using DNA sequences encoding LSU rDNA, SSU rDNA, *tef1-α*, *rpb1*, and *rpb2* (Kurtzman and Robnett [2013a](#page-143-0)).

**Life mode and distribution:** The small number of *Alloascoideomycetes* members have been primarily reported from wood, including decaying wood and oak trees (*Quercus* sp., *Fagaceae*) and on occasion associated with bark beetles (*Hylecoetus dermestoides*) and once from a larvae in West Africa (Kurtzman et al. [2011](#page-143-1)).

**Notes:** The class *Alloascoideomycetes* in subphylum *Saccharomycotina* was introduced by Groenewald et al. ([2023](#page-137-0)), who, based on the analysis of genomic data, proposed the distribution of the yeasts assigned to *Saccharomycotina* among seven classes. They provided a diagnosis



of *Alloascoideomycetes* with class-specific orthologous groups of proteins (OGs). For more details, see the entry *Saccharomycetes*. Analysis of the two yeast DNA barcodes, LSU rDNA, SSU rDNA, and the gene sequence for the D1/D2 domain of the LSU rDNA (Kurtzman and Robnett [1998](#page-143-2)) indicated that two *Ascoidea* species, *Ascoidea africana* and *Ascoidea hylecoeti* are not closely related to *Ascoidea rubescens*. Kurtzman and Robnett ([2013a](#page-143-0)) clearly divided *Ascoidea* species into two genera, *Ascoidea* and

*Alloascoidea*, where the latter was included as the sole genus in the family *Alloascoideaceae*. *Alloascoidea* was found to be homothallic and produces clusters of more than eight ascospores in an ascus (Kurtzman and Robnett [2013a](#page-143-0)). Groenewald et al. ([2023](#page-137-0)) is the latest study that provides a comprehensive background on the taxonomy of the higherlevel classifcation of *Alloascoideomycetes*. Although only two species of *Alloascoidea* are presented by the class*,* Groenewald et al. ([2023\)](#page-137-0) proposed the separation of *Alloascoidea*

<span id="page-42-0"></span>**Fig. 16** Microscopic features of taxa in *Saccharomycotina*. **a** Sep-◂ tate non-sporulating hyphae and pseudohyphae of *Alloascoidea hylecoeti* (*Alloascoideaceae*, *Alloascoideales*, *Alloascoideomycetes*). **b**–**d** *Alloascoidea africana* (*Alloascoideaceae*, *Alloascoideales*, *Alloascoideomycetes*) asci with ascospores and released ascospores (indicated with arrows). **e**, **f** *Dipodascus albidus* (*Dipodascaceae*, *Dipodascales*, *Dipodascomycetes*). **e.** Ascospores within hyphae and released ascospores. **f** Hyphae, arthroconidia and ascospores (indicated by arrow). **g** *Lipomyces starkeyi* (*Lipomycetaceae*, *Lipomycetales*, *Lipomycetes*) conjugation between a cell and its bud, ascus with ellipsoidal ascospores (indicated by arrows). **h**–**j** *Pichia membranifaciens* (*Pichiaceae*, *Pichiales*, *Pichiomycetes*) budding cells and asci with hat-shaped ascospores (indicated by arrows). **k** *Pachysolen tannophilus* (*Pachysolenaceae*, *Alaninales*, *Pichiomycetes*) budding cells, ascus with ascospores and liberated ascospores (indicated by arrow). **l**, **m** *Metschnikowia bicuspidata* (*Metschnikowiaceae*, *Serinales*, *Pichiomycetes*). **l** *M. bucuspidata* var. *californica* budding cells and needle-shaped ascospores. **m** *M. bucuspidata* var. *chathamia* budding cells and needle-shaped ascospores. **n** *Saccharomyces cerevisiae* (*Saccharomycetaceae*, *Saccharomycetales*, *Saccharomycetes*) budding cells and unconjugated, persistent ascus with globose ascospores (indicated by arrow). **o**, **p** *Phafomyces opuntiae* (*Phaffomycetaceae*, *Phafomycetales*, *Saccharomycetes*). **o** Budding cells and asci with hat-shaped ascospores (indicated by arrows). **p** Budding cells in short chains and asci with hat-shaped ascospores (indicated by arrows). **q** *Saccharomycodes ludwigii* (*Saccharomycodaceae*, *Saccharomycodales*, *Saccharomycetes*) bipolar cell division of budding cells and asci with spherical ascospores (indicated by arrows). **r**–**t** *Ascoidea rubescens* (*Ascoideaceae*, *Ascoideales*, *Saccharomycetes*). **r** Asexual cells and cluster of ascospores (indicated by arrow) **s** Asexual cells and septate hyphae, **t** Ascus with spores in diferent states of nascendi. **u** *Sporopachydermia lactativora* (*Sporopachydermiaceae*, *Sporopachydermiales*, *Sporopachydermiomycetes*) budding cells. **v** *Trigonopsis variabilis* (*Trigonopsidaceae, Trigonopsidales*, *Trigonopsidomycetes*). Budding cells with ellipsoidal and triangular morphology. Triangular cells produce ellipsoidal and triangular buds. Scale bar: **a**=5 μm, **b**–**q**, **u**, **v**=10 μm, **r**–**t**=30 μm

from its closest relative, *Sporopachydermia*, due to their long branches into two sister classes.

According to Groenewald et al. ([2023](#page-137-0)), currently, the class comprises one order *Alloascoideales*, one family *Alloascoideaceae*, and one genus *Alloascoidea*.

**Type order:** *Alloascoideales* M. Groenew., Hittinger, Opulente & A. Rokas. Stud. Mycol. 105: 7 (2023).

**Type family**: *Alloascoideaceae* Kurtzman & Robnett, FEMS Yeast Res. 13: 429 (2013).

**Type genus:** *Alloascoidea* Kurtzman & Robnett, FEMS Yeast Res. 13: 426 (2013).

**Type species:** *Alloascoidea hylecoeti* (L.R. Batra & Francke-Grosm.) Kurtzman & Robnett, FEMS Yeast Res. 13: 427 (2013).

*Dipodascomycetes* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 10 (2023).

Index Fungorum Registration Identifier 847281; Fig. [16e](#page-42-0),f

**Sexual reproduction:** *Ascomata* absent. *Asci* free or attached to hyphae, unconjugated or formed by isogamous or heterogamous conjugation of yeast cells or conjugation of gametangia, one- to many-spored, subspheroid ovoid, clavate, ellipsoid, cylindrical, tubular or acicular, an apical cell may be attached to it. Asci persistent, deliquescent or ascospores are liberating following rupture of the tip of the ascus. *Ascospores* hyaline or brown, spheroid, subspheroid, hemispheroid, ellipsoid, cylindrical, bacilliform, hat- or helmet-shaped, aseptate, smooth or rough, may be embedded in transparent coating material, ledge is present or absent. Species are homothallic or heterothallic. **Asexual reproduction:** White to cream coloured yeast-like or flamentous colonies. Spherical, ellipsoid, ovoid, elongated or apiculate budding cells present or absent. Two layered cell wall, holoblastic, multilateral budding or bipolar bud-fssion, solitary or catenate, aseptate blastoconidia present or absent. Pseudohyphae and true hyphae present or absent. Arthroconidia present or absent. Chlamydospores rarely present. Sympodial or anneleidic conidiogenesis on undiferentiated conidiogenous cells rarely present. **Physiology/biochemistry:** Fermentation is present or absent. Nitrate assimilation is variable. The diazonium blue B reaction is negative, urease is not produced. The major ubiquinone, where known, is coenzyme Q-6 or Q-9. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein families OG0005588, OG0005810, and OG0006132 (Groenewald et al. [2023\)](#page-137-0) as well as phylogenetic analyses using DNAsequences encoding LSU rDNA, SSU rDNA, *tef1-α*, *rpb1*, and *rpb2* (Kurtzman and Robnett [2013b](#page-143-3)).

**Life mode and distribution:** While some species have a restricted distribution, others occur worldwide in diferent terrestrial and aquatic ecosystems. Several species are associated in varying degrees with fowers, tree exudates, fruits, insects, and humans, or are saprobes in soil, rotten wood and other decaying plant materials. *Trichomonascus mycophagus* is a mycoparasite. *Magnusiomyces capitatus* is often recovered from the materials of patients (Kurtzman et al. [2011](#page-143-1); de Hoog et al. [2020](#page-135-0)).

**Notes:** The class *Dipodascomycetes* in the subphylum *Saccharomycotina* was introduced by Groenewald et al. [\(2023](#page-137-0)), who, based on the analysis of genomic data, proposed the distribution of the yeasts assigned to *Saccharomycotina* among seven classes. They provided a diagnosis of *Dipodascomycetes* with class-specifc orthologous groups of proteins (OGs). For more details, see the entry on *Saccharomycetes*. ITS rDNA phylogeny combined with nDNA/DNA reassociation data and physiological characters of ascomycete yeasts reproducing mainly by arthric conidiogenesis were used for species delimitation in *Geotrichum* and related genera by

de Hoog and Smith ([2004](#page-134-0)). As a result, they classifed the teleomorphs of *Geotrichum* under genera *Galactomyces* and *Dipodascus* while *Magnusiomyces* with *Saprochaete* anamorphs accommodated the related group of arthroconidium forming yeasts. Several yeast species currently assigned to *Dipodascomycetes* were studied by means of multigene phylogenetic analysis by Kurtzman and Robnett ([2007](#page-143-4)), who also proposed *Sugiyamaella*. Recent genome-scale phylogenetic studies (Shen et al. [2016](#page-154-0), [2018](#page-154-1), [2020](#page-154-2)) also included several taxa from class *Dipodascomycetes*.

According to Groenewald et al. [\(2023](#page-137-0)) the class currently comprises one order, *Dipodascales*, two families *Dipodascaceae* and *Trichomonascaceae* and 17 genera. The members of the class have been reported as important taxa in medical and other industries (e.g. *Galactomyces candidus*, *Magnusiomyces capitatus* and *Yarrowia lipolytica* are reported as human pathogens *fde* Groenewald et al. [\(2014](#page-137-1)), de Hoog et al. [\(2020\)](#page-135-0), *Yarrowia lipolytica* and *Starmerella bombicola* have been used in food and biotechnology industries *fde* Groenewald et al. ([2014\)](#page-137-1), Qazi et al. [\(2022](#page-150-0))).

**Type order:** *Dipodascales* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 10 (2023).

**Type family:** *Dipodascaceae* Engl. & E. Gilg, Syllabus der Pfanzenfamilien: 59. (1924).

**Type genus:** *Dipodascus* Lagerh., Jahrb. Wiss. Bot. 24: 549. (1892).

**Type species:** *Dipodascus albidus* Lagerh., Jahrb. Wiss. Bot. 24: 549. (1892).

*Lipomycetes* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 10 (2023).

Index Fungorum Registration Identifer 847285; Fig. [16](#page-42-0)g

**Sexual reproduction:** *Ascomata* absent. *Asci* unconjugated or formed by isogamous or heterogamous conjugation, or by fusion of buds of the same cell, or a bud with another cell, or by fusion of hyphal tips or by terminal and penultimate hyphal cells, one- to many-spored, globose, ellipsoid, pear shaped, saccate, tubular, cylindrical or acicular, persistent or deliquescent. *Ascospores* light brown, amber coloured or hyaline, spheroid, ellipsoid, reniform, allantoid or cymbiform, smooth walled or warty, or ornamented with longitudinal ridges. **Asexual reproduction:** Hyaline, often mucoid yeast-like or flamentous colonies. Spherical, ovoid, ellipsoid or elongated budding cells usually present. Multilateral budding. Pseudohyphae and true hyphae present or absent. **Physiology/biochemistry:** production of intracellular lipids and extracellular starch-like compounds, fermentation is absent, nitrate is not assimilated. The major ubiquinone, where known, is coenzyme Q-8, Q-9 or Q-10. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein families OG0010973, OG0011052, OG0008472, and OG0009553 (Groenewald et al. [2023](#page-137-0)) as well as phylogenetic analyses using either LSU rDNA or DNA sequences encoding SSU rDNA, LSU rDNA, mtSSU rDNA, and *tef1-α* (Kurtzman et al. [2007](#page-143-5)).

**Life mode and distribution:** The majority of the species assigned to *Lipomycetes* are most frequently recovered from soil, while some of them are insect-associated or are reported from tree fuxes, wood cutting and cacti (Kurtzman et al. [2011](#page-143-1); Yamazaki et al. [2020](#page-161-0)).

**Notes:** The class *Lipomycetes* in the subphylum *Saccharomycotina* was introduced by Groenewald et al. ([2023\)](#page-137-0) based on genomic data and diagnosed with class-specifc orthologous groups of proteins (OGs). For more details, see the entry *Saccharomycetes*. Currently, *Lipomyces* and its anamorphic counterpart *Myxozyma*, *Babjevia*, *Dipodascopsis* and *Kockiozyma* are assigned to a single order and single family within the class (Groenewald et al. [2023\)](#page-137-0). Based on their multigene phylogenetic analyses, Kurtzman et al. [\(2007](#page-143-5)) reassigned the species of *Kawasakia*, *Smithiozyma*, *Waltomyces* and *Zygozyma* to the genus *Lipomyces* and transferred *Babjevia anomala*, the single species in the genus, to *Dipodascopsis*. The genus *Lipomyces* was interpreted in a narrower sense by Jindamorakot et al. [\(2011](#page-141-1), [2012\)](#page-141-2) and they proposed novel monotypic genera, *Limtongia* and *Kockiozyma* for *Lipomyces* (*Zygozyma*) *smithiae* and *Lipomyces* (*Zygozyma*) *suomiensis*, respectively. Yamazaki et al. [\(2020](#page-161-0)) used multigene studies to investigate the relationships among lipomycetaceous yeasts and included in their analysis 14 undescribed species isolated from soil. They reinstated the genus *Babjevia* and described two novel *Babjevia* species. In addition, they described twelve anamorphic *Lipomyces* species. *Lipomyces starkeyi* was regarded as a species that can be used in biotechnology (McNeil and Stuart [2018](#page-147-0)).

**Type order:** *Lipomycetales* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 11 (2023).

**Type family:** *Lipomycetaceae* E.K. Novák & Zsolt, Acta Bot. Acad. Sci. Hung. 7: 97. (1961).

**Type genus:** *Lipomyces* Lodder & Kreger-van Rij, The Yeasts: A Taxonomic Study: 669. (1952).

**Type species:** *Lipomyces starkeyi* Lodder & Kreger-van Rij, The Yeasts: A Taxonomic Study: 669. (1952).

*Pichiomycetes* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 11 (2023).

Index Fungorum Registration Identifier 847268; Fig. [16h](#page-42-0)-m

**Sexual reproduction:** *Ascomata* absent. *Asci* unconjugated or formed after conjugation between a cell and its bud, or between independent cells. Asci may arise from conjugating haploid or from diploid cells, be formed on hyphae or a tube and may show a variety of shapes, such as globose, subglobose, lenticular, allantoid, elongate, clavate, sphaeropedunculate, or ellipsoidopedunculate. Asci persistent or deliquescent, with quick or slow spore release, thick- or thin-walled, usually contain one to four, more infrequently up to 16 ascospores. *Cephaloascus* forms erect and *Pachysolen* short tubular ascophores. *Ascospores* are spheroid, globose, ovoid, ellipsoid, hemispheroid, elongate, fattened, tapering at the ends, coiled nematode-like, hat-shaped, helmet-shaped, capshaped, clavate, needle-shaped, attenuated at one or both ends, occasionally swollen along one half, may contain a small oil droplet. *Ascospore* surface may be smooth, rough, or warty, with spiral ridges and may be ornamented with brims, equatorial or subequatorial ledges, membranes, be surrounded by a sheath and may or may not tend to agglutinate when released. Species are homothallic or heterothallic. **Asexual reproduction:** Cells divide by holoblastic budding, mostly multilateral on a narrow base, or bipolar on a broad or narrow base, cells occur singly, in short chains, in clusters, or may be formed at the end of a short protuberance. Cell wall is two layered. Cells may be globose, subglobose, spherical, ovoid, ellipsoid, cylindrical, elongate, ogival, apiculate, pyriform, cylindroid, lunate, triangular, or irregular and cells of *Pachysolen* may have apiculi. Pseudohyphae and branched septate or non-septate hyphae may be formed, *Metschnikowia* may form long aseptate germ tubes reminiscent of true hyphae. Hyphae of *Danielozyma* may form endoconidia, hyphal septa of *Ambrosiozyma* may have a central, plugged pore surrounded by a thickening of the wall (dolipore-like). Blastoconidia on denticles present or absent, and rarely arthroconidia may be formed. Colonies white to cream coloured, tannish-white, butyrous to hyphal, in *Kuraishia* and *Nakazawae* often mucoid due to the production of extracellular polysaccharides, with a smooth, glistening, or farinose surface. Pellicles may be formed on the surface of liquid media. *Brettanomyces*/*Dekkera* cultures grow slowly and are usually short-lived because of acetic acid production. **Physiology/biochemistry:** Fermentation of glucose and other sugars is present, weak, or absent. Xylose may be fermented, which is important for biofuel production. In *Brettanomyces/Dekkera* fermentation may be stimulated by molecular oxygen (negative Pasteur efect or Custers efect). Assimilative growth may occur with several carbohydrates, including pentoses, hexoses, disaccharides, alcohols, polyols, hexadecane, and organic acids. *Babjeviella inositovora* can utilize myo-inositol as a sole source of carbon. Nitrate assimilation is variable. Growth may require extraneous vitamins such as thiamine and biotin. Some *Ogataea* species grow at temperatures as high as 45–50 °C. The major ubiquinone, where known, is coenzyme Q-7, Q-8, and Q-9. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein family OG0000547 (Groenewald et al. [2023\)](#page-137-0) as well as phylogenetic analyses using DNA sequences encoding LSU rDNA, SSU rDNA, *tef1-α*, *rpb1*, and *rpb2*, *ACT1*, *tef2*, Mcm7 (Kurtzman and Robnett [2013b;](#page-143-3) Daniel et al. [2014](#page-134-1)).

**Life mode and distribution:** Saprobic in terrestrial and aquatic habitats, may be associated with plants, animals, particularly insects, but also warm-blooded animals including humans. Few species may be pathogenic in animals, including immune-compromised humans. While some species have a restricted distribution, *Pichiomycetes* taxa are cosmopolitan in diferent terrestrial and aquatic ecosystems. Some species are widespread in natural and anthropogenic environments, while others are specialised (Kurtzman et al. [2011;](#page-143-1) Buzzini et al. [2017](#page-132-0)). Sugar or starch-rich environments are particularly favourable substrates. Many species are associated in varying degrees with diferent plant structures such as fowers, fruits, tree fuxes, phylloplane, and animals ranging from insects to humans, or are saprophytes in soil, rotten wood, and other decaying materials, while some are reported from fruiting bodies of mushrooms. They are also reported from fermented substrates with low pH, like acid beers, wine, cocoa, and coffee. Species of the genera *Komagataella, Kuraishia, Ogataea* and *Ascoidea* show an uncommon property among yeasts: the ability to grow on methanol as a sole carbon source. *Scheffersomyces*, *Spathaspora*, and *Pachysolen* ferment xylose, the major pentose of hemicellulosic plant residues, which is a possible ecological indicator of wood decomposition. Endemic species have been recognized through extensive studies on beetle associated *Metschnikowia* species, e.g. *M. borealis* in eastern North America, *M. santaceciliae* in Central America, *M. ipomoeae*, *M. lochheadii* in the Neotropics with modern introduction to Hawaii, *M. hawaiiensis* in Hawaii, *M. continentalis* and *M. cerradonensis* in South America, *M. aberdeeniae* in Africa, and *M. hibisci* in Australia. Some species are commensals in warm-blooded animals, and may have potential as opportunistic human pathogens (Inácio and Daniel [2017\)](#page-141-3). *Candida albicans* is more frequently found in humans with and without signs of disease compared to other niches.

**Notes:** *Pichiomycetes* was introduced as one of six new classes to classify yeasts of the subphylum *Saccharomycotina* according to evolutionary divergence based on the analysis of genomic data (Groenewald et al. [2023\)](#page-137-0). The diagnosis of *Pichiomycetes* was provided as a class-specifc orthologous group of proteins. For more details, see the entry on *Saccharomycetes*. Following the nomenclatural requirement to use a single name for a single taxon, the genus *Candida,* originally used to group morphologically indistinguishable anascosporogenous yeast species, is to be reduced to a group typifed by *C. tropicalis* and situated in the *Pichiomycetes* (Daniel et al. [2014\)](#page-134-1). However, in the understanding that conclusive phylogenetic inference necessitates complete taxon sampling and genomic data, *Candida* species outside the *C. tropicalis* group and outside of *Pichiomycetes* still await their reclassifcation.

Members of *Pichiomycetes* are signifcant economically and ecologically, clinically and industrially (see Tables [12,](#page-45-0) [13](#page-45-1), and [14](#page-46-0) respectively for examples).

According to Groenewald et al. [\(2023\)](#page-137-0) the class comprises the three orders *Alaninales*, *Pichiales*, *Serinales*, accommodating fve families and 41 genera. The genus *Babjeviella* represents a highly divergent lineage in the class and was not assigned to a family.

**Type order:** *Pichiales* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 12 (2023).

**Type family:** *Pichiaceae* M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 12 (2023).

**Type genus:** *Pichia* E.C. Hansen, Centralbl. Bakteriol. Parasitenk., 2. Abth. 12: 538. (1904).

**Type species:** *Pichia membranifaciens* (E.C. Hansen) E.C. Hansen, Centralbl. Bakteriol. Parasitenk., 2. Abth. 12: 538. (1904).

*Saccharomycetes* G. Winter, Rabenh. Krypt.-Fl. Ed. 2, 1(1): 32, 68. (1884) emend. M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 13 (2023); Fig. [16n](#page-42-0)-t Index Fungorum Registration Identifer 90791.

**Sexual reproduction:** *Ascomata* absent. *Asci* unconjugated or formed by isogamous or heterogamous conjugation, oneto many-spored, spheroid ovoid or ellipsoid, persistent or deliquescent. *Ascospores* spheroid, subspheroid, ovoid, ellipsoid, cylindroid, hat-shaped, reniform, saturn-shaped, fusiform, bacilliform or acicular, aseptate or rarely 1-septate; smooth or rough; ledges, appendages, brims and terminal extensions are present or absent. Species are homothallic or

<span id="page-45-0"></span>**Table 12** Economical and ecological signifcances of *Pichiomycetes*, examples:



<span id="page-45-1"></span>



<span id="page-46-0"></span>



heterothallic. **Asexual reproduction:** White to cream coloured, greyish or yellow yeast-like or flamentous colonies. Spherical, ellipsoid, ovoid, elongated or apiculate budding cells usually present. Two layered cell wall, holoblastic, multilateral or bipolar budding. Solitary or catenate, aseptate blastoconidia present or absent. Pseudohyphae and true hyphae present or absent. Arthroconidia rarely present. **Physiology/biochemistry**: Fermentation is present or absent. Nitrate assimilation is variable. The diazonium blue B reaction is negative, urease is not produced. The major ubiquinone, where known, is coenzyme Q-5, Q-6, Q-7, Q-8 or Q-9. **Molecular analyses**: The class can be distinguished by the class-specifc protein families OG0004556 and OG0004235 (Groenewald et al. [2023](#page-137-0)).

**Life mode and distribution:** While some species have restricted distribution, the taxa assigned to *Saccharomycetes* are spread worldwide in diferent terrestrial and aquatic ecosystems (Kurtzman et al. [2011;](#page-143-1) Buzzini et al. [2017](#page-132-0)). Many species are associated with plants and inhabit, e.g. fowers and fruits, or are saprobes on decaying plant material, while some are reported from the fruiting bodies of mushrooms. *Eremothecium* species are phytopathogenic causing diseases on cotton, flax, coffee, soybean, mustard, and many other plants. They are often vectored by insects (Kurtzman and de Hoog [2011](#page-143-6)). Numerous species are saprobic in soil or dwelling in freshwater or in marine habitats. *Saccharomyces* species take an essential role in baking, brewing and

winemaking. Various species are known from fermented substrates (e.g., sauerkraut and silage) thus, important in industries (Table [15\)](#page-47-0). Several species, e.g. *Zygosaccharomyces* spp., may cause spoilage of foods and beverages. Some species are associated with warm-blooded animals, and a few, like *Nakazawaea glabratus* (*Candida glabrata*) are opportunistic human pathogens (Kurtzman et al. [2011](#page-143-1)). Three mycoparasitic genera *Endomyces*, *Helicogonium*, *Phialoascus* and an endoparasite of *Drosophila*, *Coccidiascus*, are tentatively assigned to *Saccharomycetes* as *Saccharomycetes incertae sedis* (Groenewald et al. [2023\)](#page-137-0). Most of the species assigned to these genera have not been cultivated and sequenced yet, or if cultivated (*E. scopularum*), the identity of the strains remains to be confrmed (Suh et al. [2001\)](#page-156-3).

**Notes:** The class *Saccharomycetes* was introduced by Winter (1884) for single-celled "plants" that multiply by budding and form endogen spores. Eriksson and Winka ([1997\)](#page-136-0) also proposed *Saccharomycetes* to accommodate the order *Saccharomycetales*, but it is a later homonym of Winter's class. Until recently *Saccharomycetes* has comprised all (more than 1,200) species assigned to *Saccharomycotina*. Detailed treatments of the species assigned to *Saccharomycetes* were presented in the subsequent editions of The Yeasts: A Taxonomic Study (Lodder and Kreger-van Rij [1952](#page-145-0); Lodder [1970](#page-145-1); Kreger-van Rij [1984;](#page-143-7) Kurtzman and Fell [1998](#page-143-8); Kurtzman et al. [2011](#page-143-1)). Kurtzman was among the frst to apply DNA sequence-based classifcation for *Saccharomycetes*.

<span id="page-47-0"></span>**Table 15** Industrial applications of *Saccharomycetes*



Initially, he utilized rRNA coding gene sequences (Kurtzman and Robnett [1997,](#page-143-10) [1998\)](#page-143-2), and later conducted multigene phylogenetic studies (Kurtzman [2003](#page-143-11); Kurtzman and Robnett [2003](#page-143-12), [2007](#page-143-4), [2010](#page-143-13); Kurtzman et al. [2007](#page-143-5), [2008](#page-143-14); Kurtzman and Suzuki [2010](#page-143-15); Kurtzman and Robnett [2013a,](#page-143-0) [b](#page-143-3)). In the latest decade, the classification of yeasts, including ascomycete, especially at higher ranks, has increasingly relied on genome-scale phylogenetic reconstructions (Hittinger et al. [2015](#page-140-2); Shen et al. [2016,](#page-154-0) [2018](#page-154-1), [2020\)](#page-154-2). Based on relative evolutionary divergence (RED) analyses, Li et al. [\(2021a](#page-145-2)) pointed out that the subphylum *Saccharomycotina* is under-classifed. Groenewald et al. [\(2023\)](#page-137-0) proposed a new genome-informed higher-rank classifcation for *Saccharomycotina*. They emended the diagnosis of class *Saccharomycetes*, and reassigned numerous taxa from *Saccharomycetes* to six newly erected classes; *Alloascoideomycetes*, *Dipodascomycetes*, *Lipomycetes*, *Pichiomycetes*, *Sporopachydermiomycetes* and *Trigonopsidomycetes*. They also introduced new order(s) in each class. All classes and orders were supported by taxon-specifc orthologous groups of proteins (OGs), used as diagnoses of the taxa. As summarised by Daniel et al. [\(2014\)](#page-134-1), according to phylogenetic studies, numerous *Candida* species outside the *Lodderomyces*/*Candida* clade are afliated with the class *Saccharomycetes* (sensu Groenewald et al. [2023\)](#page-137-0). Some of them, (e.g. *C. glabrata*) have already been transferred to the corresponding teleomorph genera (Takashima and Sugita [2022](#page-156-5)), while others, (e.g. *C. adriatica*) are still awaiting reassignment.

According to Groenewald et al. [\(2023\)](#page-137-0), the class currently comprises four orders; *Ascoideales*, *Phafomycetales*, *Saccharomycetales* and *Saccharomycodales*, accommodating six families and 31 genera. The order *Ascoideales* comprises taxa forming expanding mycelia, while budding cells are present or absent. *Phafomycetales* and *Saccharomycetales* are characterised by yeast cells multiplying by multilateral budding, while *Saccharomycodales* cells multiply by bipolar budding.

**Type order:** *Saccharomycetales* C. Luerssen, Grundz. Bot.: 156. 1877, emend. M. Groenew., Hittinger, Opulente & A. Rokas, Stud. Mycol. 105: 14(2023).

**Type family:** *Saccharomycetaceae* C. Luerssen, Grundz. Bot.: 160. (1877).

**Type genus:** *Saccharomyces* Meyen, Arch. Naturgesch. 4 (2): 100. (1838).

**Type species:** *Saccharomyces cerevisiae* (Desm.) Meyen, Arch. Naturgesch. 4 (2): 100. (1838).

*Sporopachydermiomycetes* M. Groenew., Hittinger, Opulente & A. Rokas Stud. Mycol. 105: 15 (2023); Fig. [16](#page-42-0)u Index Fungorum Registration Identifer 847275.

**Sexual reproduction:** Conjugation may or may not precede ascus formation. Ascospores are spheroid, subglobose, ellipsoid, or bacilliform. One to four ascospores are formed per ascus. The ascospores usually have a thick coat of refractile material that keep them together after release from the ascus at maturity. They tend to agglutinate. **Asexual reproduction:** Cell division is by multilateral budding on a narrow base. Cells are ovoid, ellipsoid or elongate, occasionally curved and occur singly, in pairs or even in small groups. Pseudohyphae and true hyphae are not formed. **Physiology/ biochemistry:** Glucose fermentation is absent or weak. Nitrate is not assimilated, myo-inositol as sole carbon source is assimilated. The major ubiquinone is coenzyme Q-9. The diazonium blue B reaction can be negative or positive. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein families OG0028621, OG0028581, OG0028722, and OG0028736 (Groenewald et al. [2023](#page-137-0)) as well as phylogenetic analyses using DNA sequences encoding LSU rDNA, mtSSU rDNA, and *COX II* sequences (Kurtzman and Robnett [2007](#page-143-4)).

**Life mode and distribution:** Cosmopolitan in distribution, on (decaying) cacti, non-cactus plant–insect interface, slimy exudates of red oaks (*Quercus rubra*, Fagaceae). A small number of *Sporopachydermiomycetes* members have primarily been reported from necrotic tissues of cacti, wood, and (sea) water (Lachance [2011b](#page-144-3)). High levels of 2-propanol and acetone are present in decaying cactus tissues. *Sporopachydermia* species can metabolise 2-propanol and acetone as carbon sources, making cactus tissues an ideal substrate.

**Notes:** The class *Sporopachydermiomycetes* was introduced by Groenewald et al. ([2023](#page-137-0)) based on the analysis of genomic data, proposed the distribution of yeasts, assigned to *Saccharomycotina,* among seven classes. They provided a diagnosis of *Sporopachydermiomycetes,* with class specifc orthologous groups of proteins (OGs). For more details see the entry *Saccharomycetes*. *Sporopachydermia* representatives form a strongly supported monophyletic clade as per LSU, MtSm and COXII analyses (Kurtzman and Robnett [2007\)](#page-143-4). Although the number of species validly described in this class is limited, Lachance [\(2011b](#page-144-3)) has listed a fair number of species with strong affinity to the genus *Sporopachydermia,* that are awaiting formal descriptions. The distribution of those phylotypes is highly infuenced by geography and less so by host-plant association (Lachance [2011b\)](#page-144-3). Groenewald et al. [\(2023](#page-137-0)) is the latest study that provides a comprehensive background on the taxonomy of the higher-level classifcation of *Sporopachydermiomycetes*. Several phylogenetic studies found that the *Sporopachydermia* clade is situated on a long branch as a sister clade to *Alloascoidea*, indicating substantial evolutionary divergence (Lachance [2011b;](#page-144-3) Kurtzman and Robnett [2013a,](#page-143-0)[b](#page-143-3); Groenewald et al. [2023\)](#page-137-0). Although a limited number of *Sporopachydermia* species are presented by the class *Sporopachydermiomycetes*, Groenewald et al. ([2023\)](#page-137-0) propose the separation of *Sporopachydermia* and its sister genus into two sister classes, *Sporopachydermiomycetes* and *Alloascoideomycetes*, respectively, due to their long branches.

**Type order:** *Sporopachydermiales* M. Groenew., Hittinger, Opulente & A. Rokas. Stud. Mycol. 105: 15 (2023).

**Type family:** *Sporopachydermiaceae* M. Groenew., Hittinger, Opulente & A. Rokas. Stud. Mycol. 105: 15 (2023). **Type genus:** *Sporopachydermia* Rodr. Mir., Antonie van Leeuwenhoek 44: 440 (1978).

**Type species:** *Sporopachydermia lactativora* Rodr. Mir., Antonie van Leeuwenhoek 44: 440 (1978).

*Trigonopsidomycetes* M. Groenew., Hittinger, Opulente & A. Rokas Stud. Mycol. 105: 16 (2023).

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**Sexual reproduction:** *Ascospores* are helical or lunate with tapering ends and arise in a bud formed by cells following isogamous conjugation. One to four *ascospores* are formed per *ascus*. Conjugation may or may not precede ascus formation. **Asexual reproduction:** The cells are either relatively small, ovoid to short cylindroid occur singly, in pairs and sometimes in short chains, or they can be arranged in thalli with a T-shaped cell. No *pseudohyphae* or *true hyphae* are produced. **Physiology/biochemistry:** Usually a narrow range of sugars are assimilated, absence of growth on nitrate as sole nitrogen source. Most simple carboxylic acids commonly tested are utilized. Only glucose can be fermented and only by some members. The major ubiquinone, where known, is coenzyme Q-9. The diazonium blue B reaction is negative. Some members can utilize 1-butanol as sole carbon source and produce extracellular proteases. **Molecular analyses:** The class can be distinguished by the following molecular characteristics. Class-specifc protein families OG0008190 and OG0008482 (Groenewald et al. [2023](#page-137-0)) as well as phylogenetic analyses using DNA sequences encoding LSU rDNA, mtSSU-rDNA, and *COX II* (Kurtzman and Robnett [2007,](#page-143-4) [2013b;](#page-143-3) Lachance and Kurtzman [2013](#page-144-4)).

**Life mode and distribution:** Members of *Trigonopsidomycetes* have a relatively wide host range (Kerrigan and Smith [2001;](#page-142-3) Kurtzman 2011; Lachance and Kurtzman [2013](#page-144-4)). They can be found in alcoholic beverages such as wine and beer, grapes and vineyards, necrotic tissue of cactus and other plants, trees, from free-living nematodes, and wood-boring beetle galleries. Many are associated with grape must, especially the *Trigonopsis* members, suggesting that *Trigonopsis* may have a distinct ecological niche that is linked to grapes. It is frequently associated with necrotic tissue of primarily cactus, although the genus *Tortisopra* has been identifed in other plants as well, most likely due to its ability to produce extracellular proteases that solubilize casein. Members of this class, particularly *Botryozyma*, have the ability to adhere to the cuticle of nematodes due to the presence of distinct T-shaped cells. Some members of this class are found in nematodes associated with beetles, and the yeast appears to have a commensal relationship, taking nutrition from the nematodes (Kerrigan and Smith [2001\)](#page-142-3).

**Notes:** The class *Trigonopsidomycetes* in the subphylum *Saccharomycotina* was introduced by Groenewald et al. ([2023](#page-137-0)), who, based on the analysis of genomic data, proposed the distribution of the yeasts assigned to *Saccharomycotina* among seven classes. They provided a diagnosis of *Trigonopsidomycetes* with class-specifc orthologous groups of proteins (OGs) (for more details see the entry *Saccharomycetes*). Members of the *Trigonopsidales* form a mildly supported monophyletic clade as per LSU, MtSm and COXII sequence analyses (Kurtzman and Robnett [2007](#page-143-4), [2013b](#page-143-3); Lachance and Kurtzman [2013\)](#page-144-4) and the phylogenomic analyses done by Shen et al. [\(2018\)](#page-154-1). Groenewald et al. [\(2023\)](#page-137-0) is the latest study that provides a comprehensive background on the taxonomy of the higher-level classifcation of *Trigonopsidales*. Phylogenetic analysis of D1/ D2 LSU rRNA gene sequences demonstrated that the two described species of *Botryozyma* are quite closely related to one another as well as to the two known species of *Ascobotryozyma* (Kerrigan and Smith [2001](#page-142-3)), and thus represent a single genus in the new single nomenclature system. The Melbourne Code assigns equal weight to anamorphic and teleomorphic genus names. Because the genus *Botryozyma* (1992) has priority of date over *Ascobotryozyma* (2001), Lachance and Kurtzman ([2013](#page-144-4)) propose assigning the species of this clade to the genus *Botryozyma*. Considering the exceptional early divergence of the basal branching point marking this order, additional data may eventually support the division of this new order. However, due to the lack of additional species, Groenewald et al. ([2023](#page-137-0)) have decided to combine the genera *Trigonopsis*, *Tortispora*, and *Botryozyma* into one order, while acknowledging that future descriptions of more species in this group may lead to the split of *Tortispora* and *Botryozyma* from *Trigonopsis* into a separate class.

The economic and ecological signifcance of members of this class is unknown, as is whether they have any application in the medical/clinical feld. However, the ability of *Trigonopsis variabilis* to produce D-amino acid oxidases that catalyse the oxidative deamination of α-amino acids to α-keto acids and ammonia (Kurtzman 2011) is of great importance to the pharmaceutical industry. Other uses for this enzyme include the synthesis of chiral intermediates to produce antihypertensive drugs (Patel [2001\)](#page-149-2) and the conversion of cephalosporin C to 7-amino cephalosporanic acid, which is a key intermediate for cephem antibiotics (Dib and Nidetzky [2008\)](#page-135-4).

According to Groenewald et al. ([2023](#page-137-0)), currently, the class comprises one order *Trigonopsidales*, one family *Trigonopsidaceae* and three genera (*Botryozyma*, *Tortispora* and *Trigonopsis*).

**Type order:** *Trigonopsidales* M. Groenew., Hittinger, Opulente & A. Rokas. Stud. Mycol. 105: 16 (2023).

**Type family:** *Trigonopsidaceae* Lachance & Kurtzman, Int. J. Syst. Evol. Microbiol. 63: 3113 (2013).

**Type genus:** *Trigonopsis* Schachner, Z. Ges. Brauwesen 52: 137 (1929).

**Type species:** *Trigonopsis variabilis* Schachner, Z. Ges. Brauwesen 52: 137 (1929).

**Subphylum** *Taphrinomycotina* O.E. Erikss. & Winka, Myconet 1(1): 11 (1997).

Index Fungorum Registration Identifer 501471.

**Notes:** Eriksson and Winka ([1997](#page-136-0)) introduced *Taphrinomycotina* to accommodate *Neolectomycetes*, *Pneumocystomycetes* (as *Pneumocystidomycetes*), *Schizosaccharomycetes* and *Taphrinomycetes*. Rosling et al. ([2011](#page-152-2)) and Čadež et al. [\(2021](#page-132-6)) introduced *Archaeorhizomycetes* and *Novakomycetes* respectively thus *Taphrinomycotina* comprises six classes.

*Archaeorhizomycetes* Rosling & T.Y. James, Science, N.Y. 333(no. 876): 879 (2011).

Index Fungorum Registration Identifer 561762.

**Morphology:** The life cycle of *Archaeorhizomycetes* is still poorly understood, and no sexual reproductive structures have yet been observed. In culture on MMN medium, the growth edges in both *Archaeorhizomyces borealis* and *A. fnlayi* are characterised by thin (0.2–2 µm in diameter) hyphae. Pale aerial hyphae are produced in the older parts of the colonies (Fig. [17\)](#page-50-0) (Rosling et al. [2011;](#page-152-2) Menkis et al. [2014](#page-147-2)). *Archaeorhizomyces fnlayi* grows faster in culture than *A. borealis* (Menkis et al. [2014](#page-147-2)). In *A. fnlayi*, the grouping of thin hyphae into chord-like bundles is common, a growth that is not observed in *A. borealis* (Fig. [17](#page-50-0)b). Based on TEM, both species form simple septa lacking pores, and some of the hyphal cell-walls are double layered and ca. 150–200 nm thick (Rosling et al. [2011](#page-152-2); Menkis et al. [2014](#page-147-2); Fig. [17c](#page-50-0),e,f). In culture, *A. fnlayi* hyphae are extensively coated by extracellular materials (Rosling et al. [2011](#page-152-2)), and the hyphal surface appears rough both in SEM and TEM (Fig. [17](#page-50-0)c). The hyphal cells have one or two nuclei (Fig. [18](#page-51-0)). Both species produce chlamydospore-like swellings in the older parts of the colony (Fig. [17](#page-50-0)). The swellings are larger and more numerous in *A. borealis* than in *A. fnlayi* (Rosling et al. [2011;](#page-152-2) Menkis et al. [2014](#page-147-2); Fig. [17](#page-50-0)). The large and old swellings are thick-walled, and the cell-wall is multi-layered (Fig. [17\)](#page-50-0). They can contain multiple nuclei (Menkis et al. [2014](#page-147-2)) (Fig. [18](#page-51-0)j-l). In addition to the large swellings, smaller terminal and intercalary swellings are produced in both species, delimited by septa (Fig. [17](#page-50-0)[,18\)](#page-51-0). *A. borealis* has more swellings and a broader variation in colony morphology than *A. fnlayi*. In addition, these swollen cells can be devoid of nuclei (Fig. [18](#page-51-0)b-c). Menkis et al. ([2014](#page-147-2)) observed that *A. borealis* grows as yeast on b-cyclodextrin amended medium.

**Life modes and distribution:** Environmental DNA (eDNA) sequences taxonomically assigned to *Archaeorhizomycetes* are frequently observed in samples of soil from forest and woodland habitats. Interestingly, the class was underrepresented across root and rhizosphere samples suggesting a weak association with plants at the class level. Individual

<span id="page-50-0"></span>**Fig. 17** SEM and TEM images of *Archaeorhizomyces fnlayi* and *A. borealis* grown on MMN. **a** Terminal and intercalary swellings and hyphae of *Archaeorhizomyces fnlayi*. **b** Chordlike hyphal bundles typical for *Archaeorhizomyces fnlayi*. **c** *Archaeorhizomyces fnlayi* hyphae with simple septa and uneven hyphal surface. **d** Terminal and intercalary swellings in *Archaeorhizomyces borealis* covered by the extracellular material (arrow). **e** *Archaeorhizomyces borealis* hyphae with simple septa without pores. **f** *Archaeorhizomyces borealis* hyphae with double cell-wall (arrow) and swellings with lipid-like bodies (L). Scale 1 μm. Scale bars **a**, **d**= 10 μm, b=3 μm, **d**, **e**= 300 nm



taxa in the class may, of course, respond diferently, as suggested by the two described species that were both cultured from roots. Further, an eDNA metabarcoding of pot cultures with *Arabidopsis* found that three *Archaeorhizomycetes* OTUs had the highest relative abundance in the rhizosphere compared to endosphere and bulk soil samples (Urbina et al. [2018](#page-157-2)). Based on the two described species that have been studied in culture it is known that while associated with roots

these fungi do not form characteristic mycorrhizal structures but can colonize roots without causing harm to the roots (Rosling et al. [2011\)](#page-152-2). Further, the two described species have broad saprotrophic potential, as indicated by their ability to grow on a range of complex carbon sources, with a predominant occurrence in the rhizosphere soil (Menkis et al. [2014](#page-147-2)). These growth patterns are consistent with sequences from the class being more frequently observed in soil samples

<span id="page-51-0"></span>**Fig. 18** Confocal laser scanning images of *Archaeorhizomy ces fnlayi* and *A. borealis*. Calcofuor white stained septa and cell walls (cyan) and propidium iodine stained of nuclei (magenta) ( **a** *A. fnlayi* hyphae with two nuclei per cell. **b** Swollen hyphae of *A. fnlayi* with often two nuclei per cell. Notice that some of the swollen cells are empty of nuclei. **c** Swollen hyphae of *A. fnlayi* with a small intercalary swell ing (arrow) with two nuclei. **d** *A. borealis* hyphae with one nucleus. **e** Multiple consecu tive swellings in *A. borealis* hyphae with one or two nuclei. **f** Intercalary *A. borealis* swelling with one nucleus). Scale bars **a**–**f**=5 μm



compared to other types of samples. It is likely that many taxa in *Archaeorhizomycetes* are free-living soil fungi and that species display diferent degrees of root association.

*Archaeorhizomycetes* is globally distributed, i.e., sequences taxonomically assigned to the class have been detected in 18% of all samples currently contained in the globalfungi.com database and include records from all continents (Větrovský et al. [2020\)](#page-158-3) and sequences of *Archaeorhizomycetes* are frequently observed in environmental soil and root samples (Schadt et al. [2003;](#page-153-1) Porter et al. [2008;](#page-150-1) Rosling et al. [2011](#page-152-2), [2013](#page-152-3)). Based on sequence similarity and clustering of published eDNA amplicon sequences, the class has been estimated to comprise at least 500 species (Menkis et al. [2014\)](#page-147-2). However, estimating species richness from eDNA metabarcoding data is not trivial and global richness estimates based on clustering at 98.5% of environmental sequences available as Short Read Archive data has suggested that *Archaeorhizomycetes* may encompass as many as 16,231 taxa (Hawksworth and Lücking [2017](#page-139-0)). The authors emphasise that the latter number is likely infated by a combination of sequencing errors and artifacts of the clustering method. More moderate estimates include the gbif.org database, which currently includes 17,131 occurrences and 767 species hypotheses for *Archaeorhizomycetes*. The curated fungal sequence database UNITE (version 9.0, Nilsson et al. [2018](#page-148-3)) currently has 765 species hypotheses classifed in the class *Archaeorhizomycetes*, based on its 1.5% sequence dissimilarity threshold. Of these, 126 are also recovered in the globalfungi.com database, which currently includes fungal eDNA amplicon data from 515 studies (Větrovský et al. [2020](#page-158-3)). Local species richness can be high, with records of 278 *Archaeorhizomyces* hypothesised species in an alpine habitat (Pinto-Figueroa et al. [2019](#page-150-2)), 80 species recovered in a subtropical evergreen shrub bog (Tian et al. [2021\)](#page-157-3) and 68 hypothesised species detected in a Swedish boreal forest site (Kalsoom Khan et al. [2020\)](#page-142-4). Irrespective of the actual species richness, the two species that have been cultured, formally described, and named, *Archaeorhizomyces borealis* Menkis et al. and *Archaeorhizomyces fnlayi* Rosling & T. James, certainly represent only a fraction of the true diversity in the class. Despite repeated efforts, no additional species have been successfully captured in pure culture, but two additional species, *A. secundus* and *A. victor*, have been proposed based on phylogenetic and ecological evidence from eDNA amplicon data (Kalsoom Khan et al. [2020\)](#page-142-4).

**Type order:** *Archaeorhizomycetales* Rosling & T.Y. James, Science, N.Y. 333(no. 876): 879 (2011).

**Type family:** *Archaeorhizomycetaceae* Rosling & T.Y. James, Science, N.Y. 333(no. 876): 879 (2011).

**Type genus:** *Archaeorhizomyces* Rosling & T.Y. James, Science, N.Y. 333(no. 876): 879 (2011).

**Type species:** *Archaeorhizomyces fnlayi* Rosling & T.Y. James, Science, N.Y. 333(no. 876): 879 (2011).

## *Neolectomycetes* O.E. Erikss. & Winka, Myconet 1(1): 8 (1997).

Index Fungorum Registration Identifer 90270; Fig. [19](#page-53-0)

*Ascomata* long, irregularly clavate, lanceolate or spathulate and consisting of a sterile zone at the bottom and a fertile zone (hymenium) on the top, yellow to bright yellowish, smooth, irregularly lobate. *Asci* cylindrical to cylindricalclavate, 8-spored. *Paraphyses* absent. *Ascospores* uniseriate, unicellular,  $5.5-8 \times 3-4$  µm, reniform, ellipsoid or ovoid, hyaline, smooth (Claudiu Chinan and Hewitt [2012](#page-133-1)).

**Life mode and distribution:** Associated with rootlets of conifers. Distribution is cosmopolitan.

**Notes:** Eriksson and Winka ([1997\)](#page-136-0) introduced this class to accommodate *Neolecta* Speg., typifed by *Neolecta favovirescens* Speg. Currently, the class comprises a single order, a family and a genus. Index Fungorum (2023) has listed four epithets (accession date: 29.06. 2023). However, Landvik et al. [\(2001](#page-144-5)) have accepted only two species (viz*., N. vitellina* (Bres.) Korf & J.K. Rogers 1971 and *N. irregularis* (Peck) Korf & J.K. Rogers 1971) except the type species.

The placement of *Neolecta* has been problematic for the last three decades. Based on *RPB2* sequence data, Liu et al. [\(1999](#page-145-3)) showed that *Neolecta vitellina* can be accommodated in the basal lineage of the *Ascomycota,* sister to *Schizosaccharomyces pombe* Lindner. Furthermore, Liu et al. ([1999\)](#page-145-3) stated that 'the ancestral ascomycetes may have been flamentous fungi with a sexual phase that produced a fruiting body'. The clade in which both taxa resided was named 'Archiascomycetes' but this was not a proper rank (See editorial comment in the Index Fungorum 2023). Landvick et al. (2001), however, showed that *Neolecta* is phylogenetically close to yeast taxa in *Saccharomycotina* and *Taphrinomycotina*. Schoch et al. [\(2009\)](#page-153-2) confirmed the placement of *Neolecta* in *Taphrinomycotina.*

**Type order:** *Neolectales* Landvik, O.E. Erikss., Gargas & P. Gust., Syst. Ascom. 11: 107–118 (1993).

**Type family:** *Neolectaceae* Redhead, Can. J. Bot. 55: 305 (1977).

**Type genus:** *Neolecta* Speg., Anal. Soc. cient. argent. 12(5): 218 (1881).

**Type species:** *Neolecta favovirescens* Speg. [as 'favo-virescens'], Anal. Soc. cient. argent. 12(5): 218 (1881).

*Novakomycetes* Dlauchy, G. Péter & Čadež, Microorganisms 9 (2, no. 301): 14 (2021).

Index Fungorum Registration Identifer 838640; Fig. [20](#page-54-0)



<span id="page-53-0"></span>**Fig.19** *Neolecta irregularis*: **a** Ascomata. **b** Asci. **c** Ascospores. **d** Cells of stipe. Scale bars: **A**=10 mm; **B**=20 µm; **C**–**D**=10 µm. (Photograph 'a' with the courtesy of Bau Tolgor. b-d was redrawn from Jin-Zhao and Tolgor 2009)

**Sexual reproduction:** *Ascomata* absent. *Asci* unconjugated or formed by heterogamous conjugation, one- or two-spored, ellipsoid, deliquescent. *Ascospores* subspheroid or ellipsoid, aseptate. **Asexual reproduction:** White to cream coloured yeast colonies formed by budding cells. Two layered cell walls, enteroblastic budding, solitary, aseptate blastoconidia. Pseudohyphae and true hyphae absent.

**Life mode and distribution:** Budding yeast originating from olive oil in Spain.

**Notes:** The class *Novakomycetes* was introduced by Čadež et al.  $(2021)$  $(2021)$  $(2021)$ . Currently, it is the smallest among the six classes assigned to the subphylum *Taphrinomycotina* (Wijayawardene et al. [2022a\)](#page-160-3). According to phylogenomic

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analyses, *Novakomycetes* is closely related to *Schizosaccharomycetes* comprising fission yeasts (Čadež et al. [2021\)](#page-132-6). The class includes a hierarchic chain of monotypic taxa: *Novakomycetales*, *Novakomycetaceae* and *Novakomyces* erected to accommodate the species *Novakomyces olei* (Čadež et al. [2021](#page-132-6)).

The single species assigned to *Novakomycetes* is budding yeast originating from olive oil; however, it may be allochthonous in this habitat, as no additional report of its occurrence has been published (Čadež et al. [2021](#page-132-6); Zullo and Ciafardini [2022\)](#page-162-0).

**Type order:** *Novakomycetales* Dlauchy, G. Péter & Čadež, in Čadež, Dlauchy, Tome & Péter, Microorganisms 9(2, no. 301): 14 (2021).



**Fig. 20** Micromorphological characters of *Novakomyces olei* (from Čadež et al. [2021](#page-132-6)). **a** Budding cell of *Novakomyces olei*. Collarette surrounding the forming bud is indicated by arrows. **b** Ascosporulat-

<span id="page-54-0"></span>**Type family:** *Novakomycetaceae* Dlauchy, G. Péter & Čadež, in Čadež, Dlauchy, Tome & Péter, Microorganisms 9(2, no. 301): 14 (2021).

**Type genus:** *Novakomyces* Dlauchy, G. Péter & Čadež, Microorganisms 9 (2, no. 301): 14 (2021).

**Type species:** *Novakomyces olei* Dlauchy, G. Péter & Čadež, in Čadež, Dlauchy, Tome & Péter, Microorganisms 9(2, no. 301): 14 (2021).

*Pneumocystomycetes* O.E. Erikss. & Winka [as 'Pneumocystidomycetes'], Myconet 1(1): 9 (1997).

Index Fungorum Registration Identifer 91007.

Non-flamentous yeast-like organisms, aggregated and parasitic cells in mammalian alveoli, thin-coated, uninucleate, divided by fission. *Asci* crassitunicate, tetra- or octosporum. *Ascospores* globose or round, 8-spored, thickwalled (Eriksson and Winka [1997](#page-136-0); Cushion et al. 2004).

**Life mode and distribution:** Parasitic on mammals as they aggregate in mammalian alveoli and *P. carinii* causes pneumonia (PCP) in immunocompromised (esp. AIDS) patients (Mills [1986](#page-147-3); Kirk et al. [2008\)](#page-142-0) and cancer patients (Sepkowitz et al. [1992;](#page-154-4) Sepkowitz [1993\)](#page-154-5).

**Notes:** The genus *Pneumocystis* P. Delanoë & Delanoë was introduced by Delanoë and Delanoë ([1912](#page-135-5)). For almost eight decades, *Pneumocystis* was regarded as a protozoan taxon (Stringer [2002\)](#page-156-6) but Edman et al. ([1988\)](#page-136-4) confrmed that *Pneumocystis* is related to *Saccharomyces cerevisiae* ing culture of *Novakomyces olei*. The remnant of an ascus formed by heterogamous conjugation is indicated by arrow. Scale bars:  $\mathbf{a} = 1 \, \mu \text{m}$ ,  **(The image 'a' was taken by Bence Rácz)** 

using rRNA. Later, Stringer et al. [\(1989](#page-156-7)) confrmed that the members of *Pneumocystis* are related to *Candida albicans*, *Neurospora crassa*, and *Cryptococcus neoformans* as well.

Eriksson ([1994\)](#page-136-5) introduced the family *Pneumocystaceae* O.E. Erikss. and the order *Pneumocystales* O.E. Erikss. to accommodate *Pneumocystis.* Later, Eriksson and Winka ([1997\)](#page-136-0) introduced the class *Pneumocystomycetes.*

*Pneumocystomycetidae* P.M. Kirk, P.F. Cannon, Stalpers & J.C. David.

Index Fungorum Registration Identifer IF 558745.

=*Pneumocystomycetidae* P.M. Kirk, P.F. Cannon, Stalpers & J.C. David, Ainsworth & Bisby's Dictionary of the Fungi, Edn 9 (Wallingford): 417 (2001); Nom. inval., Art. 38.1(a) (Shenzhen).

Index Fungorum Registration Identifer 91006.

Description: The description is identical to the Class description.

**Notes:** *Pneumocystomycetidae* P.M. Kirk et al. 2001 was invalidly published (Art. 38.1(a) (Shenzhen)). Here we validate *Pneumocystomycetidae*.

**Type order:** *Pneumocystales* O.E. Erikss. [as 'Pneumocystidales'], Syst. Ascom. 13(2): 170 (1994).

**Type family:** *Pneumocystaceae* O.E. Erikss. [as 'Pneumocystidaceae'], Syst. Ascom. 13(2): 170 (1994).

**Type genus:** *Pneumocystis* P. Delanoë & Delanoë, C. r. hebd. Séanc. Acad. Sci., Paris 155: 660 (1912).



<span id="page-55-0"></span>**Fig. 21** Microphotographs of diferent *Schizosaccharomyces* species. **a** Ascus of *S. pombe* with 4 ascospores (nuclei stained with dapi). **b** Ascus of *S. lindneri* with 8 ascospores. **c** Dividing cell of *S. pombe*.

**Type species:** *Pneumocystis carinii* P. Delanoë & Delanoë, C. r. hebd. Séanc. Acad. Sci., Paris 155: 660 (1912).

*Schizosaccharomycetes* O.E. Erikss. & Winka, Myconet 1(1): 10 (1997).

Index Fungorum Registration Identifer 90345; Fig. [21](#page-55-0)

All species are haplontic and homothallic. After mating type switching, clonal haploid cells of opposite mating type may conjugate and form a zygote. The diploid phase is restricted to the zygote, which promptly undergoes meiosis upon its formation. Asci of *Schizosaccharomyces pombe* contain 4 spores whereas the asci of all other species contain usually contain 8 spores. *S. cryophilus* is described as being very variable in the number of spores per ascus, which ranges from 1 to 8. Spores are round, ovoid, drop-shaped or reniform.

**Other characteristics:** Asexual reproduction is by polar growth and fssion. While all other species solely live as

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**d** Dividing cell of *S. lindneri.* Cell division instead of budding is one of the characteristic features that diferentiates the *Schizosaccharomycetes* from the *Saccharomycetes*

unicellular organisms *Schizosaccharomyces japonicus* and *S. versatilis* may switch to hyphal growth. Cell walls mainly consists of  $(1\rightarrow 3)$ -α-glucan with some  $(1\rightarrow 4)$  linkages,  $(1\rightarrow 3)$ -β-glucan with some  $(1\rightarrow 6)$ -β-linkages and galactomannans (Bush et al. [1974;](#page-132-7) Sietsma and Wessels [1990](#page-155-3)). Additionally, a small amount (0.5%) of glucosamine was detected (Sietsma and Wessels [1990\)](#page-155-3). In the glycoprotein the sugars are predominantly O-linked to the peptide moiety (de Mora et al. [1990\)](#page-135-6).

The major ubiquinone is either coenzyme Q9 or Q10 (Yamada et al. [1973](#page-161-3); Yamada and Banno [1987](#page-161-4)). In *Schizosaccharomyces japonicus* in a certain proportion of the strains no coenzyme was detected.

Glucose and other sugars are fermented. Urease is produced. Nitrate is not assimilated. Diazonium Blue B reaction is negative. Starch and acetic acid are not produced.

**Life modes and distribution:** All *Schizosaccharomyces* species live saprobic in terrestrial habitats. Pathogenicity

was not observed. *Schizosaccharomyces japonicus* and *S. versatilis* are most frequently isolated from forest materials while the rest of the species seem to depend on honey bees and/or solitary bees. Occasionally, *S. pombe, S. japonicus* and *S. versatilis* are present on fresh and rotting fruit. On dried fruit like raisins or dried mango pieces or dried pineapple pieces *S. octosporus* and *S. pombe* were frequently detected. In some manmade fermentative environments (kombucha, rum production or cacao bean fermentation), *S. pombe* plays an important role. *S. octosporus* seem to occur more frequently in the tropical and subtropical zones while others like *S. osmophilus* seem to be well adapted to cold climatic conditions (Brysch-Herzberg et al. [2022](#page-132-8)). The ecology of *S. cryophilus* is unknown as just one strain was isolated by now and its origin is rather unclear.

*Schizosaccharomyces japonicus* and *S. versatilis* do not exhibit a special osmotolerance while *S. octosporus, S. osmophilus* and *S. lindneri* are among the most osmotolerant yeast species. In terms of osmotolerance, *S. pombe* has an intermediate position.

**Notes:** Eriksson et al. ([1993](#page-136-6)) introduced the order *Schizosaccharomycetales* O.E. Erikss. et al. to accommodate the family *Schizosaccharomycetaceae* Beij. ex Klöcker. Eriksson and Winka [\(1997](#page-136-0)) introduced the class *Schizosaccharomycetes.* The phylogenetic placement was subject to a longlasting discourse. Although it was early concluded that the fssion yeasts diverged early from a common ancestor of the eukaryotes it could not be answered unambiguously if they are archaic ascomycetes which diverged early from the *Ascomycota* or if they diverged later after the *Ascomycota* from the lineage that leads to the Metazoa (Sipiczki 1989; Sipiczki 1995). Only after whole genome data were available open questions concerning the phylogeny of the fission yeast could be answered with higher confdence. Liu et al. ([2009](#page-145-4)) confrmed the placement of *Schizosaccharomyces* (the representative genus of *Schizosaccharomycetes*) in the subphylum *Taphrinomycotina*. Currently, the class comprises one order, one family which only contains the genus *Schizosaccharomyces*. It was proposed to divide the genus *Schizosaccharomyces* in three genera: *Schizosaccharomyces*, *Octosporomyces* (Kudrjawzew 1960) and *Hasegawaea* (Yamada and Banno [1987\)](#page-161-4). This proposal was never broadly accepted by later authors. The genus *Schizosaccharomyces* currently comprises seven species (i.e. *S. japonicus, S. versatilis, S. cryophilus, S. pombe, S. octosporus, S. lindneri* and *S. osmophilus*).

## **Type order:** *Schizosaccharomycetales* O.E. Erikss., Syst. Ascom. 11: 146 (1994).

**Type family:** *Schizosaccharomycetaceae* Beij. ex Klöcker, Lafar's Handb. Techn. Mykol., Edn 2 4: 189 (1905).

**Type genus:** *Schizosaccharomyces* Lindner, Wochenschr. Brau. 10: 1298 (1893).

**Type species:** *Schizosaccharomyces pombe* Lindner, Wochenschr. Brau. 10: 1298 (1893).

*Taphrinomycetes* O.E. Erikss. & Winka, Myconet 1(1): 11 (1997).

Index Fungorum Registration Identifer 90366; Fig. [22](#page-57-0)

**Sexual morph:** *Mycelium* enlarged, multinucleate. *Ascogenous cells* intercalarily as a bulge on the cell of the intercellular mycelium, spherical to roughly spherical (lemon and globose form). *Exosporium and mesosporium* smoothwalled. *Endosporium* thickness, three layers, with short pedicle-like basal appendages. **Asexual morph**: *Cells* ovoid to ellipsoidal or cylindrical, with few globose cells. *Budding* unipolar or bipolar budding cells. No hyphae and pseudohyphae (Bacigálová [2008\)](#page-130-2).

**Life modes and distribution:** The members of *Taphrinomycetes* can be saprobic or pathogenic on plants, soil, insect frass broad range of terrestrial and distributed in a broad range of ecosystems (worldwide) and are reported as diferent life modes. Phytopathogenic members of the class are responsible for causing galls on stems or inforescences, leaf curls and witches' brooms (Mix [1949](#page-148-4); Fonseca and Rodrigues [2011](#page-137-6)). Some members of *Taphrinomycetes* are detected as endophytic fungi in ITS rDNA libraries (Jumpponen and Jones [2009](#page-141-6), [2010](#page-141-7); Cordier et al. [2012\)](#page-134-4).

**Notes:** The class *Taphrinomycetes* was introduced by Eriksson and Winka [\(1997\)](#page-136-0), with *Taphrinales* as the type order. *Taphrinales* comprises two families viz*.*, *Taphrinaceae* and *Protomycetaceae*. *Protomycetaceae* has fve genera (i.e. *Protomyces, Protomycopsis, Saitoella, Taphridium,* and *Volkartia*) while *Taphrinaceae* has only one genus (*Taphrina*) (Wijayawardene et al. [2022a\)](#page-160-3). The type genus of *Protomycetaceae* (*Protomyces*) was frst established by Unger (Kurtzman 1833), and *P. macrosporus* as a type species. This genus has been strictly defned previously, based on morphology and the host range (*Compositae* and *Umbelliferae*) (Reddy and Kramer [1975;](#page-151-2)). Later, Wang et al. [\(2016a,](#page-159-1) [b\)](#page-159-2) found out that *Protomyces* species can be detected in other host plant families. The type genus of *Taphrinaceae* (*Taphrina*) was introduced by Fries [\(1815](#page-137-7)) to accommodate *T. populina* as a type species of the genus, which has a dimorphic lifestyle. The parasitic counterparts on plants are known as teleomorphs, and the anamorphic yeast state is known as the saprotrophic state and is classifed in the genus *Lalaria* (Mix [1949;](#page-148-4) Nutaratat et al. [2022](#page-149-3)). In 2003, Rodrigues and Fonseca [\(2003](#page-151-3)) studied the *Taphrina* species based on DNA sequence by using the LSU rRNA gene (D1/D2 domains) and ITS regions to confrm the placement of the genus *Taphrina* and <span id="page-57-0"></span>**Fig. 22 a** Leaf spots and galls on *Laserpitium latifolium* caused by *Protomycesmacrosporus*. **b**, **c**Young stage of ascogenous cells. **d**, **e** Mature ascogenous cells. Scale bar: **b**–**e**=50 μm (Redraw from Bacigálová et al. 2008)



found that *Taphrina* is closely related to the genus *Protomyces*. Later, the reclassifcation of four *Lalaria* species (*L. arrabidae, L. inositophila, L. kurtzmanii,* and *L. veronaerambellii*) was proposed and synonymized under *Taphrina*, according to the International Code of Nomenclature for Algae, Fungi, and Plants.

**Type order:** *Taphrinales* Haeckel, Systemat. Phylogen. Protist. Pfanzen, Entwurf natürl. Syst. Organism. Grund ihrer Stammesg. 1: 316 (1894).

**Type family:** *Taphrinaceae* Gäum., in Gäumann & Dodge, Comp. Morph. Fungi (London): 161 (1928).

**Type genus:** *Taphrina* Fr., Observ. mycol. (Havniae) 1: 217 (1815).

**Type species:** *Taphrina populina* Fr., Observ. mycol. (Havniae) 1: 217 (1815).

*BASIDIOBOLOMYCOTA* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Index Fungorum Registration Identifer 90741.

Mycelium is widely distributed, thick, highly branched, and soon becomes divided by septa. Conidiophores are erect, undivided, and swollen in the upper part into a thick elliptical club (basidium), on the top of which the conidium is formed. At maturity, the basidium cytoplasm forms the fruiting body at its lower part and is propelled by the ejecting plasma together with the still-attached conidium. Later, the conidium is separated from the basidium by the protrusion of a sharp column. Resting spores are formed on the mycelium by the copulation of two adjacent cells that initially grow up into beak-like projections on the common septum. The resting spores are crowned by these beaks.

**Life modes and distribution:** Saprobic and pathogenic widespread.

**Note:** The phylum *Basidiobolomycota* was proposed by Doweld ([2001\)](#page-135-7). *Basidiobolomycota* encompasses certain fungi that were traditionally classifed within the phylum '*Zygomycota*' but were reclassifed due to molecular and phylogenetic evidence (Doweld [2001](#page-135-7)). The classifcation was revised to account for monophyly and divergence time, as presented in studies conducted by Tedersoo et al. ([2018\)](#page-157-4) and Wijayawardene et al. ([2018b](#page-160-4)). Members of *Basidiobolomycota* are characterized by their unique features, particularly in their reproductive structures. They produce spores on specialized structures called basidia, which are club-shaped cells that typically bear four spores at their tips. This mode of spore production is one of the defning characteristics of the phylum and distinguishes it from other fungal groups. One of the remarkable traits of *Basidiobolomycota* is the production of multinucleate cells, known as coenocytic hyphae. Unlike most fungi, which have septate hyphae with distinct cell walls separating individual compartments, the hyphae of *Basidiobolomycota* lack septa, and multiple nuclei can exist within a single hyphal compartment (Tedersoo et al. [2018](#page-157-4); Wijayawardene et al. [2018b\)](#page-160-4).

**Controversial opinions:** Humber ([2012](#page-140-3)) established *Entomophthoromycota*, to accommodate *Basidiobolomycetes*, *Neozygitomycetes* and *Entomophthoramycetes*. This placement was accepted by Gryganskyi et al. [\(2013\)](#page-138-1). In the phylogenomic analyses of Galindo et al. ([2021](#page-137-8)), the representative taxa of *Basidiobolomycota* clustered in *Zoopagomycota*. Strassert and Monaghan ([2022a,](#page-156-8) [b\)](#page-156-9) also confrmed the results of Galindo et al. ([2021](#page-137-8)) and regarded *Basidiobolomycota* as a subphylum of *Zoopagomycota*, i.e. *Basidiobolomycotina*.

*Basidiobolomycota* comprises one subphylum (*Basidiobolomycotina*) and one class (*Basidiobolomycetes*).

*Basidiobolomycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001); Fig. [23](#page-59-0)

Index Fungorum Registration Identifer 90740.

Cells in their vegetative state possess a single nucleus and can manifest either as a septate mycelium or as yeastlike cells that originate from the insides of preceding cells, exemplifed by the 'Darmform' growth pattern. During mitosis, the nuclear membrane starts by breaking into pieces. These fragments, alongside other membranes, accumulate around a specifc nuclear area. The chromosomes, which are numerous and minuscule, get compacted and position themselves on a central metaphase plate. This plate typically embeds within the nucleolus, working in conjunction with a barrel-shaped spindle. During the interphase, these chromosomes relax and expand. The cell responsible for producing conidia, known as the conidiophore, is straightforward but showcases a distinct bulbous enlargement at its tip, directly below the emerging conidium. Conidia are spherical, with a single nucleus, and have a tiny conical outgrowth at their base. This outgrowth juts into the spore body but fips out during release. The wall layers of the conidia are indistinguishable. The release of these conidia is quite forceful, resembling a rocket's propulsion, which is triggered by a rupture in the central frail region of the sub-conidial enlargement. The upper part of this enlargement is released with the conidium and can occasionally separate during its movement. If secondary conidia are produced, they are typically elongated, sometimes bent, and may have a mucous drop at their tip. These are formed at the top of a slender, long conidiophore and are dispersed passively. Resting spores, generally zygospores, are developed within the axis of preceding cells. The nuclei of the gametangial cells divide before merging, but only a single nucleus from each cell becomes part of the zygospore. Once matured, zygospores exhibit robust, double-layered walls. Their germination directly results in a germ conidium, typically resembling the secondary type, which is elongated and passively released from a thin conidiophore (modifed description from Humber [2012\)](#page-140-3).

**Life modes and distribution:** Although *Basidiobolomycetes* are mostly known for their saprophytic lifestyle, there are a few species that can cause rare infections in humans and animals. *Basidiobolus ranarum* is the most well-known species responsible for infections, which are known as basidiobolomycosis. These infections often affect the subcutaneous tissue, gastrointestinal tract, or rhinofacial regions (Eidam 1886; Benjamin [1979\)](#page-131-3).

**Notes:** Similar to notes in the phylum.

**Type order:** *Basidiobolales* Jacz. & P.A. Jacz. (1931), Opredelitel' Gribov, I Ficomiţeti (Leningrad)**:** 8 (1931).

**Type family:** *Basidiobolaceae* Engler & Gilg (1924), Syllabus, Edn 9 & 10 (Berlin): 45 (1924).

**Type genus:** *Basidiobolus* Eidam 1886 [Amphoromorpha Thaxter 1914], Krypt.-Fl. Schlesien (Breslau) 3.1(9–16): 224 (1886) [1889].

**Type species:** *Basidiobolus ranarum*, Krypt.-Fl. Schlesien (Breslau) 3.1(9–16): 225 (1886) [1889].

*BASIDIOMYCOTA* R.T. Moore, Bot. Mar. 23(6): 371 (1980).

Index Fungorum Registration Identifer 90050.

*Basidiomycota* is a very large and diverse phylum of fungi, second to *Ascomycota* in species number (He et al. [2019](#page-139-1)). Many species produce macroscopic sexual



<span id="page-59-0"></span>**Fig. 23** Culture and micromorphological characteristics of *Basidiobolus omanensis* (CBS 146282). **a**, **b** Colony after 7 days of incubation at 28 °C. **a** MEA front. **b** PDA front. **c** Primary conidia. **d** Primary conidia arising from mycelia. **e** Young zygospore. **f** Zygospores.

**g** Mature conidia. **h** Capilloconidium. **i** Conidia with sharp-pointed papillae projection with beak. **j** Smooth mature sexual zygospore. **k** Numerous primary conidia. Scale bars **c**–**k**=10 µm

reproductive organs known as fruiting bodies, basidiocarps, or basidiomata. Karyogamy and meiosis occur in specialized structures (basidia), resulting in mostly four (one to twelve) haploid basidiospores. Basidia are aseptate or may be longitudinally, oblique, or transversely (partially) septate. Other important diagnostic characteristics include dolipore septa in several lineages such as *Agaricomycetes* (a barrelshaped cell wall swelling) between hyphal cells, and the presence of clamp connections at hyphal septa (although not all representatives produce clamp connections). The phylum includes mushroom-forming fungi, bracket fungi, pufballs, earthstars, stinkhorns, false truffles, and jelly fungi, but also

a number of microscopic fungi, such as rusts, smuts, and yeasts (Zhao et al. [2017\)](#page-161-5). Some species with feshy fruiting bodies are traditionally used for culinary purposes and/ or contain diverse bioactive compounds with medicinal or health-promoting properties (Sandargo et al. [2019;](#page-153-3) Sum et al. [2023\)](#page-156-10). *Basidiomycota* yeasts are sometimes dimorphic, switching between anamorph (unicellular yeast) and teleomorph (dikaryotic hyphal) stages (Boekhout et al. [2011](#page-132-9)).

**Life modes and distribution:** Saprobes, ectomycorrhizal, parasites (on plants, animals, humans, or fungi including lichens), lichen-forming, insect symbionts, endophytes, flamentous or yeast-like, widespread.

**Notes:** *Basidiomycota* was introduced at the phylum rank by Moore [\(1980](#page-148-5)). It is the second largest phylum in the kingdom, *Fungi*, which includes 2,106 genera belonging to 297 families (He et al. [2024](#page-139-2)). The frst taxonomic review of *Basidiomycota* (and *Entorrhizomycota*) of genera and higher ranks was published by Begerow et al. [\(2018](#page-131-4)). Sub-sequently, He et al. [\(2019](#page-139-1)) provided a complete outline, phylogenetic dating and notes for all genera and higher ranks of *Basidiomycota*.

*Basidiomycota* comprises four subphyla: *Agaricomycotina*, *Pucciniomycotina*, *Ustilaginomycotina*, and *Wallemiomycotina* (Tables [2,](#page-3-0) [16\)](#page-60-0)*.*

**Subphylum** *Agaricomycotina* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVIII (2001).

**Notes:** Doweld ([2001\)](#page-135-7) introduced *Agaricomycotina,* which currently comprises four classes (He et al. [2024\)](#page-139-2), **1.** *Agaricomycetes***; 2.** *Bartheletiomycetes***; 3.** *Dacrymycetes* and **4.** *Tremellomycetes*.

*Agaricomycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVIII (2001).

Index Fungorum Registration Identifer 501297; Fig. [24a](#page-61-0)

The morphological diversity of fruiting bodies in *Agaricomycetes* is unparalleled in any other clade of *Fungi*. In *Agaricomycetes*, the pileate-stipitate morphotype is dominant, along with the coralloid, polyporoid, corticioid, gasteroid, and other forms (Hibbett et al. [2014\)](#page-139-3). Size variation of their fruiting bodies encompasses at least four orders of magnitudes ranging from tiny cyphelloid forms, such as *Henningsomyces candidus* (Pers.) Kuntze (in the order of 1 mm) (Wei et al. [2007\)](#page-159-3), to giant polypores (*Phellinus ellipsoideus* (B.K. Cui & Y.C. Dai) B.K. Cui, Y.C. Dai & Decock), which can measure up to 1,085 cm long (Dai and Cui [2011\)](#page-134-5). Sánchez-García et al. ([2020](#page-152-4)) highlighted that morphological transitions, not nutritional modes, are the most important drivers of diversifcation across *Agaricomycetes* and that lineages with pileate-stipitate basidiomata have strongly increased diversifcation rates across all clades.

**Life modes and distribution:** *Agaricomycetes* are saprobes (e.g., *Agaricaceae*, *Polyporaceae*), and form mutualistic symbioses with a great variety of vascular plants (e.g., *Amanitaceae*, *Boletaceae*, *Russulaceae*) or insects (e.g., *Termitomyces*). Others are plant pathogens (e.g., *Heterobasidion*, *Armillaria*) (Weiß et al. [2004](#page-159-4)), human pathogens (e.g., *Schizophyllum commune* Fr.), mycophilous fungi and mycoparasites (e.g., *Squamanita*) (Liu et al. [2021;](#page-145-5) Schoutteten et al. 2024), and predators on nematodes (e.g., *Hohenbuehelia*, *Pleurotus*). Although less common, some *Agaricomycetes* species are lichenized (e.g. *Omphalina licheniformis* X.L. Wei, Z.H. Cao & R.L. Zhao) (Zhang et al. [2022d](#page-161-6)). Members of the class are widespread.

**Notes:** *Agaricomycetes* with about 30,000 species encompasses the majority of described species in *Basidiomycota,* including 22 orders, 128 families, and 1,434 genera (He et al. [2019](#page-139-1)). As a result of the complexity of morphological and ecological diversity, there are many difficulties in the delimitation of *Agaricomycetes* species. Given this, the best practice is to delimit species and publish new taxa based on as many independent lines of evidence as available, that is, by applying a so-called integrative (or polyphasic)

<span id="page-60-0"></span>**Table 16** Subphyla and classes of *Basidiomycota* Phylum *Basidiomycota*



<span id="page-61-0"></span>**Fig. 24** Diferent taxa of *Basidiomycota*. **a** *Agaricomycetes* (*Amanita hemibapha*). **b** *Bartheletiomycetes* (*Bartheletia paradoxa*). **c** *Cystobasidiomycetes* (*Cyphobasidium*), white arrow shows the fruiting bodies. **d** Dacrymycetes (*Dacryomyces spathularia*). **e** *Exobasidiomycetes* (*Exobasidium ferrugineae*). **f** *Malasseziomycetes* (*Malassezia*). **g** *Microbotryomycetes* (*Microbotrium*). **h** *Physodermatomycetes* (*Physoderma menyanthis*). **i** *Pucciniomycetes* (*Puccinia*). **j** *Tremellomycetes* (*Tremella fusiformis*). **k** *Tritirachiomycetes* (*Tritirachium oryzae*). **l** *Ustilaginomycetes* (*Mycosystema maydis*). **m** *Wallemiomycetes* (*Wallemia sebi*). Photo credits: Figure a: Stephen Axford, Figures B-M: iNaturalist (The photos have been used under the CC BY-NC 4.0 noncommercial use license)



taxonomic approach (Cao et al. [2021](#page-133-2)). The most comprehensive phylogenetic analyses of *Agaricomycetes* so far were a multilocus and genomic-scale data-based phylogeny of 5,284 taxa (Varga et al. [2019\)](#page-158-4) and a fve-locus analysis of 8,472 taxa (Sánchez-García et al. [2020](#page-152-4)).

The vast majority of wild or cultivated edible fungi include *Agaricus bisporus* (J.E. Lange) Imbach, *Tricholoma matsutake* (S. Ito & S. Imai) Singer, and *Lentinula edodes* (Berk.) Pegler, all of which are *Agaricomycetes* (Niego et al. [2023\)](#page-148-2). Similarly, a signifcant proportion of medicinal mushrooms, including *Ganoderma lucidum* (Curtis) P. Karst., and *Hericium erinaceus* (Bull.) Pers., also belong to the *Agaricomycetes* (Thongbai et al. [2015;](#page-157-5) Dai et al. [2021](#page-134-6)). However, there also are many poisonous mushrooms threatening public health. The lethal species are mainly from *Agaricales*, such as *Amanita exitialis* Zhu L. Yang & T.H. Li,

*A. phalloides* (Vaill. ex Fr.) Link, and *Galerina marginata* (Batsch) Kühner (He et al. [2022b](#page-139-4)).

**Type order:** *Agaricales* Underw., Moulds, mildews, and mushrooms. A guide to the systematic study of the Fungi and Mycetozoa and their literature (New York): 97 (1899). **Type family:** *Agaricaceae* Chevall., Fl. gén. env. Paris (Paris) 1: 121 (1826).

**Type genus:** *Agaricus* L., Sp. pl. 2: 1171 (1753).

**Type species:** *Agaricus campestris* L. [as 'campester'], Sp. pl. 2: 1173 (1753).

*Bartheletiomycetes* Thines, Mycological Progress 17 (3): 340 (2017).

Index Fungorum Registration Identifer 815512; Fig. [24](#page-61-0)b

**Asexual morph:** *The sori* produce copious conidia (slimy, hyaline 1-celled), which germinate readily to infect other newly fallen leaves. *Conidiophores* branched, hyaline, fnely verruculose in the apical parts, originating from  $a \pm com$ pact base of irregularly interwoven, hyaline to pale brown intramatrical hyphae. *Conidiogenesis* holoblastic. *Conidiogenous cells* are mostly long, terminal, or intercalary with one or two conidiogenous branches, unilocal with percurrent proliferation and rather dense annellations, and/or multilocal with distichous sympodial proliferation and densely stacked secession scars. *Conidia* hyaline, one-celled, straight, cylindrical-bacilliform or sometimes slightly broader below the middle with a short attenuate base and minutely truncate scar, uninucleate with the nuclear area in the middle, with minute guttules, fnely verruculose. The *conidiophores* are soon replaced by teliospores originating from the same basal cushion of intramatrical hyphae (Scheuer et al. [2008](#page-153-4)). **Sexual morph:** thick-walled, dark brown teliospores either single, immersed in the leaf tissues and  $\pm$  evenly dispersed (most conspicuous in the epidermis), or conglutinated in compact, erumpent, and finally  $\pm$  superficial telia. *Telia* (similar to those of rust fungi) are single or arranged in  $\pm$  circular groups. Telia often develop from conidial sori, hemispherical or cushion-like to  $\pm$  spherical. After a year of dormancy, teliospores germinate with stipitate, longitudinally septate, statismosporic phragmobasidia and produce basidiospores on each sporogenous locus (Scheuer et al. [2008\)](#page-153-4).

**Life modes and distribution:** *Bartheletia paradoxa* is apparently saprotrophic rather than endophyte, strictly hostspecifc and widely distributed fungal companion of *Ginkgo biloba.* Most probably, just like its host plant, the fungus is also a living fossil (Scheuer et al. [2008\)](#page-153-4). Its biology is unclear. Kirschner and Okuda ([2013](#page-143-16)) showed that the fungus is a pioneer colonizer in the initial stage of decomposition of *Ginkgo* litter. According to available information, although *Ginkgo* is grown worldwide, the global distribution of *B. paradoxa* is uneven. The species are generally distributed in Europe (Austria, Czech Republic, Denmark, France, Germany, Sweden, Netherlands, and the United Kingdom), Russia, Ukraine, and Asia (Japan and Korea) (Hayova and Tykhonenko [2017\)](#page-139-5).

**Note:** The family *Bartheletiaceae* was proposed to accommodate *B. paradoxa,* which was discovered on ginkgo leaf litter and remained unknown for a long time (Scheuer et al. [2008\)](#page-153-4). The first valid information on this fungus was included only by Scheuer et al. [2008](#page-153-4) (Hayova and Tykhonenko [2017](#page-139-5)). The position of *B. paradoxa* was not resolved in the initial nrSSU DNA sequence analysis, but Mishra et al. ([2018\)](#page-147-4) took a more phylogenetic approach using a large set of conserved genes and ofered a perfect solution. The new study used 67 highly conserved orthologous loci for network analysis. New data excluding negative genes strongly (but not conclusively) support that this organism is at the base of the *Agaricomycetes* and is the sister of all other members of the sub-phylum and separate from the *Ustilaginomycotina* and *Pucciniomycotina*, which together create a sister group to the whole of the *Agaricomycotina*. The new class and order *Bartheletiomycetes* and *Bartheletiales*, were introduced by Mishra et al. ([2018](#page-147-4)).

According to Wijayawardene et al. [\(2020a\)](#page-160-5), the class *Bartheletiomycetes* consists of a single order *Bartheletiales*, single family *Bartheletiaceae*, and single genus *Bartheletia. B. paradoxa* is the only species within this genus (Scheuer et al. [2008\)](#page-153-4). *Bartheletiomycetes paradoxa* reproduces sexually only on ginkgo leaves and can survive in some tissues of ginkgo trees as an asymptomatic disease (Scheuer et al. [2008](#page-153-4)).

**Type order:** *Bartheletiales* Thines, Mycological Progress 17 (3): 340 (2017).

**Type family:** *Bartheletiaceae* R. Bauer, Scheuer, M. Lutz & Grube, in Scheuer, Bauer, Lutz, Stabentheiner, Melnik & Grube, Mycol. Res. 111(11): 1269 (2008).

**Type genus:** *Bartheletia* G. Arnaud ex Scheuer, R. Bauer, M. Lutz, Stabenth., Melnik & Grube, Mycol. Res. 111(11): 1269 (2008).

**Type species:** *Bartheletia paradoxa* G. Arnaud ex Scheuer, R. Bauer, M. Lutz, Stabenth., Melnik & Grube, Mycol. Res. 112(11): 1269 (2008).

*Dacrymycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVIII (2001).

Index Fungorum Registration Identifer 501296; Fig. [25](#page-63-0) & [26](#page-64-0)

**Sexual morph:** *Basidiomata* scattered or gregarious, coalesced, displaying a wide variation in size and shape, pustulate to pulvinate or subglobose, cylindrical, subulate, rugose pilei, palmate, stipitate-pileate or sessile, strikingly long bisterigmate (forked) holobasidia, pale yellow to yellow, orange, olive, reddish brown to dark reddish brown, frm-gelatinous or soft-cartilaginous (Hibbett et al. [2007](#page-139-6)), except *Unilacryma unispora* (Shirouzu et al. [2013\)](#page-154-6). *Hymenium* amphigenous, producing numerous basidia. *Basidia* 1–3-spored, bearing a loop-like basal clamp connection; apex frequently U-shaped, sometimes Y-shaped dacrymycete, and more rarely with an apical protuberance, bearing elliptical aseptate spores at the apices of these basidia (Zamora and Ekman [2020\)](#page-161-7). *Basidiospores* 0–1-septate (exceptionally 3-septate), thin-walled, hyaline, cylindricallantoid. Spore prints whitish. Basidia and basidiospores with or without conspicuous lipid drops when old. *Hyphae* densely branching and frequently anastomoses, with a



<span id="page-63-0"></span>**Fig. 25** Basidiocarps of *Dacrymycetales*. No scale bar, modifed from Oberwinkler ([2012\)](#page-150-3), and Oberwinkler et al. ([2014\)](#page-150-4)

loop-like basal clamp attachment. Brownish intracellular and difuse parietal pigments often visible in some cortical/ marginal hyphae. **Asexual morph:** Asexual fructifcations are prevalent in *Dacrymyces stillatus*; however, infrequent or absent in other members of the *Dacrymycetes*. Fragmenting hyphae typically manifest in asexual fructifcations but may also occur sporadically in basidiocarps (Oberwinkler [2014](#page-149-4)).

**Life modes and distribution:** The members of *Dacrymycetes* occur worldwide (Brasfeld [1938;](#page-132-10) Shirouzu et al. [2016,](#page-154-7) [2017\)](#page-154-8), and are reported as brown-rot fungi (Worrall et al. [1997\)](#page-160-6), and some have been reported to degrade considerable amounts of lignin (Seifert [1983\)](#page-154-9). Gene contents from two *Calocera* genomes also suggest a brown-rotting lifestyle (Nagy et al. [2016\)](#page-148-6), and *Dacrymycetes* are the clade with the oldest stem age among *Basidiomycota* wood decomposers (e.g. *Pseudotsuga* spp., *Pinus* spp., *Juniperus* spp., Fagaceae, Nothofagaceae, Myrtaceae, Podocarpaceae, and Araucariaceae). The ancestors of this class and the *Agaricomycetes* appear to have separated during the upper Devonian or Carboniferous periods (Garnica et al. [2016](#page-137-9); Varga et al. [2019](#page-158-4)).

**Notes:** Doweld [\(2001](#page-135-7)) introduced the class *Dacrymycetes*, one of the three widely accepted classes of phylum *Basidiomycota*. Worrall et al. [\(1997\)](#page-160-6) provided a comprehensive background of the classifcation of these typically brown-rot fungi, and *Dacrymycetes* are the clade with the oldest stem age among *Basidiomycota* wood decomposers (Oberwinkler [1994](#page-149-5); Kirk et al. [2008](#page-142-0); Floudas et al. [2012\)](#page-136-7). According to Wijayawardene et al. ([2022a\)](#page-160-3), the class comprises two orders, four families, and twelve genera.

About 400 published species names, but many of these names are synonyms, at least 120 reported species have been identifed (Savchenko et al. [2021](#page-153-5)). However, predicting from recent studies that the real number of species is



<span id="page-64-0"></span>**Fig. 26** Micromorphological characters of *Dacrymycetales*. **a**–**d** Basidiospores. e Terminal cells of cortical/marginal hyphae, branched and gelatinised in *Dacryonaema macnabbii*. **f** Bisterigmate basidia in *Dacrymyces estonicus.* **g** Unisterigmate basidium in *Unilacryma unispora.* **h** Hyphidia, simple in *Dacrymyces ovisporus*. Modified from Zamora and Ekman  $(2020)$  $(2020)$ . Scale bars: **a**,  $c = 5 \mu m$ , **b**, **d**–  $h = 10 \mu m$ 

certainly larger, probably over 200 (Savchenko et al. [2021](#page-153-5)). Except *Unilacryma unispora*, which contains unisterigmate basidia, members of the class may be easily diferentiated from other basidiomycetes by their bisterigmate Y-shaped basidia (Wells [1994](#page-159-5); Shirouzu et al. [2013](#page-154-6)). *Dacrymycetes* are distinguished by dolipore septa with imperforate parenthesomes, or rarely with a single pore on an ultrastructural level (Maekawa [1987](#page-146-4); Oberwinkler [1994](#page-149-5); Shirouzu et al. [2013\)](#page-154-6). In terms of macromorphology, the class is dominated by "jelly fungi" with pustulate-pulvinate, cupulate, dendroid, and spathulate basidiocarps coloured with carotenoids in various yellow and orange colours (Goodwin [1953;](#page-137-10) Czeczuga [1980;](#page-134-7) Zamora and Ekman [2020](#page-161-7)). As a result, they have been used as taxonomic characters both in *Ascomycota* (Eckblad [1968](#page-136-8); Schumacher [1990](#page-154-10); Baral [2004\)](#page-130-3) and *Basidiomycota* (Arpin and Fiasson [1971](#page-130-4); Olariaga et al. [2015;](#page-149-6) Zamora and Ekman [2020\)](#page-161-7). Shirouzu et al. [\(2017](#page-154-8)) provided the most complete phylogeny in terms of included vouchers with both DNA data and morphological data from fruitbodies. Some of the provisionally recognized species were found to be polyphyletic or to have extremely lengthy branches within their clade, indicating that there is signifcant sequence variation. Zamora and Ekman [\(2020\)](#page-161-7) provided multi loci phylogeny of the class *Dacrymycetes* and revealed fve lineages that putatively represent fve diferent families, four of which were previously accepted and named.

He et al. [\(2024\)](#page-139-2) recognized only one order *Dacrymycetales* in *Dacrymycetes.*

**Type Order:** *Dacrymycetales* Henn., in Engler & Prantl, Nat. Pflanzenfam., Teil. I (Leipzig) 1(1\*\*): 96 (1897) [1900].

**Type family:** *Dacrymycetaceae* J. Schröt. [as 'Dacrymycetini'], in Cohn, Krypt.—Fl. Schlesien (Breslau) 3.1(25–32): 399 (1888).

**Type genus:** *Dacrymyces* Nees [as 'Dacryomyces'], Syst. Pilze (Würzburg): 89 (1816) [1816–17].

**Type species:** *Dacrymyces stillatus* Nees, Syst. Pilze (Würzburg): 89 (1816) [1816–17].

*Tremellomycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow):73 (2001).

Index Fungorum Registration Identifer: 90764; Fig. [24](#page-61-0)k

Dimorphic fungi, yeasts, hyphae and/or complex fruiting bodies. Fruiting bodies are gelatinous or absent, parenthesomes sacculate or absent, basidia septate or nonseptate (Hibbett et al. [2007](#page-139-6)).

**Life modes and distribution:** The members of *Tremellomycetes* are a nutritionally heterogeneous group comprising saprobes, human and animal parasites, mycoparasites and lichen-inhabiting species and distributed worldwide (Sampaio [2004;](#page-152-5) Boekhout et al. [2011;](#page-132-9) Cooper [2011;](#page-134-8) Millanes et al. [2011;](#page-147-5) Weiss et al. 2014; Schoutteten et al. 2024). Species from the genera *Cryptococcus*, *Naganishia*, *Papiliotrema*, and *Trichosporon* have been reports as human and animal pathogens (Kurtzman et al. [2011](#page-143-1)). Several *Tremella* and *Naematelia* species are mycoparasites and have been cultivated for food (Schmidt et al. [2011;](#page-153-6) Ma et al. [2021;](#page-146-5) Kuddus and Aguilar [2022;](#page-143-17) Linnakoski et al. [2023\)](#page-145-6). Some taxa have been exclusively reported in agriculture and industrial applications (Sun et al. [2020](#page-156-11); Palmieri et al. [2021](#page-149-7); Yan et al. [2022](#page-161-8)).

**Notes:** The name *Tremellomycetes* was circumscribed by Hibbett et al. [\(2007](#page-139-6)). The members of *Tremellomycetes* dimorphic taxa, but the majority of species is only known from yeast morphs. Fruiting bodies of the hyphal morph in this class either develop intrahymenially or comprise complex fruiting bodies, often with a gelatinous texture (Millanes et al. [2011,](#page-147-5)

Weiss et al. 2014). The anamorphic *Tremellomycetes* have been found in the yeast morph, including *Bullera*, *Cryptococcus*, *Dioszegia* and *Fellomyces* among others (Nakase [2000](#page-148-7); Litvinseva et al. 2006; Boekhout et al. [2011;](#page-132-9) Kurtzman et al. [2011\)](#page-143-1). Among the sexually typifed taxa, the spore-producing basidia are often formed in gelatinous basidiocarps or gall deformations on other fungi (Chen et al. [2001](#page-133-3); Sampaio [2004](#page-152-5); Millanes et al. [2011\)](#page-147-5). Phylogenetic analyses of rRNA gene (SSU, ITS, and LSU sequence data) of Hibbett et al. ([1998,](#page-139-7) [2007\)](#page-139-6), Matheny et al. ([2006\)](#page-146-6), and Millanes et al. ([2011\)](#page-147-5) accommodated this class in subphylum *Agaricomycotina.*

He et al. [\(2019\)](#page-139-1) provided the divergence time estimates for the various orders and families in the class, which comprises fve orders, 18 families, and 54 genera. According to Wijayawardene et al.  $(2022a)$  $(2022a)$  $(2022a)$ , the class comprises six orders (*Chionasterales*, *Cystoflobasidiales*, *Filobasidiales*, *Holtermanniales*, *Tremellales*, and *Trichosporonales*), 20 families and 73 genera. Various genera have a presumed afnity to *Tremellomycetes* based on morphological and ultrastructural similarities, namely Anastomyces, Celatogloea, *Heteromycophaga*, Phragmoxenidium, Sigmogloea, Sirotrema, Tremellina, *Trichosporonoides*, and *Xenolachne*. *Since viable cultures and/or DNA sequence data are lacking* 

*for these genera, they are currently classifed as Tremellomycetes incertae sedis.*

The members of *Tremellomycetes* are widely used in different applications (Table [17\)](#page-65-0) while some are medicinally important taxa, either as source of pharmacological compounds or as infectious agents (Table [18](#page-65-1)).

**Type order:** *Tremellales* Fr. [as 'Tremellinae'], Syst. mycol. (Lundae) 1: 2 (1821).

**Type family:** *Tremellaceae* Fr. [as 'Tremellini'], Syst. mycol. (Lundae) 1: lv (1821).

**Type genus:** *Tremella* Pers., Syn. meth. fung. (Göttingen) 2: 622 (1801).

**Type species:** *Tremella mesenterica* Retz., K. svenska Vetensk-Akad. Handl., ser. 1 30: 249 (1769).

**Subphylum** *Pucciniomycotina* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 45 (2006).

**Notes:** Based on multilocus phylogenetic reconstructions and ultrastructural markers, Bauer et al. [\(2006](#page-131-5)) introduced *Pucciniomycotina* to accommodate *Agaricostilbomycetes*, *Atractiellomycetes*, *Classiculomycetes*, *Cryptomycocolacomycetes*, *Cystobasidiomycetes*, *Microbotryomycetes*, *Mixiomycetes* and

<span id="page-65-0"></span>**Table 17** *Tremellomycetes* taxa used in diferent industries

<b>Species</b> Industry	References
Brewing of beer Mrakia gelida	Linnakoski et al. (2023)
Phaffia rhodozyma Astaxanthin production for the feed industry and dietary supple- ment	Schmidt et al. $(2011)$
Enzyme production for the food industry Tausonia pullulans	Kuddus and Aguilar $(2022)$
Cosmetic, daily chemical, and food industries Tremella fuciformis	Ma et al. (2021), Yuan et al. (2022)
Naematelia aurantialba Food industries, source of bioactive polysaccharides, and medici- nal purposes	Sun et al. (2020), Yan et al. (2022)
Biocontrol agent used in the field and postharvest Papiliotrema terrestris	Palmieri et al. (2021)

<span id="page-65-1"></span>**Table 18** Clinically important members of *Tremellomycetes*



<span id="page-66-0"></span>**Fig. 27** *Agaricostilbum palmicola.* **a** Habit. **b** Conidiophore tip showing the emergence of conidia, and **c** Conidia×2800. **d** Branching of young conidiophore  $\times 850$ . Scale bars:  $a=1$  mm, **b**,  $c=5$  µm (redrawn from Wright [1970\)](#page-160-7)



*Pucciniomycetes.* Subsequent studies by Schell et al. [\(2011\)](#page-153-7) and Wang et al. ([2015b\)](#page-159-6) introduced *Tritirachiomycetes* and *Spiculogloeomycetes* respectively. Currently, *Pucciniomycotina* consists of ten classes comprising organisms with a diverse range of morphological features and ecological strategies. Members of Pucciniomycotina have a type A structure of the 5.8S rDNA, mannose as the major sugar component of the cell wall, and are characterised by septal pores without membrane caps, often referred to as simple septal pores.

## *Agaricostilbomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Progr. 5(1): 45 (2006).

Index Fungorum Registration Identifer 501474; Fig. [27](#page-66-0)

*Agaricostilbomycetes* are a highly diverse class of fungi in terms of morphology and ecology. Currently, the class comprises one order, fve families, and more than 70 species distributed over 13 genera (Aime et al. [2014;](#page-130-6) Wang et al. [2015b](#page-159-6); Oberwinkler [2017;](#page-149-9) Begerow et al. [2018;](#page-131-4) He et al. [2019;](#page-139-1) Li et al. [2020;](#page-145-8) Cao et al. [2021\)](#page-133-2). The monophyly of the class was recognised by various phylogenetic studies (Bauer et al. [2006](#page-131-5); Aime et al. [2006;](#page-130-7) Wang et al. [2015b\)](#page-159-6). These studies often recovered long terminal branches in this class, indicating large phylogenetic distances between the representatives. At the ulturastructural level, the class is characterised by septal pores with simple morphology and without associated microbodies, separation of nucleoplasmic spindle pole bodies, and metaphasic spindle pole bodies that occur intranuclearly (Bauer et al. [2006;](#page-131-5) Begerow et al. [2018](#page-131-4)). Most species are exclusively known as asexually reproducing yeast morphs, whereas a few species are characterised by a dimorphic lifecycle, alternating between a unicellular yeast morph and a multicellular flamentous morph in which sexual structures are developed. It is likely that all *Agaricostilbomycetes* are dimorphic organisms, and that the hyphal morph for most species remains to be discovered in natural or laboratory conditions. For example, Fonseca et al. [\(2000](#page-137-11)) reported the development of a hyphal morph with transversally septate basidia in *Kondoa aeria* and *K. malvinella* in culture conditions after several months of inoculation. Basidiomata are usually small and comprise stilboid (*Chionosphaera*, *Crittendenia*, *Sterigmatomyces* and *Stilbum*) and pustulate forms (*Mycogloea* spp.). The studied dimorphic species in this class display a large diversity in the structures for sexual reproduction. *Stilbum* spp. develop transversally septate basidia producing statismospores. *Chionosphaera* spp. and *Crittendenia* spp. produce holobasidia with statismospores (Kirschner et al. [2001](#page-143-19)). *Kondoa* spp. produce transversally septate basidia producing ballistospores (Fonseca et al. [2000](#page-137-11)). *Mycogloea* spp. produce transversally septate basidia that become detached from the basidiogenous cell, after which sterigmata and ballistospores are formed (Bandoni [1998;](#page-130-8) Kirschner et al. [2003](#page-143-20)).

**Life modes and distribution:** *Agaricostilbomycetes* have been isolated from a large diversity of substrates in marine and terrestrial habitats. Yeast morphs are often considered to act as saprobes. Hyphal morphs were isolates from soil (*Cystobasidiopsis nirenbergiae*), palm litter (*Sterigmatomyces* spp.), lichens (*Crittendenia* spp.), and fruitbodies of other fungi (*Bensingtonia* spp. and *Kondoa* spp.). Whether the fungicolous and lichenicolous species are parasites of their respective hosts remains unknown (Millanes et al. [2021](#page-147-8)). Haustorial cells are often seen as an indication of mycoparasitism and were reported from *Stilbum vulgare* and *Kondoa myxariophila*, but only in laboratory conditions (Seifert et al. [1992](#page-154-11); Li et al. [2020](#page-145-8); Schoutteten et al. 2024).

**Notes:** Bauer et al. [\(2006](#page-131-5)) introduced the class *Agaricostilbomycetes* to accommodate two orders, *Agaricostilbales* Oberw. & R. Bauer and *Spiculogloeales* R. Bauer et al. However, *Spiculogloeales* was only represented by two representatives (*Mycogloea* sp. and *Spiculogloea* sp.), and the authors already suggested that in the future the clade should be separated into its own class (Bauer et al. [2006](#page-131-5)). Wang et al. ([2015b\)](#page-159-6) showed that the *Sporobolomyces subbruneus* clade clustered with *Spiculogloea* sp., and introduced the class *Spiculogloeomycetes* for this lineage.

*Agaricostilbum* species are presumed saprobic fungi on palm leaves that were isolated from various palm hosts. Oberwinkler [\(2017](#page-149-9)) critically discussed the suggested synonymy of *Agaricostilbum palmicola* and *A. pulcherrimum* that was proposed by Brady et al. [\(1984](#page-132-11)) without providing details to support the synonymy. Based on DNA sequence data, Wang et al. ([2015b\)](#page-159-6) combined *A. pulcherrimum* in the genus *Sterigmatomyces*. Oberwinkler ([2017](#page-149-9)) emphasised that this genus is in need of reinvestigation, combining phylogenetic and morphological markers from a broad sampling of various palm hosts.

*Mycogloea nipponica* was shown to be the flamentous morph of *Kurtzmannomyces* sp. (Kirschner et al. [2003](#page-143-20)). Several other *Kurtzmannomyces* yeast species are known, as well as various *Mycogloea* species (Bandoni [1998](#page-130-8)). However, the overlap in data derived from these fungi is minimal, and many teleomorph-anamorph links remain to be discovered. Because of the lack of DNA sequence data, it remains unclear if all known *Mycogloea* species belong to *Agaricostilbomycetes*.

**Type order:** *Agaricostilbales* Oberw. & R. Bauer, Sydowia 41: 240 (1989).

**Type family:** *Agaricostilbaceae* Oberw. & R. Bauer, Sydowia 41: 240 (1989).

**Type genus:** *Agaricostilbum* J.E. Wright, Mycologia 62(4): 679 (1970).

**Type species:** *Agaricostilbum palmicola* J.E. Wright, Mycologia 62(4): 680 (1970).

*Atractiellomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiß & Oberw., Mycol. Progr. 5(1): 45 (2006)

Index Fungorum Registration Identifer 501476

*Atractiellomycetes* is a heterogenous class of which the representatives are only known from flamentous morphs, yeast budding is not known from this class. At the ultrastructural level, *Atractiellomycetes* are characterised by the presence of symplechosomes. These structures were described as stacked plate-like cisternae derived from the endoplasmatic reticulum, interconnected by hexagonally arranged flaments, and connected to mitochondria, but their origin and function remain unknown (Bauer and Oberwinkler [1991a](#page-131-6)). Symplechosomes are synapomorphic for this class, but only a few species were investigated for their ultrastructural

properties. Septal pores are complex and are associated with either microbodies or atractosomes (Bauer et al. [2006](#page-131-5)).

*Atractiellomycetes* comprise the order *Atractiellales*, the families *Atractogloeaceae*, *Hoehnelomycetaceae* and *Phleogenaceae*, ten genera, and more than 60 known species (Aime et al. [2018a;](#page-130-9) Begerow et al. [2018;](#page-131-4) Schoutteten et al. [2018](#page-153-8); Spirin et al. [2018](#page-155-4); Cao et al. [2021](#page-133-2)).

The documented diversity of basidiomata in this class is high, with resupinate (*Bourdotigloea*, *Helicogloea*, *Saccosoma*), stilboid (*Atractiella*, *Phleogena*) and pycnidioid (*Basidiopycnis*, *Proceropycnis*) structrures. Basidia are transversally septate, contain laterally attached probasidia for karyogamy (*Bourdotigloea*, *Helicogloea*, and *Saccosoma*), and usually produce ballistospores (Baker [1936\)](#page-130-10). The formation of statismospores is known from *Atractogloea* and *Phleogena* (Oberwinkler and Bandoni [1982a](#page-149-10), [b](#page-149-11), [c](#page-150-5)). Some *Atractiellomycetes* are known from a teleomorphic and an anamorphic morph, of which the anamorps were formerly assigned to the genera *Hobsonia*, *Infundibura*, *Leucogloea*, and *Pleurocolla* (Kirschner [2004;](#page-143-21) Spirin et al. [2018](#page-155-4)). Some species are known exclusively from the anamorph, e.g. *Proceropycnis pinicola* (Oberwinkler et al. [2006a](#page-149-12), [b](#page-150-6)).

**Life modes and distribution:** *Atractiellomycetes* have a worldwide distribution, covering Northern temperate and tropical /sub-tropical regions. Members of the class usually produce inconspicuous basidiomata or conidiomata, often on woody or herbaceous materials, sometimes overgrowing other fungi (Bauer et al. [2006](#page-131-5); Aime et al. [2018a;](#page-130-9) Spirin et al. [2018\)](#page-155-4). Most *Atractiellomycetes* are considered saprobic, associated withdecaying vegetation, or with bark beetle galleries in decaying conifer logs (Bauer et al. [2006](#page-131-5); Oberwinkler et al. [2006a,](#page-149-12) [b](#page-150-6)). However, recent studies indicated that members of this class might be associated with plant roots and may be mycorrhizal or endorrhizal (Kottke et al. [2010;](#page-143-22) Bonito et al. [2017\)](#page-132-12). One species is reportedly involved in in orchid mycorrhiza (Kottke et al. [2010](#page-143-22)), *Atractiella rhizophila* and *Proceropycnis hameedii* were found to be endophytes of *Populus* roots, and *Atractidochium hillariae* was recovered as an endophyte of *Pinus taeda* needles (Bonito et al. [2017;](#page-132-12) Aime et al. [2018a\)](#page-130-9).

**Note:** Most species in *Atractiellomycetes* are known from one or a few collections only, and DNA sequence data is often limited to ribosomal markers, or completely lacking (Schoutteten et al. [2018](#page-153-8); Cao et al. [2021](#page-133-2)). Also, only a few cultures are available of *Artactiellymycetes*, and spore germination often occurs through the formation of microconidia (Bauer et al. [2006\)](#page-131-5). Spirin et al. [\(2018\)](#page-155-4) performed a taxonomic study on the resupinate, corticioid members of the *Atractiellomycetes*, focussing on the species formerly assigned to *Helicogloea* and *Saccoblastia*. He delineated three genera based on combined phylogenetic and morphological characters. *Helicogloea* spp. are characterised by efbulate hyphae and the absence of cystidia, *Bourdotigloea* spp. are characterised by efbulate hyphae and the presence of cystidia, and *Saccosoma* spp. are characterised by clamped hyphae and the absence of cystidia. However, the current sampling remains low, and sequencing of more well-studied material is necessary to further test and refne the proposed species delimitation in this group (Malysheva et al. [2020](#page-146-7)).

**Type order:** *Atractiellales* Oberw. & Bandoni, Can. J. Bot. 60(9): 1740 (1982).

**Type family:** *Atractiellaceae* R.T. Moore, Mycotaxon 59: 8 (1996).

**Type genus:** *Atractiella* Sacc., Syll. fung. (Abellini) 4: 578 (1886).

**Type species:** *Atractiella brunaudiana* (Sacc.) Sacc., Syll. fung. (Abellini) 4: 579 (1886).

*Classiculomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 46 (2006).

Index Fungorum Registration Identifer 501479.

*Classiculomycetes* were introduced by Bauer et al. [\(2006\)](#page-131-5) to accommodate a clade with two species isolated from aquatic habitats, namely *Classicula fuitans* and *Jaculispora submersa*. More recently, Qiao et al. [\(2018\)](#page-150-7) described a second *Classicula* species from China. The class comprises one order (*Classiculales*), one family (*Classiculaceae*), two genera and three species (Bauer et al. [2006;](#page-131-5) Begerow et al. [2018;](#page-131-4) He et al. [2019\)](#page-139-1). At the ultrastructural level, the septal pores have a simple morphology, and are associated with microbodies (Bauer et al. [2003](#page-131-7), [2006](#page-131-5)). All species were originally isolated and described as anamorphic fungi, with typical navicular conidia characterised by three or four setose appendages, clamped hyphae, and the presence of basally clamped haustorial cells. All three species are known from flamentous morphs only, no yeast budding is known. *Naiadella fuitans* was originally isolated as an anamorphic fungus from leaf litter in a freshwater stream in Canada (Marvanová and Bandoni [1987](#page-146-8)). Bauer et al. ([2003](#page-131-7)) reported the production of the sexual state of *Naiadella fuitans*, which is characterised by transversally septate basidia producing ballistospores and for which the name *Classicula fuitans* was introduced. These basidia were only observed in pure culture after soaking the fungus in water. No basidiospore germination was observed (Bauer et al. [2003\)](#page-131-7). Aime et al. [\(2018b](#page-130-11)) called for the protection of the name *Classicula*, based on which the nomenclature of the higher taxa in the class is based.

**Life modes and distribution:** *Classicula* and *Jaculispora* were isolated from aquatic environments and are especially associated with leaf litter in freshwater habitats (Bauer et al. [2003](#page-131-7)). The conidia with broadly diverging setose branches in the three known species resemble those of other aquatic hyphomycetes and may be an adaptation to survival and dispersal in such environments (Ingold [1979](#page-141-8)). Also, the long and small basidiospores produced by *Classicula fuitans* could be an adaptation to water dispersal (Bauer et al. [2003](#page-131-7)). Plant host preferences have been tested for *Jaculispora submersa*, and suggest an affinity for oak leaves (Prokhorov and Bodyagin [2007](#page-150-8)). There are indications that *Classicula* and *Jaculispora* species are mycoparasites because of the presence of basally clamped haustorial cells. Such haustorial cells are known from many basidiomycetous mycoparasites, and along the contact surface between the haustorial cell and host cell one or more fusion pores are produced (Bauer [2004](#page-131-8); Bauer et al. [2006](#page-131-5); Schoutteten et al. 2024). However, the haustorial cells in *C. fuitans* were only observed to attach to hyphae of their own mycelium, which was interpreted as self-parasitism, but no fusion pores were observed (Bauer et al. [2003](#page-131-7)). It remains unclear if these structures are also formed in natural conditions, and if, and with which other fungi mycoparasitic associations are established.

**Notes:** The species diversity of this class may be higher than currently known because freshwater ecosystems have always been under-sampled niches for fungi (Cao et al. [2021](#page-133-2)). Generally, *Jaculispora submersa* is distributed in aquatic environments in Jamaica (Hudson and Ingold [1960](#page-140-4)) and is also found in Europe, Great Britain, Canada, North America (Cole and Kendrick [1981\)](#page-133-4), and India (Borse et al. [2015](#page-132-13)). *Classicula fuitans* found in Canada, Czechia, and India (Borse et al. [2015\)](#page-132-13). *Classicula sinensis* was only recorded in China (Qiao et al. [2018\)](#page-150-7).

**Type order:** *Classiculales* R. Bauer, Begerow, Oberw. & Marvanová, Mycologia 95(4): 763 (2003).

**Type family:** *Classiculaceae* R. Bauer, Begerow, Oberw. & Marvanová, Mycologia 95(4): 763 (2003).

**Type genus:** *Classicula* R. Bauer, Begerow, Oberw. & Marvanová, Mycologia 95(4): 757 (2003).

**Type species:** *Classiculafluitans* R. Bauer, Begerow, Oberw. & Marvanová, Mycologia 95(4): 757 (2003).

*Cryptomycocolacomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 46 (2006).

Index Fungorum Registration Identifer 90612; Fig. [28](#page-69-0)

*Cryptomycocolacomycetes* is comprising the order *Cryptomycocolacales*, family *Cryoptomycocolacaceae*, and the two monotypic genera *Cryptomycocolax* and *Colacosiphon* (Oberwinkler and Bauer [1990a](#page-149-13), [b](#page-150-9); Kirschner et al. [2001](#page-143-19); Bauer et al. [2006;](#page-131-5) Begerow et al. [2018\)](#page-131-4). The class was introduced by Bauer et al. ([2006](#page-131-5)) based on molecular phylogenetic reconstructions combined with ultrastructural and morphological observations. Both *Cryptomycocolax abnormis* and *Colacosiphon fliformis* are colacosome-interacting mycoparasites associated with ascomycetous hosts. At the <span id="page-69-0"></span>**Fig. 28** *Cryptomycocolax abnorme*. **a** Habit sketch of fructifcation growing in and on sclerotia of the host; one sclerotium longitudinally sectioned. **b** Basidial development (**a**); globose probasidium (**b**, **c**); apical elongation of young basidium (**d**); two celled-basidium (phragmobasidial stage). **c** Basidial apices of mature, one-celled holobasidia with sessile basidiospores at diferent development stages. **d** Old apex of basidium with bases of spore attachments. **e** Sectional view of sclerotium with hyphae of the host, unclamped and hyphae and basidia of the parasite with clamps; note botryose structures of cellular interactions. The basidia are in diferent development stages. **f** Germination of basidiospores by hyphae. **g** Yeast-like budding of basidiospores. Scale bars: a=1 mm;  $b-g=10 \mu m$  (re-drawn from Oberwinkler and Bauer [1990a,](#page-149-13) [b](#page-150-9))



ultrastructural level, both species are characterised by simple septal pores surrounded by woronin-like bodies, which are reminiscent of the woronin bodies found in *Ascomycota*. *Colacosiphon abnormis* was found to contain two distinct types of colacosomes with diferences in their morphology and spatial organisation. In *C. fliformis* on the other hand, only one type of colacosomes was reported, which is presumably the same as one of the types found in *C. abnormis*, and in mycoparasites of *Microbotryomycetes*. The colacosomes are mostly organised in vesicular cells, which are invaginate by host hyphae. Along the contact surface, a dense formation of colacosomes is observed, resulting in botryose structures in these cells. Reproductive structures in these two species are considerably diferent from each other and from other *Basidiomycota*. The sporogenous cells are elongated and produce a variable number of sessile spores. Oberwinkler and Bauer ([1990a](#page-149-13), [b\)](#page-150-9) performed an excellent study of the lifecycle of *C. abnormis* by using transmission electron microscopy (TEM), and observed meiosis in the elongated cells, which they interpreted as basidia. During basidium development, a short phragmobasidial stage is present in which the basidium is transversally one-septate. Later in the development, the top cell is abscised and the basal cell elongates, eventually producing sessil spores apically. The nature of the sporogenous cells in *Colacosiphon* was not studied in such detail by TEM and remains unclear (Kirschner et al. [2001](#page-143-19)). Therefore, these structures are tentatively interpreted as conidiogenous cells for asexual reproduction. Yeast budding was microscopically observed in *C. abnormis*, but not in *C. fliformis* (Oberwinkler [2017](#page-149-9)).

**Life modes and distribution:** Both species in this class are colacosome-interacting mycoparasites, associated with ascomycetous hosts. *Cryptomycocolax abnormis* was isolated as a slimy layer overgrowing pyrenomycetes on a decaying branch of *Circium subcoriaceum* in Costa Rica (Oberwinkler and Bauer [1990a](#page-149-13), [b\)](#page-150-9). *Colacosiphon fliformis* was isolated from bark beetles (*Ips sexdentatus*) retrieved from decaying *Pinus sylvestris* logs in Germany. This species was obtained in co-culture with other fungi, in which it was found in a parasitic interaction with an ascomycetous host (Kirschner et al. [2001](#page-143-19)).

**Notes:** Currently, only DNA sequence data from the D1/ D2 region of the LSU ribosomal nuclear DNA are available for these two species. Weiß et al. [\(2004\)](#page-159-4) discussed the

possibility that *Cryptomycocolacomycetes* is a sister to all other *Basidiomycota*, but the phylogenetic analyses rendered low bootstrap support for this scenario. Also in later studies, the phylogenetic position of this class was unresolved (e.g. Bauer et al. [2006\)](#page-131-5). It can be expected that more diversity within this class remains to be discovered, and genome-scale data will help to determine the evolutionary relationships of this class and the species it comprises.

**Type order:** *Cryptomycocolacales* Oberw. & R. Bauer, Mycologia 82(6): 672 (1990).

**Type family:** *Cryptomycocolacaceae* Oberw. & R. Bauer, Mycologia 82(6): 672 (1990).

**Type genus:** *Cryptomycocolax* Oberw. & R. Bauer, Mycologia 82(6): 672 (1990).

**Type species:** *Cryptomycocolax abnormis* Oberw. & R. Bauer [as 'abnorme'], Mycologia 82(6)**:** 672 (1990).

*Cystobasidiomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 46 (2006).

Index Fungorum Registration Identifer 501480.

The class *Cystobasidiomycetes* was proposed by Bauer et al. [\(2006](#page-131-5)) and comprises six orders (*Buckleyzymales*, *Cyphobasidiales*, *Cystobasidiales*, *Erythrobasidiales*, *Naohideales*, and *Sakagushiales*), eight families (*Buckleyzymaceae*, *Cyphobasidiaceae*, *Cystobasidiaceae*, *Erythrobasidiaceae*, *Microsporomycetaceae*, *Naohideaceae*, *Sakaguchiaceae*, and *Symmetrosporaceae*), 16 genera, and over 100 species (Wang et al. [2015b;](#page-159-6) Begerow et al. [2018](#page-131-4); He et al. [2019;](#page-139-1) Li et al. [2020\)](#page-145-8). At the ultrastructural level, *Cystobasidiomycetes* have simple septal pores which are surrounded by cylindrical organelles with a reticulate surface, called cystosomes.

The cystosome seems to be synapomorphic for this class, but they were only reported from a few dimorphic representatives. However, the cystosome was not found in the septal pores of *Naohideales*. Cell walls of *Cystobasidiomycetes* lack fucose in the sugar composition (Takashima et al. [2000](#page-156-12); Bauer et al. [2006](#page-131-5)). The vast majority of *Cystobasidiomycetes* is exclusively known from a yeast morph, although it is likely that most of them have dimorphic life cycles. The flamentous morphs of *Cystobasidiomycetes* are usually characterised by transversally septate basidia, often subtended by a thin- or thick-walled probasidium. Reproductive structures in *Bannoa* and *Erythrobasidium* are single-celled, but it remains unclear whether it concerns conidiophores or real basidia (Bauer et al. [2006;](#page-131-5) Oberwinkler [2017](#page-149-9); Begerow et al. [2018](#page-131-4)). Dimorphic species are found in *Bannoa*, *Cystobasidium*, *Naohidea*, *Occultifur*, and *Sakaguchia*. Several species were assigned to genera in *Cystobasidiomycetes* based on morphological resemblances, such as the (swollen) probasidium in combination with transversally septate basidia. However, their assignment to this class remains to be confrmed by molecular data. Basidiospore germination is either with hyphae, secondary spore formation, or yeast budding. Yeast colonies are usually orange-pinkish pigmented, except in *Nahoideale* which is characterised by whitish yeast colonies.

**Life modes and distribution:** *Cystobasidiomycetes* have been discovered and isolated from various substrates in terrestrial and marine habitats, including air, freshwater aquatic environments, soil, beetle guts, mammals, sea sponges, other fungi, and plants (Libkind et al. [2010;](#page-145-9) Oberwinkler [2017](#page-149-9); Begerow et al. [2018;](#page-131-4) Jiang et al. [2024](#page-141-9)). Several species in this class are (intrahymenial) mycoparasites characterised by haustorial cells and interact with nanometer fusionpores, establishing cytoplasmic contact between host and mycoparasite (Bauer [2004;](#page-131-8) Bauer et al. [2006;](#page-131-5) Oberwinkler [2017](#page-149-9)). *Occultifur* and *Cyphobasidium* mycoparasites produce typical tremelloid haustorial cells, the haustoria of *Naohidea sebacea* invaginates host cells. *Cyphobasidium* species are found in association with lichen thalli, but the nature of this association is still unclear (Millanes et al. [2016;](#page-147-9) Diederich et al. [2022\)](#page-135-8). Some species from the genera *Cyrenella* and *Microsporomyces* were isolated as endothallic species in *Cladonia* lichens (Kachalkin et al. [2024\)](#page-142-6)

**Notes:** The class *Cystobasidiomycetes* was introduced by Bauer et al. ([2006](#page-131-5)) to include fungi without fucose in their cell walls, and spindle pole bodies that are intranuclear during the metaphase.

**Type order:** *Cystobasidiales* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Progr. 5(1): 46 (2006).

**Type family:** *Cystobasidiaceae* Gäum., Vergl. Morph. Pilze (Jena): 411 (1926).

**Type genus: Cystobasidium** (Lagerh.) Neuhoff, Bot. Arch. 8: 272 (1924).

**Type species:** *Cystobasidium lasioboli* (Lagerh.) Neuhof, Bot. Arch.: 273 (1924).

*Microbotryomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Progr. 5(1): 47 (2006).

Index Fungorum Registration Identifer 501489; Fig. [24g](#page-61-0). The class *Microbotryomycetes* was introduced by Bauer et al. [\(2006](#page-131-5)) as a highly supported clade in molecular phylogenetic reconstructions. It currently accommodates eight accepted orders (*Curvibasidiales*, *Heitmaniales*, *Heterogastridiales*, *Kriegeriales*, *Leucosporidiales*, *Microbotryales*, and *Sporidiobolales*), 13 families (*Camptobasidiaceae*, *Curvibasidiaceae*, *Chrysozymaceae*, *Colacogloeaceae*, *Heitmaniaceae*, *Heterogastridiaceae*, *Kriegeriaceae*, *Leucosporidiaceae*, *Microbotryaceae*, *Mycogloiocolacaceae*, *Rosettozymaceae*, *Sporidiobolaceae*, and *Ustilentylomataceae*), more than 46 genera, and more than 300 species (Begerow et al. [2018](#page-131-4); He et al. [2019](#page-139-1); Li et al. [2020](#page-145-8); Schoutteten et al. [2024](#page-153-9); Jiang et al. [2024\)](#page-141-9). The class is mainly circumscribed by phylogenetic analyses, in which it forms a well-supported monophyletic clade (Aime et al. [2006;](#page-130-7) Bauer et al. [2006](#page-131-5); Wang et al. [2015b](#page-159-6); Li et al. [2020](#page-145-8); Schoutteten et al. [2024\)](#page-153-9).

*Microbotryomycetes* is a class with largely distinct ecological, morphological, and ultrastructural features, and currently, no synapomorphic characters are known that unify all members of this class. Basidiome morphology ranges from effused to pycnidioid, and many mycoparasites develop intrahymelially in the host basidiome and are barely visible or invisible to the naked eye (Toome and Aime [2014](#page-157-6); Malysheva et al. [2021](#page-146-9)). At the ultrastructural level, Microbotryomycetes are characterised by simple septal pores, and intranuclear metaphasic spindle pole bodies (Bauer et al. [2006\)](#page-131-5). Most organisms are known as dimorphic parasites of plants and fungi, but a large number of species are exclusively known as saprobic yeast morphs. These dimorphic fungi alternate between a dikaryotic parasitic flamentous stage, and a haploid saprobic yeast stage. Some exceptions are known, such as *Heterogastridium pycnidioideum* which is only known from a flamentous morph, and for which no yeast budding was observed (Oberwinkler et al. [1990](#page-149-14)). Many parasites in *Microbotryomycetes* do not produce basidiomata, but produce reproductive structures in the tissues of their host organisms (Schoutteten et al. [2024,](#page-153-9) [2024](#page-153-9)). Phytoparasites usually produce sori in specifc plant tissues (Kemler et al. [2020\)](#page-142-7), whereas mycoparasites often develop intrahymenially in the hymenium of their host fungus and are only visible as a slimy layer in fresh condition (Schoutteten et al. [2024\)](#page-153-9). Several species produce peculiar structures, such as the stalked basidiophores in *Krieglsteinera*, the stilboid structures in *Pycnopulvinus*, and the pycnidioid basidiomata of *Heterogastridium*. Basidia are usually transversally septate (except in *Camptobasidium*), and arise from thick-walled diploid teliospores in Microbotryales. In various genera of mycoparasites, the basidia arise directly from hyphae (e.g. *Colacogloea* spp.), or thin-walled probasidia (e.g. *Slooffia micra*). Basidiospores germinate by hyphae, secondary spore formation, or yeast budding.

**Life modes and distribution:** The ecological diversity in this class is large. The largest group of currently known species belongs to the dimorphic phytoparasitic *Microbotryales*. Phytoparasitism occurs in *Kriegeriales* (*Kriegeria eriophori*) and *Microbotryales*. A second ecologically relevant group are the dimorphic mycoparasites. Schoutteten et al. ([2024\)](#page-153-9) showed that mycoparasitism is phylogenetically diverse in this class, with representatives occurring in various orders and families. Mycoparasites in *Microbotryomycetes* engage in colacosome interaction, which is a unique interaction mechanism only found in this class and *Cryptomycocolacomycetes* (Bauer et al. [2006;](#page-131-5) Schoutteten et al. [2024,](#page-153-9) [2024](#page-153-9)). Colacosomes, also known as lenticular bodies, are small structures (0.5  $\mu$ m diam.) produced in mycoparasite hyphae that are adjacent to host hyphae, and comprise a membranebound electron dense core, and an electron transparent sheet engulfng the core (Bauer and Oberwinkler [1991b](#page-131-9)). A secondary cell wall layer is separating the colacosome from the mycoparasite cytoplasm. Colacosomes penetrate the outer cell wall, and the cell wall of the host, which eventually degrades. How the nutrient transfer from host to parasite takes place remains unknown. A third large group of organisms comprises the *Microbotryomycetes* that were isolated as yeasts from a multitude of habitats and substrates, but of which the ecological functions and trophic strategies remain largely unknown. Several of these yeasts are psychrophilic, and were isolated from extreme habitats such as glaciers and crater lakes in Patagonian, Arctic and Antarctic regions (de Garcia et al. [2020;](#page-134-9) Perini et al. [2021](#page-149-15)). It can be expected that many of the *Microbotryomycetes* only known from yeast morphs are in fact dimorphic fungi, and that the flamentous morphs remain to be discovered. Possibly these flamentous morphs are parasitic stages of plants, fungi, or other organisms, but this remains to be investigated.

**Notes:** The phylogenetic distances in *Microbotryomycetes* are usually large, and phylogenetic reconstructions are often characterised by long terminal tip lengths (Bauer et al. [2006](#page-131-5); Wang et al. [2015b](#page-159-6); Schoutteten et al. [2024\)](#page-153-9). Due to the isolated position of various clades, several orders and families were introduced to accommodate a single genus. Although this causes some degree of taxonomic rank infation, this highlights the phylogenetic distance to other known taxa (Schoutteten et al. [2024](#page-153-9)). This is likely refecting the large morphological and ecological diversity, and it is to be expected that only a small fraction of the real diversity in this class is currently known. All previously published phylogenetic studies of this class failed to resolve the relationships between the diferent orders and families. More sampling and genome-scale datasets will be necessary to resolve the relationships within *Microbotryomycetes*. Based on currently known diversity, sampling strategies should focus on extreme habitats, soil, phylloplanes, and colacosome-interacting mycoparasites of other fungi.

**Type order:** *Microbotryales* R. Bauer & Oberw., in Bauer, Oberwinkler & Vánky, Can. J. Bot. 75: 1309 (1997).

**Type family:** *Microbotryaceae* R.T. Moore, Mycotaxon 59: 17 (1996).

**Type genus:** *Microbotryum* Lév., Annls Sci. Nat., Bot., sér. 3 8: 372 (1847).

**Type species:** *Microbotryum antherarum* (DC.) Lév. 1847 (Designated by Pfeifer, Nomenclator Botanicus (Cassellis) 2: 303. 1873) (*fde* Index Fungorum 2023).

*Mixiomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 47 (2006).


<span id="page-72-0"></span>**Fig. 29** *Mixia osmundae* on *Osmunda japonica*. **a**–**c** Ultrathin sections of showing the sporogenous cell from which spores are enteroblastically produced (redrawn from Nishida et al. [1995\)](#page-148-0)

Index Fungorum Registration Identifer 501490; Fig. [29](#page-72-0)

**Asexual morph:** Sporogenous cells developing on the surface of the host epidermal cells, the host cell walls are thickened at the contact area with the hyphal cells; sporogenous cells fusiform, oblong, ellipsoid, pyriform, or bacilliform, without septa. Hyphal cells are multinucleate with electrontransparent, non-fbrillate cell walls; numerous spores are produced exogenously, enteroblastically and simultaneously from the sporogenous cells. The spores are hyaline and ellipsoidal. At the surface of the sporogenous cell, after the release of the spores, numerous tiny denticles were observed using light microscopy. The denticles are cuneiform, ca. 0.5 μm in diameter at the tip and were slightly sunken at the center **Sexual morph:** Unknown (Nishida et al. [1995](#page-148-0); Bauer et al. [2006\)](#page-131-0).

**Life modes and Distribution:** Parasitic on ferns *Osmunda regalis*, *O. japonica* and *Osmundastrum cinnamomeum* (Nishida et al. [1995](#page-148-0); Sugiyama et al. [2018\)](#page-156-0).

**Notes:** The class *Mixiomycetes* was introduced by Bauer et al. [\(2006](#page-131-0)). According to Wijayawardene et al. ([2020a\)](#page-160-0) and He et al. ([2024](#page-139-0)), *Mixiomycetes* comprises one order, *Mixiales*, which include a single family *Mixiaceae* (type: *Mixia*).

*Mixia osmundae* is the sole known species in the *Mixiomycetes*. For more than 80 years, it was identifed as an ascomycete called *Taphrina osmundae*, partly because the sporogenous cells of *Mixia* have some superficial resemblance to the asci produced by certain *Ascomycota* (Aime et al. [2014](#page-130-0)). However, subsequent revaluation based on examination of cell wall ultrastructure and the observation of enteroblastic budding led to its reclassifcation as a *Basidiomycetes* (Boekhout et al. [2011\)](#page-132-0).

According to Wang et al. [\(2015a](#page-159-0)), *Mixiomycetes* is placed phylogenetically as a sister of *Spiculogloeales* which is an order of *Agaricostilbomycetes*. Other analyses by Wang et al. ([2015a](#page-159-0)), Hibbett et al. [\(2007](#page-139-1)), and Aime et al. [\(2014](#page-130-0)) found various placements for *Mixiomycetes*, where it was not sister to *Spiculogloeales*. The identifcation of new taxa within the monotypic class *Mixiomycetes*, as well as the place of additional taxa and sequences from subphylum *Pucciniomycotina*, will be necessary to complete the phylogenetic placement of genus *Mixia*.

**Type order:** *Mixiales* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Progr. 5(1): 47 (2006).

**Type family:** *Mixiaceae* C.L. Kramer, Stud. Mycol. 30: 159 (1987).

**Type genus:** *Mixia* C.L. Kramer, Mycologia 50(6): 924 (1959) [1958].

**Type species:** *Mixia osmundae* (Nishida) C.L. Kramer, Mycologia 50(6): 924 (1959) [1958].

*Pucciniomycetes* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw., Mycol. Prog. 5(1): 48 (2006).

Index Fungorum Registration Identifer 501494; Fig. [30](#page-73-0)

With over 8000 known species, *Pucciniomycetes* is the largest class of *Pucciniomycotina*. This class was introduced by Bauer et al. [\(2006](#page-131-0)), and currently comprises fve orders: *Helicobasidiales Pachnocybales*, *Platygloeales*, *Pucciniales*, and *Septobasidiales*. There exists much ecological and morphological diversity, and no synapomorphic characters are known for this class (Aime et al. [2014;](#page-130-0) Begerow et al. [2018](#page-131-1)). At the ultrastructural level, *Pucciniomycetes* have septal pores which can be occluded by a plug, and are mostly associated with microbodies that delineate an organelle-free zone. Hyphae of *Pucciniomycetes* are clampless. Basidia of *Pucciniomycetes* are usually transversally septate. Basidiospores usually germinate by hyphae, yeast budding is known only from some *Septobasidiales* species (Ober-winkler [2017\)](#page-149-0). The majority of the species belongs to the *Pucciniales*, which are the obligate phytoparasitic rust fungi (Aime et al. [2014](#page-130-0)). The order *Pachnocybales* is monotypic, and contains the presumably saprobic *Pachnocybe ferruginea*. This species is characterised by holobasidia producing



<span id="page-73-0"></span>**Fig. 30** Examples of the members of *Pucciniales*. **a** *Aecidium*-type aecia gathered on leaves. **b** Aeciospores of the family *Pucciniaceae*. **c** Teliospores of the family *Pucciniaceae*. **d**, **g** Telia on leaves. **e**, **h** Urediniospores of the family *Phragmidiaceae*. **f** Teliospores of the

family *Phragmidiaceae*. **i** Teliospores of the *Uredinineae incertae sedis*. **j** Uredinia and telia on leaves. **k** Urediniospores of the family *Melampsoraceae*. **l** Urediniospores of the family *Coleosporiaceae*. Scale bars: **a**, **d**, **g**, **j**=0.5 mm, **b**, **c**, **e**, **f**, **h**, **i**, **k**, **l**=10 µm

sessile statismospores. *Septobasidiales* are entomopathogenic fungi producing dense hyphal mats covering scale insects on their plant hosts. It is generally believed that *Septobasidiales* species parasitize in the scale insects, but the nature of this association remains to be better understood (Aime et al. [2014](#page-130-0)). *Septobasidiales* produce transversally septate basidia, and the basidiospores can germinate by yeast budding. *Platygloeales* are all characterised by transversally septate basidia, and mainly comprise parasites of ferns and mosses (e.g. *Eocronartium*, *Herpobasidium*, *Jola*, *Insolibasidium*, *Platycarpa*, and *Ptechelitium*), but also some saprobic fungi such as *Platygloea disciformis*. *Helicobasidiales* species have a dual ecological strategy. The sexual morphs of these species are characterised by transversally septate basidia, and are known as the '*Helicobasidium* stage', in which they are phytoparasitic. The asexual morphs are mycoparasites of rusts (*Pucciniales*), and were previously classifed in the genus *Tuberculina*. During the mycoparasitic stage, these fungi interact with their host fungus via micrometer fusion-pore interaction, in which migration of nuclei and mitochondria from the mycoparasite to the host cell was observed (Bauer [2004;](#page-131-2) Bauer et al. [2004;](#page-131-3) Lutz [2004](#page-146-0)).

**Notes:** Much diversity in the *Pucciniomycetes* remains to be discovered. The best-investigated group is the phytoparasitic *Pucciniales* because of their economic importance (Aime et al. [2014](#page-130-0)). The known diversity in the other orders is much smaller, and few taxonomic treatments are available covering the genera in these groups. The number of currently accepted *Helicobasidiales* species is low compared to the number of potential host species in the order *Pucciniales*. Many names are available for *Tuberculina*, and this complex genus is in urgent need of a taxonomic revision in which the host range of the diferent species is assessed (Table [19](#page-74-0)).

**Type order:** *Pucciniales* Caruel, Atti R. Accad. Naz. Lincei, Mem. Cl. Sci. Fis. Matem. Nat., sér. 5: 246 (1881). **Type family:** *Pucciniaceae* Chevall. [as 'Puccinieae'], Fl. gén. env. Paris (Paris) 1: 413 (1826).

**Type genus:** *Puccinia* Pers., Neues Mag. Bot. 1: 118 (1794).

**Type species:** *Puccinia graminis* Pers. 1794 (Designated by Laundon, Mycol. Pap. 99: 15. 1965).

*Spiculogloeomycetes* Q.M. Wang, F.Y. Bai, M. Groenew. & Boekhout (2015).

Index Fungorum Registration Identifer: 813101.

*Spiculogloeomycetes* belongs to the subphylum *Pucciniomycotina* (Bauer et al. [2006;](#page-131-0) Wang et al. [2015b](#page-159-1); Oberwinkler [2017](#page-149-0)). The class comprises a single order, *Spiculogloeales*, accommodating the genera *Meniscomyces*, *Phyllozyma*, and *Spiculogloea* (Wang et al. [2015b](#page-159-1); Oberwinkler [2017;](#page-149-0) Li et al. [2020\)](#page-145-0). The majority of the species belonging to *Spiculogloeomycetes* are known only from a yeast morph and no flamentous morphs with sexual structures are known from the species in *Meniscomyces* and *Phyllozyma*, which were all isolated from plant surfaces. Yeast colonies are usually yellow to pale yellowish-brown, occasionally pinkish-orange, budding cells are present, and hyphae and pseudohyphae are present or not. Ballistoconidia are fusiform, ellipsoidal, or sometimes sickle-shaped in *Meniscomyces* spp.

*Spiculogloea* species on the other hand are intrahymenial mycoparasites known from flamentous morphs only and generate hyphae with clamp collections, three-septate auricularioid basidia that are spiculated at the surface producing ballistospores, and sometimes conidiophores with cyanophilous conidia (Roberts [1996,](#page-151-0) [1997](#page-151-1); Langer and Oberwinkler [1998](#page-144-0); Bauer et al. [2006;](#page-131-0) Oberwinkler [2017](#page-149-0); Schoutteten et al. [2018,](#page-153-0) 2024). The basidiospores can germinate by hyphae, yeast budding, or secondary spore production. Furthermore, *Spiculogloea* species are characterised by the presence of clamped haustorial cells, of which the globular apex attaches to hyphae of the host fungus. Along the contact surface, one or more nanopores are produces, establishing a cytoplasmic bridge between the host and mycoparasite (Weiß et al. [2004;](#page-159-2) Bauer et al. [2006](#page-131-0)). Cultures and DNA sequence data are unavailable for most of the *Spiculogloea* species, and thus, their systematic placement remains to be investigated. Several phylogenetic reconstructions found the



<span id="page-74-0"></span>

clustering of a *Mycogloea* isolate in *Spiculogloeomycetes*, although only LSU and SSU sequence data are available and the obtained support is usually low (Weiß et al. [2004](#page-159-2); Bauer et al. [2006;](#page-131-0) Wang et al. [2015b\)](#page-159-1). Hence, it is unclear whether this *Mycogloea* truly belongs to *Spiculogloeomycetes*, or is a member of a diferent class in *Pucciniomycotina*.

**Life modes and distribution:** *Spiculogloeomycetes* are comprised of teleomorphic members (*Spiculogloea* spp.) that produce basally clamped transversally three-septate basidia and clamped tremelloid haustorial cells. *Spiculogloea* spp. are intrahymenial mycoparasites of corticioid fungi and resupinate jelly fungi (*Agaricomycetes*), and interact with their host fungi by means of nanofusionpore interaction. These mycoparasites are mainly known from North-America and Europe, but it can be expected that these mycoparasites follow the distribution of their hosts (Bandoni et al. [2002;](#page-130-1) Spirin et al. [2016](#page-155-0); Oberwinkler [2017](#page-149-0); Schoutteten et al [2018](#page-153-0), 2024). Some taxa are isolated as anamorphic yeast colonies from plant surfaces (mainly the phyllosphere), and their ecological role remains poorly understood (e.g. *Meniscomyces* spp. and *Phyllozyma* spp.). These species were isolated from various parts of the world, including South-America, Japan and China (Wang et al. [2015b](#page-159-1); Oberwinkler [2017](#page-149-0); Li et al. [2020](#page-145-0)).

**Notes:** The placement of *Spiculogloeomycetes* was confrmed by a phylogenetic analysis using seven genes in which it formed a well-supported lineage sister to *Mixiomycetes* within *Pucciniomycotina* (Bauer et al. [2006](#page-131-0); Wang et al. [2015b](#page-159-1)). *Spiculogloeomycetes* are poorly studied, and the little overlap between the teleomorphic and anamorphic data allows for further studies.

**Type order:** *Spiculogloeales* R. Bauer, Begerow, J.P. Samp., M. Weiss & Oberw. 2006.

**Type family:** *Spiculogloeaceae* Denchev, Mycol. Balcanica 6: 87 (2009).

**Type genus:** *Spiculogloea* P. Roberts, Mycotaxon 60: 112 (1996).

**Type species:** *Spiculogloea occulta* P. Roberts, Mycotaxon 60: 113 (1996).

*Tritirachiomycetes* Aime & Schell, in Schell, Lee & Aime, Mycologia 103(6): 1339 (2011).

Index Fungorum Registration Identifer: 561067; Fig. [31](#page-76-0)

**Asexual morph:** Multinucleate hyphae, simple pore septa, conidiophores that are subhyaline to dematiaceous and subhyaline to dematiaceous sympodial conidiogenous cells that occur in whorls and bear conidia on an elongated rachis. **Sexual morph:** Unknown (Schell et al. [2011\)](#page-153-1).

**Life modes and distribution:** The members of *Tritirachiomycetes* are worldwide in distribution and reported as saprophytes (Moraes et al. [2010](#page-148-1); Schell et al. [2011](#page-153-1); Naseri et al. [2013\)](#page-148-2), biotrophic mycosymbionts (Beguin [2010](#page-131-4)), endophytes (Vieira et al. [2011;](#page-158-0) Bezerra et al. [2013](#page-132-1); Qadri et al. [2014\)](#page-150-0), and as human opportunistic pathogens (Martínez-Herrera et al. [2015;](#page-146-3) Vanam et al. [2018;](#page-158-1) Bezerra et al. [2020\)](#page-132-2), e.g. *Tritirachium oryzae* is important species that causes human infections (Martínez-Herrera et al. [2015](#page-146-3); Vanam et al. [2018](#page-158-1)).

**Notes:** The class *Tritirachiomycetes* was introduced by Schell et al. ([2011](#page-153-1)). Multi-gene phylogenetic analyses (LSU, SSU, ITS, and  $tef$ - $I\alpha$ ) and septal pore ultrastructure confrmed its placement in *Pucciniomycotina* (Schell et al. [2011\)](#page-153-1). He et al. ([2019](#page-139-2)) provided the divergence time estimates for *Tritirachiomycetes*. Wijayawardene et al. [\(2022a\)](#page-160-1) and He et al. ([2024](#page-139-0)) accepted the class *Tritirachiomycetes* comprises with one order (i.e. *Tritirachiales*), one family (*Tritirachiaceae*), two genera (*Paratritirachium* and *Tritirachium*), and six species.

**Type order:** *Tritirachiales* Aime & Schell, in Schell, Lee & Aime, Mycologia 103(6): 1339 (2011).

**Type family:** *Tritirachiaceae* Aime & Schell, in Schell, Lee & Aime, Mycologia 103(6): 1339 (2011).

=*Tritirachiaceae* Locq., Mycol. gén. struct. (Paris): 208 (1984).

**Type genus:** *Tritirachium* Limber, Mycologia 32(1): 24 (1940).

**Type species:** *Tritirachium dependens* Limber, Mycologia 32: 26 (1940).

**Subphylum** *Ustilaginomycotina* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVIII (2001).

**Notes:** Doweld ([2001](#page-135-0)) introduced *Ustilaginomycotina* that currently consists of fve classes; 1. *Exobasidiomycetes*; 2. *Malasseziomycetes*; 3. *Moniliellomycetes*. 4. *Peribolosporomycetes* and 5. *Ustilaginomycetes* (He et al. [2024\)](#page-139-0).

*Exobasidiomycetes* Begerow, M. Stoll & R. Bauer, Mycologia 98(6): 908 (2007) [2006].

Index Fungorum Registration Identifer 501484; Fig. [24e](#page-61-0) Simple spores and interaction apparatus (Bauer et al. [1997](#page-131-5); Begerow et al. [2006](#page-131-6)).

**Life modes and distribution:** Parasitic smut fungi (e.g. infecting Asparagaceae, Melanthiaceae, and Xanthorrhoeaceae), yeasts, widespread.



<span id="page-76-0"></span>**Fig. 31** *Tritirachium* sp. **a**–**c** *T. dependens.* **d**, **e** *T. album*. Scale bars: **a**–**e**=10 μm (redrawn from Limber [1940](#page-145-2))

**Notes:** Bauer et al. [\(1997](#page-131-5)) introduced *Exobasidiomycetidae* to accommodate *Doassansiales* R. Bauer & Oberw., *Entylomatales* R. Bauer & Oberw., *Exobasidiales* Henn., *Georgefscheriales* R. Bauer et al., *Graphiolales* Oberw., *Microstromatales* R. Bauer & Oberw., and *Tilletiales* Haeckel. However, Begerow et al. [\(2006](#page-131-6)) introduced *Exobasidiomycetes* and regarded *Exobasidiomycetidae* as a synonym of the former. Begerow et al. [\(2006\)](#page-131-6) also introduced *Ceraceosorales* Begerow et al. as a new order in *Exobasidiomycetes*. Subsequent studies by Wang et al. ([2015a](#page-159-0)) and Li et al. [\(2022](#page-145-1)) introduced *Golubeviales* Q.M. Wang et al. and *Franziozymales* Q.M. Wang et al. hence, currently, the class comprises ten orders.

**Type order:** *Exobasidiales* Henn., in Engler & Prantl, Nat. Pfanzenfam., Teil. I (Leipzig) 1(1\*\*): 103 (1898) [1900]. **Type family:** *Exobasidiaceae* J. Schröt. [as 'Exobasidiacei'], in Cohn, Krypt.-Fl. Schlesien (Breslau) 3.1(25–32): 413 (1888) [1889].

**Type genus:** *Exobasidium* Woronin, Verh. Naturf. Ges. Freiburg 4(4): 397 (1867).

**Type species:** *Exobasidium vaccinii* (Fuckel) Woronin, Verh. Naturf. Ges. Freiburg 4(4): 397 (1867).

*Malasseziomycetes* Denchev & T. Denchev, Index Fungorum 145: 1 (2014).

Index Fungorum Registration Identifer 550570; Fig. [24f](#page-61-0) Zoophilic members of the *Ustilaginomycotina* have a monopolar budding yeast phase showing percurrent or sympodial proliferation of the buds. Yeasts are lipid-dependent or lipophilic (excluding the case of *Malassezia pachydermatis*), with a multilayered cell wall and a helicoidal evagination of the plasma membrane. **Sexual morph:** Unknown (Denchev and Denchev [2014\)](#page-135-1).

**Life modes and distribution:** Parasitic on the skin of warmblooded animals (Gupta et al. [2004](#page-138-0); Begerow et al. [2006\)](#page-131-6) and widespread (Kirk et al. [2008](#page-142-0)).

**Notes:** Denchev and Denchev [\(2014](#page-135-1)) introduced *Malasseziomycetes* to accommodate the single-order *Malasseziales*. Currently, the class accommodates one order, one family, and one genus.

**Type order:** *Malasseziales* R.T. Moore [as 'Malesseziales'], Bot. Mar. 23(6): 371 (1980).

**Type family:** *Malasseziaceae* Denchev & R.T. Moore, Mycotaxon 110: 379 (2009).

**Type genus:** *Malassezia* Baill., Traité Bot. Méd. Crypt.: 234 (1889).

**Type species:** *Malassezia furfur* (C.P. Robin) Baill., Traité Bot. Méd. Crypt.: 234 (1889).

*Moniliellomycetes* Q.M. Wang, F.Y. Bai & Boekhout [as 'Monilielliomycetes'], in Wang, Theelen, Groenewald, Bai & Boekhout, Persoonia 33: 46 (2014).

Index Fungorum Registration Identifer 823105; Fig. [32](#page-78-0)

**Asexual morph:** Chlamydospore formation of *Moniliella* occurred when the yeast is transferred from nutrient-rich to poor media, or in nitrogen-depleted media (de Hoog et al. [2011;](#page-135-2) Thanh et al. [2013](#page-157-0)). Pseudohyphae and chlamydospores are sometime present, cell walls are multi-lamellar. Hyphal septa typically possess dolipores with an arch of endoplasmic reticulum, but 'micropore'-like structures may also be present (de Hoog et al. [2011\)](#page-135-2). Sugars are fermented by most species. Nitrate is assimilated. Urease and diazonium blue B (DBB) reactions are positive. Coenzyme Q-9 is present. Xylose and fucose are absent from whole-cell hydrolysates (Wang et al. [2014\)](#page-159-3). **Sexual morph:** Unknown. *Colonies* are restricted, smooth, velvety or cerebriform, cream then paleolivaceous or black-brown, yeast-like growth, budding cells are ellipsoidal and form terminally on true hyphae that disarticulate with arthroconidia (Wang et al. [2014\)](#page-159-3).

**Life modes and distribution:** The members of *Moniliellomycetes* are distributed in a broad range of ecosystems in North and South America, Asia, and Europe in diferent life modes. Yeasts in *Moniliellomycetes* have been isolated from man-made products such as sweet fruit sauce, sweet pickles, acetic acid-containing products, honey, jam, leaf-cutting bees, honey bees, brood cells of the honey bees, pollen in honeycomb, bufalo milk, fowers (e.g. *Ipomoea pes-caprae* and *Calotropis gigantea*), fermented food, rotting wood, meat processing environments, and air (Rosa et al. [2009;](#page-152-0) de Hoog et al. [2011](#page-135-2); Thanh et al. [2012](#page-157-1), [2013;](#page-157-0) Lei et al. [2022](#page-144-1); Pipiková et al. [2023](#page-150-1)), or pathogenetic on humans and cats (McKenzie [1984](#page-147-0); Guarro et al. [1999](#page-138-1)). Species may be osmophilic and lipophilic, and are used to produce erythritol, a low-calorie sweetener (de Hoog et al. [2011](#page-135-2)).

**Notes:** The monotypic class *Moniliellomycetes* belongs to the subphylum *Ustilaginomycotina*, and has only one order (*Moniliellales*) and one family (*Moniliellaceae*), with *Moniliella* as the type genus (Wang et al. [2014\)](#page-159-3). The phylogeny of *Moniliella*, was not well understood for a long time (de Hoog et al. [2011](#page-135-2)). The genus was initially thought to be closely related to *Tremellomycetes* in *Agaricomycotina* (de Hoog [1979](#page-134-0); Guarro et al. [1999](#page-138-1)). *Moniliella* includes black, yeast-like fungi in the *Basidiomycota*, but morphologically similar black, yeast-like fungi also occur in *Ascomycota* (Yurlova et al. [2008](#page-161-0)). Despite their relative uniformity in morphological appearance and physiological properties, species of *Moniliella* exhibit a high degree of genetic divergence (Rosa et al. [2009\)](#page-152-0).

**Type order:** *Moniliellales* Q.M. Wang, F.Y. Bai & Boekhout, in Wang, Theelen, Groenewald, Bai & Boekhout, Persoonia 33: 46 (2014).

**Type family:** *Moniliellaceae* Q.M. Wang, F.Y. Bai & Boekhout, in Wang, Theelen, Groenewald, Bai & Boekhout, Persoonia 33: 46 (2014).

**Type genus:** *Moniliella* Stolk & Dakin, Antonie van Leeuwenhoek 32: 399 (1966).

**Type species:** *Moniliella acetoabutans* Stolk & Dakin [as 'acetoabutens'], Antonie van Leeuwenhoek 32: 400 (1966).

*Peribolosporomycetes* Witfeld, M.A. Guerreiro, H.D.T. Ngyuen & Begerow, in Witfeld, Guerreiro, Nitsche, Wang, Ngyuen & Begerow, Mycol. Progr. 22(4, no. 30): 10 (2023).

Index Fungorum Registration Identifer 843630.

Class of mesophilic, heat resistant, and osmotolerant basidiomycetes with slow hyphal growth and high phenotypic variability. Characteristically, triangular-shaped chlamydospores are distally produced on coiled hyphae. Sympodial, ovoid conidia are produced. Relatively simple septal pores, without thickened septum membranes. Parenthesomes, forming distinct septal pore caps are absent, but low contrasted elements indicate non-membranous structures around the pore openings and within the pore. Based on the analysis of 38 protein-coding genes, orthology analysis, and septal pore type analysis, this class is placed within the *Ustilaginomycotina* (Witfeld et al. [2023](#page-160-2)).

**Life modes and distribution:** From soil, associated with *Pinus resinosa*. Known from Nova Scotia (Witfeld et al. [2023](#page-160-2)).

**Notes:** Witfeld et al. ([2023](#page-160-2)) introduced *Peribolospora* Witfeld et al. (type: *P. baueri* Witfeld et al.) and showed that this genus has a distinct phylogenetic lineage in *Ustilaginomycotina*. Hence, they introduced new family (*Peribolosporaceae*), order (*Peribolosporales*) and class (*Peribolosporomycetes*).



<span id="page-78-0"></span>**Fig. 32** Morphology of *Moniliella pyrgileucina* (redrawn from Thanh et al. [2018\)](#page-157-2)

**Type order:** *Peribolosporales* Witfeld, M.A. Guerreiro, H.D.T. Ngyuen & Begerow, in Witfeld, Guerreiro, Nitsche, Wang, Ngyuen & Begerow, Mycol. Progr. 22(4, no. 30): 11 (2023).

**Type family:** *Peribolosporaceae* Witfeld, M.A. Guerreiro, H.D.T. Ngyuen & Begerow, in Witfeld, Guerreiro, Nitsche, Wang, Ngyuen & Begerow, Mycol. Progr. 22(4, no. 30): 10 (2023).

**Type genus:** *Peribolospora* Witfeld, M.A. Guerreiro, H.D.T. Ngyuen & Begerow, in Witfeld, Guerreiro, Nitsche, Wang, Ngyuen & Begerow, Mycol. Progr. 22(4, no. 30): 11 (2023).

**Type species:** *Peribolospora baueri* Witfeld, M.A. Guerreiro, H.D.T. Ngyuen & Begerow, in Witfeld, Guerreiro, Nitsche, Wang, Ngyuen & Begerow, Mycol. Progr. 22(4, no. 30): 11 (2023).

*Ustilaginomycetes* Warm., Haandb. syst. Bot., Edn 2 (Kjøbenhavn): 57 (1884).

Index Fungorum Registration Identifer: 816414.

Free-living saprobic phase with yeast and pathogenic on many plants, typically grasses of the family Poaceae, shrubs, some ferns, and conifers, or parasites on animals*.*

**Sexual morph:** *Sori* surrounding stems, inforescence, ovaries, and organs, long-cylindrical, circular, subcircular, angular to tubular or curved, hypertrophied sometimes bearing small remnants of deformed glumes or as leaf spots, whitish to brown or yellow-green with or without peridium, peridium greyish to brown splitting longitudinally to expose powdery spore mass, sori not usually present in all organs, partitioning cells may be present or absent (Piepenbring [2001](#page-150-2); Shivas et al. [2006;](#page-154-0) McTaggart et al. [2016\)](#page-147-1). *Columellae* sometimes present or absent, stout or woody, fagelliform or fliform. *Spore balls* colourless to pigmented, irregular to globose to slightly fattened when connected to adjacent balls, composed of compacted masses of spores, friable or persistent with or without cortex, lobed, and with a central part consisting of parenchymatous sterile cells in the air canals of the mesophyll. *Teliospores* correspond to probasidia, thick-walled, dark-coloured, sometimes smooth-walled, usually ornamented with warts and ridges, arranged singly or in pairs, multicellular or in pairs, in groups or balls, elongate, globose, subglobose, sometimes broadly ellipsoidal, ovoid, subcuneiform, boomerang shaped, sometimes irregular, collapsed, hemispherically cupped, sub-angular, oblong, club-shaped to oval, threelayered (perisporium, episporium, endosporium), pale olivaceous brown to reddish brown in colour (Rattan et al. [1995](#page-151-2); Vánky et al. [2009;](#page-158-2) Hu et al. [2018\)](#page-140-0). Teliospores germinate into a cylindrical, four-celled basidium that buds as basidiospores. *Basidia* club-shaped, phragmobasidial, or holobasidial, with or without septa, respectively. *Basidiospores* numerous, fliform, uninucleate, mostly ballistosporic, and others are passively released (Bauer et al. [2001;](#page-131-7) Hu et al. [2018](#page-140-0)).

**Asexual morph:** hyphomycetous or yeast-like. **Hyphomycetous** *Conidiophores* solitary or aggregated, semimacronematous to micronematous. *Conidiogenous cells* monoblastic, basally clamped, single conidia derived by completely transferring protoplasm. *Conidia* navicular, obclavate, or fusiform to ellipsoidal on short sterigmata-like structures with 3–4 branches, sometimes narrow at the base, along with many bristle-like distal branches. *Chlamydospores* present. (Marvanova and Bandoni 1987; Mekha et al. [2014\)](#page-147-2). **Yeast** budding yeast-like phase, with the presence of hyphae and pseudohyphae, cells spherical, ellipsoidal to fusoid-shaped on short denticles, cream to light salmon in colour (Piątek et al. [2017\)](#page-150-3).

**Life modes and distribution:** The members of *Ustilaginomycetes* are distributed in a wide range of habitats and are reported to follow diferent life modes (Boekhout [1995](#page-132-3); Avis et al. [2001\)](#page-130-2). They are mostly phytopathogenic, mostly infecting members of the grass families *Poaceae* and *Cyperaceae*, and are responsible for causing smut diseases and huge economic losses (e.g., *Ustilago maydis* and *U. nuda*) (Kruse et al. [2017\)](#page-143-0) (Table [20](#page-79-0)). *Melanotaenium oreophilum* and *M. selaginellae* are two smuts found to be associated with *Selaginella* (Bauer et al. [1999\)](#page-131-8). Rarely, some yeast morphs of the members have also been reported to infect humans (Teo and Tay [2006](#page-157-3); McNeil and Palazzi [2012\)](#page-147-3) (Table [21](#page-80-0)). Some of the members have been isolated from the leaves, fruits, and fowers of plants; there are also reports of isolation from human secretory fuid or blood (Sugita et al. [2003](#page-156-2); Arendrup et al. [2014\)](#page-130-3). *Moesziomyces antarcticus* has been isolated from soil and plants and rarely from human blood (Boekhout [2011](#page-132-4)). *Moesziomyces aphidis* was frst isolated from an aphid and was later also isolated from water (Boekhout [2011](#page-132-4)). Rarely, a few species of *Ustilaginomycetes* parasitize ferns or conifers (Bauer et al. [1999](#page-131-8), [2001\)](#page-131-7). Riess et al. ([2016](#page-151-3)) reported *Uleiella chilensis* to be parasitic on gymnosperms like *Araucaria araucana*.

**Notes:** Warming ([1884\)](#page-159-4) introduced the class *Ustilaginomycetes* of the phylum *Basidiomycota*. Bauer et al. [\(1997\)](#page-131-5), on the basis of ultrastructural markers, revealed that *Ustilaginomycetes* form three lineages belonging to the phyto-parasitic members that lack dolipore with multilayered pore-bearing caps. Bauer et al. ([2006\)](#page-131-0) and Hibbett et al. ([2007\)](#page-139-1), in their phylogenetic classifcation of fungi, renamed the clade *Ustilaginomycetes* as in Kirk et al. [\(2008\)](#page-142-0) as *Ustilaginomycotina* to minimize confusion between the names of the taxon and informal terms. Tedersoo et al. [\(2018](#page-157-4)), in their classifcation of the fungi, placed the class *Ustilaginomycetes* under the subphylum *Ustilaginomycotina*.

Hyde et al.  $(2011)$  $(2011)$  $(2011)$ , in an initial attempt to compile asexually typifed genera among the members of the Kingdom *Fungi*, incorporated anamorphic fungi in the class <span id="page-79-0"></span>**Table 20** Examples of economically important *Ustilaginomycetes*



*Ustilaginomycetes* according to the natural system of classifcation. Later, Wijayawardene et al. [\(2012](#page-159-5), [2017b](#page-160-3), [2021b\)](#page-160-4) published an updated checklist along with notes and entries on the pleomorphic genera in *Ustilaginomycetes*. Aime et al. [\(2018](#page-130-4)) recommend the use of generic names in place of synonymous alternate asexually typifed morph generic names for several members of *Ustilaginomycetes*, such as *Anthracoidea* for *Crotalia*, *Mycosarcoma* or *Ustilago* for *Pseudozyma*, *Thecaphora* for *Rhombiella* and *Thecaphorella*, and *Urocystis* for *Paepalopsis*.

According to Kijpornyongpan et al. ([2018\)](#page-142-1), the class comprises three orders: *Ustilaginales*, *Urocystidales*, and *Uleiellales* (Bauer et al. [1997;](#page-131-5) Riess et al. [2016](#page-151-3)). The order *Ustilaginales* contains eight families, 49 genera, and 851 species (Kirk et al. [2008\)](#page-142-0). *Urocystidales* has six families and 400 taxa (Begerow et al. [2006](#page-131-6), [2014\)](#page-131-9), and the order *Uleiellales* is comprised of one family (*Uleiellaceae*), a single genus (*Uleiella*), and two species (Riess et al. [2016](#page-151-3)). There are no recognized subclasses under *Ustilaginomycetes*.

Many pleomorphic taxa among the members of *Ustilaginomycetes* have been reported (Kirk et al. [2008](#page-142-0)). Wang et al. [\(2015a](#page-159-0)) proposed taxonomic revisions to adopt one name (between two generic names) when the links between the morphs are confrmed in *Ustilaginomycetes*. Since the discontinuation of dual names for fungi (Hawksworth et al. [2011\)](#page-139-3), Wang et al. [\(2015a\)](#page-159-0) and Kruse et al. ([2017\)](#page-143-0) have made important publications dealing with the recent nomenclatural changes.

Phylogenetic analyses of the rRNA gene (SSU, ITS, LSU, *atp6*, and *tub* sequence data) of Hibbett et al. ([2007\)](#page-139-1) and Begerow et al. [\(2006](#page-131-6)) accommodated this class in subphylum *Ustilaginomycotina.* He et al. [\(2019\)](#page-139-2) provided the divergence time estimates for the various orders and families in the class *Ustilaginomycetes*. According to Wijayawardene et al. ([2022a\)](#page-160-1), the class comprises four orders (*Uleiellales*, *Urocystidales*, *Ustilaginales*, and *Violaceomycetales*), 15 families, and 68 genera. However, two genera,

## <span id="page-80-0"></span>**Table 21** Medicinally/clinically signifcant *Ustilaginomycetes* in humans



*Capitulocladosporium* and *Eriocortex* are classifed in *Ustilaginomycetes* genera *incertae sedis.*

**Type order:** *Ustilaginales* Bek., Kurs Botaniki 1: 221 (1864).

**Type family:** *Ustilaginaceae* Tul. & C. Tul. [as 'Ustilagineae'], Annls Sci. Nat., Bot., sér. 3 7: 14 (1847).

**Type genus:** *Ustilago* (Pers.) Roussel, Fl. Calvados, Edn 2: 47 (1806).

**Type species:** not indicated (see editorial comments in Index Fungorum 2023).

**Subphylum** *Wallemiomycotina* Doweld, Index Fungorum 73: 1 (2014).

**Notes:** Doweld ([2014j\)](#page-135-3) introduced *Wallemiomycotina* to accommodate a single phylum, i.e. *Wallemiomycetes*. He et al. [\(2024\)](#page-139-0) accepted this placement.

*Wallemiomycetes* Zalar, de Hoog & Schroers, Antonie van Leeuwenhoek 87(4): 322 (2005).

Index Fungorum Registration Identifer 501496.

**Sexual morph:** *Basidiomata* absent; *Basidia* singly or in clusters, irregular in shape, arising from primary cells with a probasidium; single sterigmata or sterigmata-like cells present; *basidiospores* produced by some genera; single basidiospore in one basidium; globose, rarely subglobose; hirsute; hyaline; double-walled; browning on maturity (Nguyen et al. [2013\)](#page-148-4). **Asexual morph: hyphomycetous.** *Conidiophores* solitary or aggregated, unbranched, erect or sometimes sympodial, smooth-walled, and little constricted below the apex. *Conidiogenous cells* continuing from conidiophores, verruculose, cylindrical, biaxially extending, disarticulating basipetally, four arthrospore-like conidia, forming chains. *Conidia* single-cell, verruculose or spherical, short cylindrical, thick-walled, and pale brown (Zalar et al. [2005](#page-161-1); Jančič et al. [2015](#page-141-0); Díaz-Valderrama et al. [2017\)](#page-135-4).

**Life modes and distribution:** The members of *Wallemiomycetes* are cosmopolitan in distribution and are xerophilic and halophilic in nature. The members are airborne and found to be associated with solar salts, salt crystals, dried plant parts such as hay and seeds, dried food, and are also responsible for food spoilage and agricultural aerosols (Zalar et al. [2005](#page-161-1); Díaz-Valderrama et al. [2017](#page-135-4); Zajc and Gunde-Cimerman [2018](#page-161-2)). Some strains are known to cause subcutaneous skin infections in humans (de Hoog et al. [2000](#page-134-1)) and hypersensitivity pneumonitis (farmer's lung disease), caused by *Wallemia sebi* (Lappalainen et al. [1998](#page-144-2); Roussel et al. [2005](#page-152-1)).

**Notes:** The class *Wallemiomycetes* was introduced by Zalar et al. [\(2005\)](#page-161-1) and is one of the smallest classes of *Basidiomycota*. Matheny et al. ([2006\)](#page-146-4) attempted to resolve the phylogenetic position of *Wallemiomycetes* using nuclear protein-coding genes. Zhao et al. ([2017\)](#page-161-3) tried to resolve the phylogenetic obscurity of the class based on the concepts of molecular data.

The class *Wallemiomycetes* is placed in phylogenetic trees in slightly diferent ways depending on which gene regions are analysed and which phylogenetic inference method is used (Zalar et al. [2005](#page-161-1); Matheny et al. [2006](#page-146-4)). Analyses of the parsimony of the nrDNA data revealed that *Wallemiomycetes* is an early diverging branch of the *Basidiomycota* and that it occupies a basal position close to *Entorrhizomycetidae* (Matheny et al. [2006\)](#page-146-4). He et al. [\(2024\)](#page-139-0) confrmed that *Wallemiomycetes* belongs to the subphylum *Wallemiomycotina.* The class comprises two orders, *Geminibasidiales* and *Wallemiales* (Wijayawardene et al. [2022a](#page-160-1); He et al. [2024](#page-139-0)).

Some members of *Wallemiomycetes* are medicinally important taxa. *Wallemia sebi* causes allergies, bronchial asthma, farmer's lung disease (in humans), chronic ulcerative skin lesions, and fatal livestock toxicosis (*fde* Cole and Samson [1979](#page-133-0); Hanhela et al. [1995;](#page-138-2) Lappalainen et al. [1998,](#page-144-2) [2012;](#page-144-3) Reboux et al. [2001;](#page-151-4) Roussel et al. [2005](#page-152-1); Zajc and

Gunde-Cimerman [2018\)](#page-161-2). *Wallemia muriae* causes farmer's lung disease and bronchial asthma in humans (*fde* Zajc and Gunde-Cimerman [2018](#page-161-2)). *Wallemiomycetes mellicola* causes phaeohyphomycosis on the foot in immunocompetent human (Zajc and Gunde-Cimerman [2018](#page-161-2)).

**Type order:** *Wallemiales* Zalar, de Hoog & Schroers, Antonie van Leeuwenhoek 87(4): 322 (2005).

**Type family:** *Wallemiaceae* R.T. Moore, Rhizoctonia Species, Taxonomy, Molecular Biology, Ecology, Pathology and Disease Control (Dordrecht): 20 (1996).

**Type genus:** *Wallemia* Johan-Olsen, Skr. VidenskSelsk. Christiania, Kl. I, Math.-Natur. (no. 12): 6 (1887).

**Type species:** *Wallemia ichthyophaga* Johan-Olsen [as 'ichthiophaga'], Skr. VidenskSelsk. Christiania, Kl. I, Math.- Natur.(no. 12): 6 (1887).

*BLASTOCLADIOMYCOTA* T.Y. James, Mycologia 98(6): 867 (2007) [2006].

Index Fungorum Registration Identifer 501464.

Thallus monocentric or polycentric, may form hyphae, varies greatly in size and extent or degrees of complexity; zoospore with a single posteriorly directed whiplash-type fagellum possessing a lipid-flled organelle often called a side-body complex (SBC) composed of several lipid droplets, characterized by the presence of distinctive anteriorly located membrane-bounded ribosomal nuclear cap adjacent to the cone-shaped nucleus which abuts the kinetosome, cytoplasmic microtubules extending anteriorly from the proximal (upper) part of the kinetosome (the basal body of the fagellum) to surround the nuclear envelopes, absence of electron opaque plug in the transitional zone at the base of zoospore fagellum. Asexual reproduction through unifagellated zoospores produced in zoosporangium, sexual reproduction by planogamete (sexual gamete) fusion and life cycle with sporic meiosis displaying alternation of gametophytic and sporophytic generations (James et al. [2006b\)](#page-141-1).

**Life mode and distribution:** Saprobic on a wide variety of animal and plant substrates in both aquatic and terrestrial habitats. Some representatives are parasites of algae, fungi, plants and invertebrates of freshwater and soil environments, and occasionally found to be facultatively anaerobic that grow under foul conditions (Sparrow [1960](#page-155-1); Karling [1977](#page-142-2); James et al. [2006a,](#page-141-2) [b](#page-141-1), [2014;](#page-141-3) Hibbett et al. [2007;](#page-139-1) Powell [2017](#page-150-4)).

**Notes:** James et al. [\(2006b\)](#page-141-1) introduced the phylum *Blastocladiomycota* since it formed a monophyletic clade, distinct from other early divergent lineages in the Kingdom *Fungi*. *Blastocladia* Reinsch (Reinsch [1877](#page-151-5)) the type genus of the phylum *Blastocladiomycota* was accommodated in its own family, *Blastocladiaceae* by Petersen ([1910](#page-150-5)). Later, the Order *Blastocladiales* was introduced by Kanouse ([1927](#page-142-3)). Fitzpatrick ([1930\)](#page-136-0) regarded *Blastocladiales* in the class *Phycomycetes* (i.e. 'A class formerly used for *Fungi* and now treated in Chromista (q.v.) and some fungi (*Chytridiomycota* and *Zygomycota*) *fde* Kirk et al. [2008](#page-142-0)). However, Alexopoulos (1966) accommodated *Blastocladiales* in *Chytridiomycetes*.

Tehler ([1988](#page-157-6)) used the term '*Blastocladiomycota'*, as a division of 'Subkingdom *Eumycota'* but did not provide type details or descriptions thus invalid. Doweld [\(2001\)](#page-135-0) introduced *Blastocladiomycota* Doweld [2001](#page-135-0) but invalid (Art. 38.1(a) (Shenzhen); 'Published as 'nom. nov.', based on *Allomycotina* Caval.-Sm., 1998 (David, Constancea 83.16: 5) *fde* Index Fungorum 2023). Hence, James et al. ([2006b\)](#page-141-1) provided proper diagnosis and description of the phylum thus, *Blastocladiomycota* T.Y. James 2006 [2007] is the correct name. Subsequent studies agreed with the conclusions in James et al. ([2006b](#page-141-1)) and confrmed its phylogenetic placement in the fungal tree (e.g. Hibbett et al. [2007\)](#page-139-1). Porter et al. [\(2011](#page-150-6)) showed the family placements of *Blastocladiomycota* but did not assign/ mention any subphyla or classes but their tree clearly showed the separation of *Physodermataceae*. Agreeing with Porter et al. [\(2011\)](#page-150-6), Tedersoo et al. [\(2018\)](#page-157-4) introduced Subphylum *Blastocladiomycotina* Tedersoo et al. and the Class *Physodermatomycetes* Tedersoo et al. Following studies by Wijayawardene et al. [\(2018b,](#page-160-6) [2020a\)](#page-160-0) and Voigt et al. ([2021\)](#page-158-3) also agreed with the higher-level classifcation proposed by Tedersoo et al. [\(2018](#page-157-4)).

Currently comprises one subphylum (*Blastocladiomycotina*) and two classes.

*Blastocladiomycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Isonym: *Blastocladiomycetes* T.Y. James, Mycologia 98 (6): 868 (2007).

Index Fungorum Registration Identifer 90742; Fig. [33](#page-82-0)

Thallus monocentric, polycentric or flamentous (sometimes presenting pseudosepta); zoospore with a single posteriorly oriented whiplash fagellum that lacks electron-opaque shield present in the kinetosome transition region, characterized by the presence of lateral membrane cisterna known as side-body complex (SBC), membrane-bound ribosomal aggregation known as a ribosomal cap (nuclear cap) containing a single cone-shaped nucleus that terminates adjacent to the kinetosome, cytoplasmic microtubules with anteriorly radiating roots leading from the proximal portion of the kinetosome continuing on to wrap around a nucleus; resting sporangium darkly pigmented and usually ornamented with spines or other projections. Asexual reproduction through posteriorly unifagellate chemotactic zoospores (fagellated asexual spores), sexual reproduction through planogamete fusion with a sporic meiosis type of life cycle involving



<span id="page-82-0"></span>**Fig. 33** Microscopic characteristics of *Allomyces* spp. (*Blastocladiaceae*, *Blastocladiales*, *Blastocladiomycetes*). **a** Mature thallus of *Allomyces neomoniliformis* with some empty zoosporangia and resistant sporangia. **b** Young developing thallus of *Allomyces moniliformis.* **c** Dark brown and thick-walled resistant sporangia of *Allo-*

alternation of a haploid gametophytic (which produces gametes) with a diploid sporophytic (produces asexual spores) generation (James et al. [2006b\)](#page-141-1).

**Life mode and distribution:** The members of *Blastocladiomycetes* are widely distributed in a broad range of aquatic and terrestrial ecosystems throughout the world and are reported commonly as saprotrophs, feeding on decomposing organic matter, with diferent types of life cycle. This very large and morphologically diverse class also contains many parasites of various soil inhabiting small invertebrates, their eggs and fungi in water-logged soil or water (Sparrow [1960](#page-155-1); Karling [1977](#page-142-2); James et al. [2014;](#page-141-3) Powell [2017](#page-150-4)).

*myces neomoniliformis.* **d**–**f** Hypha with thin-walled zoosporangia in *Allomyces moniliformis.* **g**–**i** Thin-walled zoosporangia with papillae of *Allomyces moniliformis*. Scale bars:  $a = 60 \mu m$ ,  $b = 100 \mu m$ ,  $c = 40 \text{ µm}, d - f = 100 \text{ µm}, g - i = 50 \text{ µm}$ 

**Notes:** Doweld ([2001](#page-135-0)) introduced this class to accommodate *Blastocladia* Reinsch [1877](#page-151-5) (*Blastocladiaceae*, *Blastocladiales*). James et al. ([2006b](#page-141-1)) also introduced *Blastocladiomycetes* but it is an isonym of *Blastocladiomycetes* Doweld. Currently, the class comprises three orders i.e. **1.** *Blastocladiales* (families *Blastocladiaceae* H.E. Petersen, *Catenariaceae* Couch, *Paraphysodermataceae* Doweld, and *Sorochytriaceae* Dewel); **2.** *Callimastigales* Doweld (families: *Callimastigaceae* Fonseca); **3.** *Catenomycetales* Doweld (*Catenomycetaceae* Doweld, *Coelomomycetaceae* Couch).

**Type order:** *Blastocladiales* H.E. Petersen [as 'Blastocladiineae'], Bot. Tidsskr. 29: 357 (1909).

**Type family:** *Blastocladiaceae* H.E. Petersen, Bot. Tidsskr. 29: 357 (1909).

**Type genus:** *Blastocladia* Reinsch, Jb. wiss. Bot. 11(2): 291 (1877).

**Type species:** *Blastocladia pringsheimii* Reinsch [as 'pringsheimi'], Jb. wiss. Bot. 11(2): 367 (1877).

*Physodermatomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Divers. 90: 148 (2018).

Index Fungorum ID: 553995; Fig. [34](#page-84-0)

Thallus monocentric to polycentric, eucarpic, epibiotic with slightly branched and limited rhizoids generally arising from a single axis near the base of the thin-walled sporangium, with endogenous (intracellular) development; Golgi apparatus or dictyosome with stacked cisternae and vesicles in sporangium; zoospores produces bipolar growth; the posteriorly unifagellate zoospores present a characteristic nuclear cap of ribosomes (membrane-bound aggregated ribosomes) with adjacent nucleus attached to centriole; resting sporangium thick-walled and typically darkly pigmented; germination and development of sporangium through protruding endosporangium (Tedersoo et al. [2018\)](#page-157-4).

**Life mode and distribution:** Representatives within this class are common obligate parasites of microalgae, pteridophytes and angiosperms in aquatic and marshy habitats throughout the world (Sparrow [1960\)](#page-155-1).

**Notes:** Tedersoo et al. ([2018](#page-157-4)) introduced this class to accommodate *Physoderma* Wallr. (*Physodermataceae*, *Physodermatales*). Characterized by bearing both monocentric thallus with an endobiotic polycentric thallus. Spores germinate producing zoospores that infect the specifc host. The initial infestation generates the monocentric, epibiotic zoosporangium that is attached to endobiotic rhizoids that are limited to a single host cell where the zoosporangium is characterized as *Rhizidium* or *Phlyctochytrium*. The zoospores liberate and infect healthy new host cells, where the infestation transmits through number of generations. The internally proliferous sporangia produce a second fush of zoospores after the liberation of the frst round. The zoospores develop into an endobiotic polycentric thallus which is often extensive. The highly branched thallus with rhizoids infects a large number of host cells. These rhizoids possess intercalary cells, which contain septa are evident for sexual reproduction. The endobiotic thallus produces large, dark-coloured resting spores with thick-wall and with the shape of host cells. Resting spores appear as formed by intercalary cells (Sparrow [1962;](#page-155-2) Karling [1977](#page-142-2); James et al. [2014;](#page-141-3) Powell [2017](#page-150-4)).

**Type order:** *Physodermatales* Caval.-Sm., Eur. J. Protist. 49:157.

**Type family:** *Physodermataceae* Sparrow, Mycologia 44(6): 768 (1952).

**Type genus:** *Physoderma* Wallr., Fl.crypt. Germ. (Norimbergae) 2: 192 (1833).

**Type species:** *Physoderma maculare* Wallr., Fl. crypt. Germ. (Norimbergae) 2: 192 (1833).

*CALCARISPORIELLOMYCOTA* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Divers. 90: 152 (2018).

Index Fungorum Registration Identifer 554019.

Members are characterized with a branched thallus with septate hyphae. The vegetative hyphae smooth and thin, thin-walled and hyaline. In the case of presence of sporangiophores are hyaline, smooth, simple and originated from undiferentiated hyphae. The sporangia are unispored, ellipsoid with or without a tiny columella. Spores hyaline, thinwalled, smooth, uninucleate, ovoid, with a rounded base. At the presence of chlamydospores, characterized with single cell, thick spiny walls, elongated to globose, born laterally on short hyphae. The sexual morph is unknown (Hirose et al. [2012](#page-139-4); Zhao et al. [2022\)](#page-162-0).

**Life modes and distribution:** Saprobic in soil and nonnematophagous in soils.

**Notes:** Tedersoo et al. ([2018\)](#page-157-4) showed that *Calcarisporiella thermophila*, the type species of *Calcarisporiella* and *Echinochlamydosporium variabile*, the type species of *Echinochlamydosporium* formed a distinct phylogenetic lineage in the Kingdom *Fungi*. Hence, Tedersoo et al. ([2018\)](#page-157-4) introduced *Calcarisporiellomycota*.

*Calcarisporiellomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 152 (2018).

Index Fungorum Registration Identifer: 554020; Fig. [35](#page-85-0) Similar to the phylum description.

**Notes:** *Calcarisporiellomycetes* comprises only one order and one family i.e. *Calcarisporiellales*, *Calcarisporiellaceae*, resepectively. *Calcarisporiella* and *Echinochlamydosporium* are the genera contained by *Calcarisporiellomycetes*. *Calcarisporiella* is a monotypic anamorphic genus and was previously considered to be an ascomycetous anamorphic genus belong to *Pezizomycotina* based on morphology of the anamorph. Besides, the spore formation process, and the presence of septate hyphae were other concerns for the revisions of the classifcation (Zhao et al. [2005](#page-161-4); Kirk et al. [2008](#page-142-0)).

<span id="page-84-0"></span>**Fig. 34** *Physoderma marsileae* **a** Petiole showing very gall histology and normal tissue. Resting spores (1), endodermis (2), protoxylem (3), pericycle (4),×100. **b** Leafet showing gall histology and normal tissue and expanded portion of leafet and row of resting spores (5), and one-half of palisade cells (6),×100. **c** Mature resting spores (7) within host cells, showing rhizoidal system (8) and turbinate organs (9). Also large central body in resting spore (10),×1790. **d** Resting spore showing wrinkled area (11) and concave area (12) with antler-like haustorial processes (13),×1790. **e** Mature resting spore showing wrinkled area and supraequatorial ring of pits  $(14)$ ,  $\times$ 1790 (redrawn from Brewster [1952](#page-132-6))



**Type order:** *Calcarisporiellales* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 152 (2018).

**Type family:** *Calcarisporiellaceae* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 152 (2018).

**Type genus:** *Calcarisporiella* de Hoog, Stud. Mycol. 7: 68 (1974).

**Type species:** *Calcarisporiella thermophila* (H.C. Evans) de Hoog, Stud. Mycol. 7: 68 (1974).

*CHYTRIDIOMYCOTA* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001);

=*Caulochytriomycota* Doweld, Index Fungorum 49: 1 (2014).

Index Fungorum Registration Identifer 550346; Fig. [36](#page-86-0)

Fungi with eucarpic, monocentric or polycentric thallus, endogenous or exogenous development; some with plasmodial thallus; asexual reproduction usually by zoospores posteriorly unifagellate, nine ciliary props, microbody–lipid globule complex (MLC) consisting of a cisterna which may



<span id="page-85-0"></span>**Fig. 35** *Calcarisporiella thermophila* (Basionym=*Calcarisporium thermophile*) (holotype, CBS 279.70). **a** Conidiogenous cells. **b** Conidia. **c** Infated hyphal cell in culture at 45 °C. Scale bar **a**, **b**, **c**=10 mm (redrawn from de Hoog [1974](#page-134-2))

be simple or fenestrated, microbodies and mitochondria associated with lipid globules; Golgi apparatus with stacked cisternae; nuclear envelope fenestrated at poles during mitosis; resting spores, when present, formed asexually or sexually by conjugation of thalli (Hibbett et al. [2007](#page-139-1); Powell and Letcher [2014](#page-150-7); Adl et al. [2019](#page-129-0)).

**Life modes and distribution:** Saprotrophic in aquatic or terrestrial ecosystems, or facultative and obligate parasites of algae, amphibians, fungi, protists and plant tissues.

**Notes:** Doweld ([2001\)](#page-135-0) introduced the phylum *Chytridiomycota*. Several major studies, before 2010 concluded that *Chytridiomycota* taxa are polyphyletic (e.g. James et al. [2006a,](#page-141-2) [b\)](#page-141-1). Thus, Hibbett et al. [\(2007\)](#page-139-1) restricted the phylum to include only two classes, i.e. *Chytridiomycetes* and *Monoblepharidomycetes*. Tedersoo et al. ([2018\)](#page-157-4) however elevated *Monoblepharidomycetes* to phylum as *Monoblepharomycota* thus not belonging to *Chytridiomycota*. Furthermore, Tedersoo et al. [\(2018\)](#page-157-4) introduced eight new classes in *Chytridiomycota*; viz*., Cladochytriomycetes* Tedersoo et al., *Lobulomycetes* Tedersoo et al., *Mesochytriomycetes* Tedersoo et al., *Polychytriomycetes* Tedersoo et al., *Rhizophlyctidomycetes* Tedersoo et al., *Rhizophydiomycetes* Tedersoo et al., *Spizellomycetes* Tedersoo et al. and *Synchytriomycetes* Tedersoo et al. Interestingly, James et al. ([2006b\)](#page-141-1) showed that the classes introduced by Tedersoo et al. [\(2018](#page-157-4)) are well-separated orders in their analyses, except for *Mesochytriomycetes* and *Lobulomycetes*. Seto et al. ([2017\)](#page-154-1) confrmed the results



<span id="page-86-0"></span>**Fig. 36** *Chytridiomycota* (*Chytridiales*). **a** Resting spore of *Asterophlyctis sarcoptoides* on shrimp exoskeleton. **b** Zoosporangium of *Phlyctochytrium aureliae* on snake skin. **c**, **d** *Chytriomyces hyalinus* on onion skin. **c** Zoosporangium. **d** Resting spore. **e**, **f** *Entophlyctis luteolus* on onion skin. **e** Zoosporangium. **f** Resting spore. **g**, **h** *Fayochytriomyces spinosus* on onion skin. **g** Zoosporangium. **h** Empty

of James et al. [\(2006b\)](#page-141-1) and confrmed the separation of *Mesochytriales* and *Lobulomycetales*.

Tedersoo et al. ([2018\)](#page-157-4) introduced the Subkingdom *Chytridiomyceta* Tedersoo et al. to accommodate *Chytridiomycota* along with *Monoblepharomycota* and *Neocallimastigomycota*. Furthermore, Tedersoo et al. [\(2018](#page-157-4)) introduced *Chytridiomycotina* Tedersoo et al., the only subphylum in *Chytridiomycota*.

Recent higher-level classifcation studies showed that *Caulochytrium protostelioides* (only representative taxon of

zoosporangium. **i** Zoosporangium discharge of *Rhizidium crepaturum* on shrimp exoskeleton. **j** Zoosporangium discharge of *Rodmanochytrium sphaericum* on shrimp exoskeleton. **k**, **l** *Rhizoclosmatium globosum* on shrimp exoskeleton. **k** Empty zoosporangium. **l** Resting spore. *Ap*apophysis; *Op*operculum. Scale bars: **a**–**l**=10 µm

*Caulochytriomycota*) nested in *Chytridiomycota* (Ahrendt et al. [2018](#page-130-5); Strassert and Monaghan [2022a,](#page-156-3) [b](#page-156-4); supplementary tree). The type species lacks genomic data but here we agree with previous studies and our phylogenomic analysis and regard *Caulochytriomycota* as a synonym of *Chytridiomycota* and regard *Caulochytriomycetes* as a class in *Chytridiomycota.* Hence, currently, *Chytridiomycota* comprises ten classes.

*Caulochytriomycetes* Doweld, Index Fungorum 49: 1 (2014).

Index Fungorum Registration Identifer 550345 (Fig. [37](#page-131-10)). Sporangia thin-walled, of two types, sessile and hollowstalked aerial. Zoospores posteriorly unifagellate, fagellum laterally inserted, fagellar insertion subapical to parabasal. Rhizoplast striated, attaches the proximal end of the kinetosome to a specialized region of the nuclear envelope; no nuclear cap; microbody-lipid globule complex comprises several lipid globules, branching microbody, membrane cisternae and circular mitochondria. Pulsating vacuolar system associated with the fagellar apparatus and endoplasmic reticulum with dictyosomes and coated vesicles (Doweld [2014a\)](#page-135-5).

**Life modes and distribution:** Saprobic (Voos and Olive [1968\)](#page-158-4) and parasitic on the mycelium of *Cladosporium* on *Acalypha hispida* (Olive [1980\)](#page-149-3).

**Notes:** Doweld ([2014a\)](#page-135-5) introduced this class to accommodate *Caulochytrium* (*Caulochytriaceae, Caulochytriales*). Traditionally, *Caulochytrium* was accommodated in *Chytridiomycota* by several studies (e.g. Subramanian [1974](#page-156-5); Olive [1980](#page-149-3), [1983\)](#page-149-4). However, Hibbett et al. [\(2007](#page-139-1)) mentioned the placement of the genus is uncertain. Doweld [\(2014a\)](#page-135-5) introduced the subphylum *Caulochytriomycotina* and *Caulochytriomycota*. However, Galindo et al. ([2021\)](#page-137-0) and Strassert and Monaghan ([2022a](#page-156-3), [b](#page-156-4)) and our analyses (Fig. [1\)](#page-10-0) showed that *Caulochytrium protosteliodes* (non-type species) resided in *Chytridiomycota*.

**Type order:** *Caulochytriales* Doweld, Index Fungorum 49: 1 (2014).

**Type family:** *Caulochytriaceae* Subram., Curr. Sci. 43: 723 (1974).

**Type genus:** *Caulochytrium* Voos & L.S. Olive, Mycologia 60(3): 731 (1968).

**Type species:** *Caulochytrium gloeosporii* Voos & L.S. Olive, Mycologia 60(3): 731 (1968).

*Chytridiomycetes* Caval.-Sm., Biol. Rev. 73:246. 1998, emend. Tedersoo et al., Fungal Divers. 90: 148 (2018).

Index Fungorum Registration Identifer 816490.

Thallus eucarpic, monocentric or polycentric, endobiotic, interbiotic or epibiotic; plasmodial thallus sometimes present. Zoosporangium endogenous or exogenous to the zoospore cyst, inoperculate or operculate, apophysate or non-apophysate. Zoospores, when present, posteriorly unifagellate with a non-fagellum centriole and a fagellum kinetosome, typically with kinetosome-associated structures; the microbody-lipid globule complex (MLC) cisterna is adjacent to the lipid globule. Resting spores, when present, formed asexually or sexually by conjugation of thalli (Sparrow [1960](#page-155-1); Doweld [2014g](#page-135-6); Powell and Letcher [2014](#page-150-7); Letcher et al. [2018](#page-145-3); Adl et al. [2019](#page-129-0); Voigt et al. [2021](#page-158-3)).

**Life modes:** Saprobic in aquatic or terrestrial ecosystems, or usually facultative and obligate parasites of plants, animals, protists and algae.

**Notes:** Cavalier-Smith ([1998\)](#page-133-2) provided a brief Latin description of *Chytridiomycetes* which was emended by Tedersoo et al. ([2018](#page-157-4)). The class comprises four orders; viz*., Chytridiales* Cohn, *Nephridiophagales* Doweld, *Polyphagales* Doweld and *Saccopodiales* Doweld (Wijayawardene et al. [2020a,](#page-160-0) [2022a;](#page-160-1) Voigt et al. [2021](#page-158-3)). *Chytridiales* is composed of four families based on ultrastructural and molecular studies, *Asterophlyctaceae*, *Chytridiaceae*, *Chytriomycetaceae* and *Pseudorhizidiaceae*, with 26 genera and more than 330 species. Members of *Phlyctochytriaceae* and *Phlyctorhizaceae* were transferred to other families within the order (James et al. [2006b](#page-141-1); Vélez et al. [2011](#page-158-5); Letcher et al. [2012](#page-144-4); Vélez et al. [2013](#page-158-6); Letcher and Powell [2014](#page-144-5); Leshem et al. [2016](#page-144-6); Letcher and Powell [2018](#page-144-7); Letcher et al. [2018;](#page-145-3) Powell et al. [2018](#page-150-8); Seto and Degawa [2018;](#page-154-2) Jesus et al. [2021](#page-141-4)). The members of *Scherfelliomycetaceae* are lacking sequences thus, classifcation is uncertain.

*Nephridiophagales*, erected by Doweld ([2014g](#page-135-6)), is composed of *Nephridiophagaceae*, with four genera and 19 species (Wijayawardene et al. [2018b;](#page-160-6) Voigt et al. [2021](#page-158-3)). *Polyphagales* was created by Doweld ([2014h\)](#page-135-7) to include *Polyphagaceae*, currently with two genera and 16 species (Index Fungorum 2023). *Saccopodiales* was also erected by Doweld [\(2014i\)](#page-135-8) to accommodate *Saccopodiaceae*, with one monotypic genus.

**Type order:** *Chytridiales* Cohn, Jber. schles. Ges. vaterl. Kultur 57: 279 (1879).

**Type family:** *Chytridiaceae* Bek., Kurs Botaniki 1: 271 (1864).

**Type genus:** *Chytridium* A. Braun, Betracht. Erschein. verjüng. Natur, Edn 2 (Leipzig): 198 (1851).

**Type species:** *Chytridium olla* A. Braun, Betracht. Erschein. verjüng. Natur, Edn 2 (Leipzig): 198 (1851).

*Cladochytriomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 148 (2018).

Index Fungorum Registration Identifer 553997; Fig. [38](#page-101-0)

Thallus eucarpic, monocentric or polycentric; rhizoids catenulate, isodiametric or tapering. Zoospore chytridioid but with a cord-like microtubular root between the kinetosome and fenestrated cisterna, composed of up to 25 microtubules interconnected by linkers; a cisterna, microbody, and mitochondrion closely associated with the lipid globule; mostly saprotrophic or pathogenic on algae (extracted from Tedersoo et al. [2018\)](#page-157-4).

**Life modes:** Mostly saprotrophic or pathogenic on algae.

**Notes:** James et al. [\(2006b\)](#page-141-1) and Seto et al. ([2017](#page-154-1)) recognized *Cladochytriales* as a distinct clade in *Chytridiomycota.* Hence, Tedersoo et al. ([2018\)](#page-157-4) upgraded *Cladochytriales* (including fve families i.e. *Catenochytridiaceae* Doweld, *Cladochytriaceae* J. Schröt., *Endochytriaceae* Sparrow ex D.J.S. Barr, *Nowakowskiellaceae* Sparrow ex Mozl.-Standr. and *Septochytriaceae* Mozl.-Standr.) to class level. Voigt et al. ([2021](#page-158-3)) also agreed with the new class, *Cladochytriomycetes* and confrmed its placement in *Chytridiomycota*.

Currently, the class comprises only one order, i.e. *Cladochytriales* Mozl.-Standr.

**Type order:** *Cladochytriales* Mozl.-Standr., Mycol. Res. 113(4): 502 (2009).

**Type family:** *Cladochytriaceae* J. Schröt., in Engler & Prantl, Nat. Pfanzenfam., Teil. I (Leipzig) 1(1): 80 (1897). **Type genus:** *Cladochytrium* Nowak., in Cohn, Beitr. Biol. Pf. 2: 92 (1877) [1876].

**Type species:** *Cladochytrium tenue* Nowak., Beitr. Kenntn. Chytr.: 92 (1877) [1876].

*Lobulomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 148 (2018).

Index Fungorum Registration Identifer 553999; Fig. [39](#page-102-0)

Thallus monocentric, eucarpic, with endogenous development; zoospore with opaque fagellar plug, anterior or posterior plug extensions; one or two lipid globules; lacking microtubule root, Golgi apparatus, striated inclusion, and electron opaque bodies near kinetosome (extracted from Simmons et al. [2009\)](#page-155-3).

**Life modes and habitats:** Saprobes, present in soil, dung, marine and freshwater habitats (Simmons et al. [2009,](#page-155-3) [2012](#page-155-4)).

**Notes:** Simmons et al. [\(2009\)](#page-155-3) showed that the new genus, *Lobulomyces* D. R. Simmons (type species: *Lobulomyces angularis* (Longcore) D. R. Simmons) has a distinct phylogenetic lineage in the Phylum *Chytridiomycota*, thus, introduced the family *Lobulomycetaceae* D. R. Simmons and the order *Lobulomycetales* D. R. Simmons, to accommodate the genus. Subsequent studies by Simmons et al. ([2012](#page-155-4)) (who introduced *Alogomyces* D.R. Simmons & Letcher) and Seto and Degawa ([2015](#page-154-3)) (who introduced *Cyclopsomyces* K. Seto & Degawa) confrmed the phylogenetics placement of *Lobulomycetales* in *Chytridiomycota*.

Tedersoo et al. [\(2018\)](#page-157-4), however, introduced *Lobulomycetes* Tedersoo et al., a separate class to accommodate the subclade of *Lobulomycetales* in the Phylum *Chytridiomycota*. Wijayawardene et al. [\(2020a](#page-160-0)) and Voigt et al. ([2021\)](#page-158-3) agreed with this conclusion.

**Type order:** *Lobulomycetales* D.R. Simmons, Mycol. Res. 113(4): 453 (2009).

**Type family:** *Lobulomycetaceae* D.R. Simmons, Mycol. Res. 113(4): 453 (2009).

**Type genus:** *Lobulomyces* D.R. Simmons, Mycol. Res. 113(4): 454(2009).

**Type species:** *Lobulomyces angularis* (Longcore) D.R. Simmons, Mycol. Res. 113(4): 454 (2009).

*Mesochytriomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 148 (2018).

Index Fungorum Registration Identifer 553998.

Thallus simple, with inoperculate, monocentric, epibiotic sporangium having endogenous development and slightly branched rhizoids near the sporangial base; zoospore centriole at an angle of ca. 30 degrees to kinetosome (extracted from Tedersoo et al. [2018\)](#page-157-4).

**Life modes and distribution:** Parasites of freshwater algae, cosmopolitan.

**Notes:** The order *Mesochytriales* Doweld was introduced by Doweld ([2013b](#page-135-9)) with the family *Mesochytriaceae* Doweld. Karpov et al. [\(2014c](#page-142-4)) also introduced *Mesochytriales* Karpov & Aleoshin but it is an isonym. Moreover, Karpov et al. [\(2014c\)](#page-142-4) confrmed the phylogenetic placement of the order *Mesochytriales* and *Gromochytriales* Karpov & Aleoshin, in *Chytridiomycota*. However, Tedersoo et al. ([2018\)](#page-157-4) established a new class, *Mesochytriomycetes* to accommodate *Mesochytriales* and *Gromochytriales.* This arrangement was accepted by Voigt et al.  $(2021)$  $(2021)$ .

Currently, the class comprises two orders, viz*., Mesochytriales* and *Gromochytriales.*

**Type order:** *Mesochytriales* Doweld, Index Fungorum 45: 1 (2013).

**Type family:** *Mesochytriaceae* Doweld, Index Fungorum 45: 1 (2013).

**Type genus:** *Mesochytrium* B.V. Gromov, Mamkaeva & Pljusch, Nova Hedwigia 71(1–2): 159 (2000).

**Type species:** *Mesochytrium penetrans* B.V. Gromov, Mamkaeva & Pljusch, Nova Hedwigia 71(1–2): 159 (2000).

*Polychytriomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 148 (2018).

Index Fungorum Registration Identifier 554000; Fig. [40c](#page-103-0)-e

Thallus polycentric or monocentric; monocentric species with multiple rhizoidal axes. Motile zoospores spherical, usually  $>4$  um diam., with or without flagellar plug and kinetosome spur; 0–3 microtubule roots present; nonfagellated centriole equal to or longer than diameter and attached to kinetosome throughout its length; cultures grow on chitin (extracted from Tedersoo et al. [\(2018\)](#page-157-4), which was prepared using Longcore and Simmons ([2012\)](#page-146-5) and Powell and Letcher [\(2014\)](#page-150-7)).

**Life modes and distribution:** Mostly in soil and freshwater, cosmopolitan.

**Notes:** Longcore and Simmons [\(2012](#page-146-5)) introduced the order *Polychytriales* in *Chytridiomycota* since *Polychytrium* Ajello, the type of the order, is segregated from the other members of the phylum. Tedersoo et al. ([2018](#page-157-4)) upgraded the order to a class.

**Type order:** *Polychytriales* Longcore & D.R. Simmons, Mycologia 104(1): 279 (2012).

**Type family:** *Polychytriaceae* Doweld, Index Fungorum 89: 1 (2014).

**Type genus:** *Polychytrium* Ajello, Mycologia 34(4): 442 (1942).

**Type species:** *Polychytrium aggregatum* Ajello, Mycologia 34(4): 442 (1942).

*Rhizophlyctidomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 148 (2018).

Index Fungorum Registration Identifer 554001.

Monocentric and eucarpic thallus; sporangium inoperculate or endo-operculate, interbiotic, with one or several discharge apparatus and multiple rhizoids; zoospores with a non-fagellated centriole positioned at an acute angle to the kinetosome, without microtubule roots and microtubules; organelles of the 'microbody-lipid globule complex (MLC) loosely arranged (Powell and Letcher [2014\)](#page-150-7).

**Life modes and distribution:** Commonly saprotrophic on cellulosic substrates in the terrestrial ecosystems.

**Notes:** *Rhizophlyctidomycetes* was erected by Tedersoo et al. [\(2018\)](#page-157-4). This class is composed of the order *Rhizophlyctidales*, which was proposed based on molecular monophyly and zoospore ultrastructure of *Rhizophlyctis rosea*-like isolates by Letcher et al. [\(2008a,](#page-144-8) [b](#page-144-9)). Four monophyletic families, with diferent types of ultrastructural organization of the zoospores, are into this order, *Arizonaphlyctidaceae*, *Borealophlyctidaceae*, *Rhizophlyctidaceae* and *Sonorophlyctidaceae*, which have four genera, *Arizonaphlyctis*, *Borealophlyctis*, *Rhizophlyctis* and *Sonoraphlyctis*, respectively, and around 30 species (Powell and Letcher [2014](#page-150-7)).

**Type order:** *Rhizophlyctidales* Letcher, in Letcher, Powell, Barr, Churchill, Wakefield & Picard, Mycol. Res. 112(9): 1034 (2008).

**Type family:** *Rhizophlyctidaceae* H.E. Petersen [as 'Rhizophlyctaceae'], Bot. Tidsskr. 29: 357 (1909).

**Type genus:** *Rhizophlyctis* A. Fisch., in Winter, Rabenh. Krypt.-Fl., Edn 2 (Leipzig) 1(4): 114 (1892).

**Type species:** *Rhizophlyctis rosea* (de Bary & Woronin) A. Fisch., in Winter, Rabenh. Krypt.-Fl., Edn 2 (Leipzig) 1(4): 122 (1892).

*Rhizophydiomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T.May, M. Ryberg & Abarenkov, Fungal Diversity 90: 149 (2018).

Index Fungorum Registration Identifer 554002; Fig. [40a](#page-103-0)

Thallus monocentric; ribosomes enclosed by a system of double membranes; mitochondria, microbodies, lipid globules, and membrane cisterna are typically associated as a microbody-lipid globule complex. The nonfagellated centriole and kinetosome lie parallel or slightly angled toward each other and are connected by fbrillar material. The base of the fagellum proper lacks an electron-opaque plug (extracted from Tedersoo et al. [2018](#page-157-4) who modifed the descriptions from Letcher et al. [2006](#page-144-10); Powell and Letcher [2014](#page-150-7)).

**Life modes and habitats:** parasites and saprobes mostly in soil and freshwater.

**Notes:** The order *Rhizophydiales* was introduced by Letcher et al. [\(2006](#page-144-10)) and was accepted by James et al. [\(2006b\)](#page-141-1) and Seto et al. [\(2017](#page-154-1)). Tedersoo et al. ([2018](#page-157-4)) introduced *Rhizophydiomycetes* to accommodate *Rhizophydiales*. Compare to other classes in *Chytridiomycota*, *Rhizophydiomycetes* is rich in genera and families (James et al. [2006b;](#page-141-1) Wijayawardene et al. [2020a](#page-160-0)).

**Type order:** *Rhizophydiales* Letcher, in Letcher, Powell, Churchill & Chambers, Mycol. Res. 110(8): 908 (2006). **Type family:** *Rhizophydiaceae* Letcher, in Letcher, Powell,

Churchil & Chambers, Mycol. Res. 110(8): 909 (2006).

**Type genus:** *Rhizophydium* Schenk ex Rabenh., Flora Europaea algarum aquae dulcis et submarinae 3: 280 (1868) [1867].

**Type species:** *Rhizophydium globosum* (A. Braun) Rabenh., Flora Europaea algarum aquae dulcis et submarinae 3: 280 (1868) [1867].

*Spizellomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 149 (2018).

Index Fungorum Registration Identifer 554003; Fig. [40](#page-103-0)b

Thallus monocentric, eucarpic; sprorangium inoperculate; nucleus of zoospores associated directly or indirectly with kinetosome; rumposomes absent; replacement of the translation elongation factor 1-alpha gene by elongation factor-like gene in genome (extracted from Tedersoo et al. [\(2018\)](#page-157-4) who provided the modifed descriptions from Barr  $(1980)$  and Powell and Letcher  $(2014)$  $(2014)$  $(2014)$ ).

**Life modes and habitats:** mostly saprobes in soil and parasites of animals, fungi and stramenopiles.

**Notes:** Barr ([1980](#page-131-11)) introduced *Spizellomycetaceae* D.J.S. Barr and *Spizellomycetales* D.J.S. Barr to accommodate *Spizellomyces* D.J.S. Barr. The phylogenetic placement of *Spizellomycetales* in *Chytridiomycota* was broadly accepted (e.g. James et al. [2006b;](#page-141-1) Seto et al. [2017](#page-154-1)). Tedersoo et al. [\(2018](#page-157-4)) upgraded the order to class rank. Currently, the class consists of one order and two families.

**Type order:** *Spizellomycetales* D.J.S. Barr, Can. J. Bot. 58: 2384 (1980).

**Type family:** *Spizellomycetaceae* D.J.S. Barr, Can. J. Bot. 58(22): 2384 (1980).

**Type genus:** *Spizellomyces* D.J.S. Barr, Can. J. Bot. 58(22): 2384 (1980).

**Type species:** *Spizellomyces punctatus* (W.J. Koch) D.J.S. Barr [as 'punctatum'], Can. J. Bot. 58(22): 2384 (1980).

*Synchytriomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 149 (2018).

Index Fungorum Registration Identifer 554004.

Thallus endobiotic, holocarpic, in a form of a resting spore or sorus surrounded by a membrane, colonial in some stages of development; zoospores posterior, unifagellate; with a single lipid globule surrounded by cisternae of endoplasmic reticulum and microbodies; gamma-like vesicles present; nuclear cap lacking; two kinetosomes almost in parallel, transversely striated; dictyosome solitary, associated with posterior rumposome; fagellar apparatus comprises kinetosome and secondary centriole; fagellar terminal plate biconcave if present (extracted from Tedersoo et al. [2018](#page-157-4) who provided the description based on Doweld [\(2014b](#page-135-10)) and Powell and Letcher [\(2014](#page-150-7))).

**Life modes and habitats:** mostly pathogens of terrestrial plants.

**Notes:** James et al. [\(2006a,](#page-141-2) [b](#page-141-1)) and Sekimoto et al. [\(2011\)](#page-154-4) confrmed the placement of *Synchytrium* in *Chytridiomycota*.

Doweld [\(2014b](#page-135-10)) introduced the order *Synchytriales* Doweld in *Chytridiomycota*. Powell and Letcher ([2014\)](#page-150-7), Smith et al. [\(2014](#page-155-5)) and Longcore et al. ([2016\)](#page-146-6) accepted *Synchytriales* as a distinct order in *Chytridiomycetes*. Tedersoo et al. ([2018\)](#page-157-4) upgraded *Synchytriales* to a class as *Synchytriomycetes.* Wijayawardene et al. ([2020a,](#page-160-0) [2022a\)](#page-160-1) and Voigt et al. ([2021\)](#page-158-3) agreed with the conclusion of Tedersoo et al. ([2018](#page-157-4)).

Currently, *Synchytriomycetes* comprises only one order, i.e. *Synchytriales*.

**Type order:** *Synchytriales* Doweld, Index Fungorum 92: 1 (2014).

**Type family:** *Synchytriaceae* J. Schröt., in Engler & Prantl, Nat. Pfanzenfam., Teil. I (Leipzig) 1(1): 71 (1897).

**Type genus:** *Synchytrium* de Bary & Woronin, Verh. Naturf. Ges. Freiburg 3(2): 46 (1863).

**Type species:** *Synchytrium taraxaci* de Bary & Woronin, Verh. Naturf. Ges. Freiburg 3(2): 46 (1863).

*ENTOMOPHTHOROMYCOTA* Humber, Mycotaxon 120: 481 (2012).

Index Fungorum Registration Identifer 564375.

Conidiophores rising from mycelia or host body, usually phototropic, simple or apically branched, with an apical conidiogenous cell giving rise to a single conidium, or with septa and each cell forming a single conidium. Primary conidia mostly forcibly discharged, with outer wall continuous from that of conidiogenous cells, and inner wall invaginating to form two-layer septa separating conidia and conidiogenous cells. Secondary conidia usually formed, forcibly discharged from short secondary conidiophores and usually similar in shape to primary conidia, or passively dispersed from long, thin secondary conidiophores and usually distinct from primary conidia. Resting spores with thick and 2-layer walls, colored or hyaline, smooth or decorated. Zygospores or azygospores laterally or in the axis of the parental cells. Nuclear numbers more than two in mature spores. Mature spores forming germ conidiophores and germ conidia (secondary-conidia-like) or forming germ mycelia and germ conidia (primary-conidia-like).

**Life modes and distribution:** Saprobes in soil or litter, primary pathogens of arthropods (insect, mites, spiders) or other soil invertebrates (nematodes, tardigrades), or highly specifc pathogens of desmid algae, fern gametophytes. Arthropod pathogens may form specialized organs: rhizoids with or without diferentiated holdfasts may anchor hosts to substrates, and cystidia may perforate host cuticle and facilitate the emergence of conidiophores.

**Notes:** Hibbett et al. ([2007](#page-139-1)) recognized *Entomophthoromycotina* Humber as a distinct subphylum along with three other subphyla *Mucoromycotina*, *Kickxellomycotina*, and *Zoopagomycotina*. Humber ([2012\)](#page-140-3) elevated this subphylum to a phylum rank as *Entomophthoromycota*, including three classes, viz*., Basidiobolomycetes* Humber, *Neozygitomycetes* and *Entomophthoromycetes.* The placement of *Basidiobolomycetes* in *Entomophthoromycota* is not accepted in subsequent studies (e.g. Sekimoto et al. [2011](#page-154-4); Gryganskyi et al. [2013\)](#page-138-4), but placed in the phylum *Basidiobolomycota* instead (Tedersoo et al. [2018](#page-157-4)). And thus, *Entomophthoromycota* currently comprises two classes *Neozygitomycetes* and *Entomophthoromycetes* (Gryganskyi et al. [2013](#page-138-4); Tedersoo et al. [2018](#page-157-4); Wijayawardene et al. [2018b,](#page-160-6) [2020a](#page-160-0); Voigt et al. [2021](#page-158-3))*.* Moreover, Tedersoo et al. ([2018](#page-157-4)) considered that *Entomophthoromycota* is closely related to *Kickxellomycota* and *Zoopagomycota*, and introduced the subkingdom *Zoopagomyceta* to accommodate these three phyla.

**Controversial opinions:** It is worth to note that Spatafora et al. [\(2016](#page-155-6)) reassigned the subphylum *Entomophthoromycotina* into the phylum *Zoopagomycota.* Galindo et al. [\(2021](#page-137-0)) and Strassert and Monaghan [\(2022a](#page-156-3), [b](#page-156-4)) agreed on this placement. Therefore, the high-level taxonomic status of this fungal group needs more evidence to be clarifed.

*Entomophthoromycetes* Humber, Mycotaxon 120: 486 (2012).

Index Fungorum Registration Identifer 564381; Fig. [41](#page-104-0)

Vegetative growth as coenocytic mycelia or rod-like to variably shaped hyphal bodies, walled; if wall-less, amoeboid. Conidiophores simple or digitately branched and with a single apical conidiogenous cell, or unbranched, septate and forming one conidium per cell. Primary conidia unitunicate or bitunicate, variously shaped, uni- to multinucleate, with fat, conical or rounded basal papilla, forcibly discharged by papillar eversion in most genera. Secondary conidia more or less similar in shape to primary conidia and forcibly discharged if formed on short secondary conidiophore, or elongate and passively dispersed if formed on elongated capillary secondary conidiophore. Nuclei with a small nucleolus, with interphasic heterochromatin present in *Entomophthoraceae* only, intranuclear mitosis, with a small lateral metaphase plate, with interphasic chromosomes partly condensed and stained readily in *Entomophthoraceae* but euchromatic in other families. Resting spores globose to subglobose, formed as zygospores or azygospores.

**Life mode and distribution:** Obligately pathogenic for invertebrates (Entomophthoraceae, Meristacraceae, some Ancylistaceae), saprobic (some Ancylistaceae), or phytopathogenic (Completoriaceae and Ancylistaceae).

**Notes:** The class *Entomophthoromycetes* was introduced by Humber [\(2012\)](#page-140-3) to accommodate the order *Entomophthorales* (*Ancylistaceae*, *Completoriaceae*, *Entomophthoraceae* and *Meristacraceae*), which was supported by Gryganskyi et al. [\(2013](#page-138-4)). Recently, Nie et al. [\(2022](#page-148-5)) introduced two new genera in *Entomophthorales*, i.e. *Capillidium*, typifed by *Capillidium heterosporum* (Drechsler) B. Huang & Y. Nie and composed of other six new combinations, and *Neoconidiobolus*, typifed by *Neoconidiobolus thromboides* (Drechsler) B. Huang & Y. Nie. Subsequently, *Capillidiaceae*, *Conidiobolaceae* and *Neoconidiobolaceae,* three other families of *Entomophthorales*, was established by Gryganskyi et al. ([2022](#page-138-5)).

**Type order:** *Entomophthorales* G. Winter [as 'Entomophthoreae'], Rabenh. Krypt.-Fl., Edn 2 (Leipzig) 1.1: 74 (1880) [1884].

**Type family:** *Entomophthoraceae* A.B. Frank, Synopsis der Drei Naturreiche. Ein Handbuch …, Edn 2: 1258 (1874).

**Type genus:** *Entomophthora* Fresen., Bot. Ztg. 14: 883 (1856).

**Type species:** *Entomophthora muscae* (Cohn) Fresen., Bot. Ztg. 14: 883 (1856).

*Neozygitomycetes* Humber, Mycotaxon 120: 485 (2012). Index Fungorum Registration Identifer 564377.

Vegetative hyphal bodies rod-like, walled or protoplastic, elongating until  $\pm$  synchronous mitosis, with daughter cells separated by septa. Nuclear number in all cell types strongly regulated, usually 3–5 in vegetative cells and conidia and 2 in resting spores. Mitoses intranuclear,  $\pm$  synchronous in any cell. Nuclei fusoid at metaphase with central, fusoid spindle, without nucleus-associated mitotic organelles. Chromosomes euchromatic during interphase. Conidiophores simple, forming apical conidiogenous cell and one conidium. Primary conidia subglobose to broadly ovoid, with a short basal papilla, comparatively fat; forcibly discharged to short distance by papillar eversion. Secondary conidia usually forming quickly after primary conidial discharge, most commonly forming as capilliconidia. Resting spores buding from a short conjugation bridge between rounded-up gametangia.

**Life modes and distribution:** Pathogens of either aphids or mites.

**Notes:** Humber [\(2012](#page-140-3)) recognized the family *Neozygitaceae* as a distinct lineage in '*Zygomycota*' and thus introduced the order *Neozygitales* and the Class *Neozygitomycetes* in the newly introduced Phylum *Entomophthoromycota*.

**Type order:** *Neozygitales* Humber, Mycotaxon: 486 (2012). **Type family:** *Neozygitaceae* Ben Ze'ev, R.G. Kenneth & Uziel, Mycotaxon 28(2): 321 (1987).

**Type genus:** *Neozygites* Witlaczil, Arch. Mikr. Anat. 24: 599 (1885).

**Type species:** *Neozygites fresenii* (Nowak.) Remaud. & S. Keller, Mycotaxon 11(1): 332 (1980).

=*Neozygites aphidis* Witlaczil, Arch. Mikr. Anat. 24: 599 (1885).

*ENTORRHIZOMYCOTA* R. Bauer, Garnica, Oberw., Riess, Weiß & Begerow, PLoS ONE 10(7): e0128183, 10 (2015).

Index Fungorum Registration Identifer 808783; Fig. [42](#page-105-0)

Hyphae frst grew intercellularly between cortical cells, and develop to intracellular hyphae inside host tissue, forming intracellular septate coils bearing terminal teliospores. Teliospores germinate internally, becoming four-celled; hyphae with regular septa, with or rarely without dolipores, without Woronin bodies or membrane caps. Clamp connections absent. Haustoria present (modifed from Bauer et al. [2015](#page-131-12); Tedersoo et al. [2018](#page-157-4)).

**Life modes and distribution:** Phytoparasitic by forming root galls in Cyperaceae or Juncaceae or rarely in Eudicodyledons.

**Notes:** Over the last two decades molecular phylogenies have gradually elevated the *Entorrhiza* to a new phylum. Bauer et al. ([1997,](#page-131-5) [2001](#page-131-7)) initially regarded *Entorrhiza* as a basal lineage inside *Ustilaginomycotina* on the basis of ultrastructure and morphology, and the frst molecular studies to include rDNA of *Entorrhiza* have focused exclusively on the members of *Ustilaginomycotina*, recovering the basal position of *Entorrhiza* to the rest of the group (Begerow et al. [2006](#page-131-6)). Further analyses with rDNAs using more inclusive samplings of fungal species inferred an even deeper placement of *Entorrhiza*, concluding that *Entorrhiza* should be viewed either as a sister group of the *Basidiomycota* or an *incertae sedis* among *Basidiomycota* (Matheny et al. [2006](#page-146-4); Hibbett et al. [2007\)](#page-139-1).

A five-gene analysis by Bauer et al.  $(2015)$  $(2015)$ , which employed a concatenated dataset of rDNAs and partial RNA polymerase II subunit sequences, has affirmed that *Entorrhiza* represents a distinct early lineage of higher fungi, warranting an introduction of a new fungal phylum *Entorrhizomycota*. The fve-gene analysis reconstructed *Entorrhiza* as a sister group to the rest of *Dikarya*, although the alternative topology, with *Entorrhiza* sister to *Basidiomycota*, could not be rejected by a statistical signifcance test (Bauer et al. [2015\)](#page-131-12). To this day, the exact placement of *Entorrhizomycota* within *Dikarya* remains uncertain. Transcriptomic or genomic data for this phylum are lacking, and the available data are limited to the rDNA sequences and the partial RPB genes from Bauer et al. [\(2015](#page-131-12)).

*Entorrhizomycetes* Begerow, M. Stoll & R. Bauer, Mycologia 98(6): 908 (2007) [2006].

Index Fungorum Registration Identifer 501482.

Description is similar to the phylum description.

**Notes:** Bauer et al. [\(2015](#page-131-12)) introduced *Entorrhizomycetes* as the type class of *Entorrhizomycota*. Currently, *Entorrhizomycetes* comprises two orders *Entorrhizales* R. Bauer & Oberw. and *Talbotiomycetales* K. Riess et al.

**Type order:** *Entorrhizales* R. Bauer & Oberw., in Bauer, Oberwinkler & Vánky, Can. J. Bot. 75: 1311 (1997).

**Type family:** *Entorrhizaceae* R. Bauer & Oberw., in Bauer, Oberwinkler & Vánky, Can. J. Bot. 75(8): 1311 (1997).

**Type genus:** *Entorrhiza* C.A. Weber, Bot. Ztg. 42: 378 (1884).

**Type species:** *Entorrhiza cypericola* (Magnus) C.A. Weber, Bot. Ztg. 42: 378 (1884).

*GLOMEROMYCOTA* C. Walker & A. Schüßler, in Schüßler, Schwarzott & Walker, Mycol. Res. 105(12): 1416 (2001).

Index Fungorum Registration Identifer 90511; Fig. [43](#page-106-0)

The core of the symbiotic trade-off is represented by arbuscules, specifc structures produced by AMF in plant root cells, where the fungi supply minerals and water in exchange for 20% of the carbon fxed by the plants. **Sexual morph:** Unknown. **Asexual morph:** acaulosporoid, ambisporoid, entrophosporoid, entrosacculoid, gigasporoid, glomoid, otosporoid or scutellosporoid glomerospores. Glomerospores are multinucleate, globose, subglobose, ellipsoid, ovoid or irregular, hyaline or pigmented, ranging from 30 to 1.050 µm, consisting of one to four walls with one to multiple layers that may or may not stain in Melzer's reagent (Fig. [44\)](#page-107-0). *Glomeromycota* produce extraradical mycelium, intraradical hyphae and arbuscules. Intraradical vesicles a storage and propagative structure—are also formed by members of all orders except *Gigasporales*. Instead, *Gigasporales* form extraradical auxiliary cells, which also function as storage (de La Providencia et al. [2005;](#page-135-11) de Souza et al. [2005](#page-135-12)), but not as propagative structures. Species of *Pacispora* (*Diversisporales*) form both intraradical vesicles and extraradical auxiliary cells. The intraradical mycelium stains strongly or weakly with trypan blue. The mycelium is coenocytic and hyphal anastomosis, which play a key role in genetic exchange among compatible individuals and in the formation of interconnected mycorrhizal networks, have been observed in a limited number of isolates. Extraradical glomerospores are single, in epigeous or sub-hypogeous loose or compact clusters or glomerocarps.

**Life modes and distribution:** Arbuscular Mycorrhizal Fungi (AMF) are obligate biotrophic symbionts, forming arbuscular mycorrhiza with ca. 80% of terrestrial and an unknown number of aquatic plants, and one species forming symbiosis with cyanobacteria of the genus *Nostoc* (Fig. [43](#page-106-0)). The phylum is worldwide distributed, with occurrences in all continents.

**Notes:** The class *Glomeromycetes* (= *Glomomycetes*) was described by Cavalier-Smith ([1998](#page-133-2)) in the phylum '*Zygomycota*' with two orders (*Glomerales* and *Endogonales*) as sister clades. Schüßler et al. ([2001](#page-154-5)) erected the phylum *Glomeromycota*, raising *Glomeromycetes* to the rank of phylum. Oehl et al. ([2011a\)](#page-149-5), using spore morphology and molecular analyses of the ribosomal gene sequences, concluded that *Glomeromycota* constitutes a phylum formed by three classes (*Archaeosporomycetes*, *Glomeromycetes*, *Paraglomeromycetes*).

Currently, six orders, 17 families, 49 genera, and approximately 360 species are recognized in *Glomeromycota* (Błaszkowski et al. [2022;](#page-132-7) Silva et al. [2022b;](#page-155-7) Wijayawardene et al. [2022a](#page-160-1); Chen et al. [2023a,](#page-133-3) [b;](#page-133-4) Corazon-Guivin et al. [2023](#page-134-3); da Silva et al. [2023\)](#page-134-4). Previous phylogenomic analyses (Galindo et al. [2021;](#page-137-0) Mikhailov et al. [2022;](#page-147-4) Strassert and Monaghan [2022a,](#page-156-3) [b](#page-156-4)) did not solve the ambiguities about the phylogenetic position of the phylum. Despite the conficting results of the Bayesian inference and the disagreement between the ML and Bayesian trees, eventually the reconstructions obtained in the present work with the CAT-PMSF method placed the *Glomeromycota* phylum sister to *Dikarya* with nearly full support.

*Archaeosporomycetes* Sieverd., G.A. Silva, B.T. Goto & Oehl, Mycotaxon 116: 374 (2011).

Index Fungorum Registration Identifer 519686.

**Asexual morph:** acaulosporoid, entrophosporoid and/ or glomoid glomerospores. *Archaeosporomycetes* include dimorphic and, to a lesser extension, monomorphic species, producing acaulosporoid (produced laterally on the neck of a sporiferous saccule), entrophosporoid (produced intercalary the neck of a sporiferous saccule) and glomoid spores (formed at the tip of sporogenous hyphae, occasionally intercalary). Only one species, *Geosiphon pyriforme* (Kütz) Wettst, produces a distinct symbiotic association with cyanobacteria, or blue-green algae, of the genus *Nostoc* (Fig. [43](#page-106-0)), with no evidence, so far, of symbiosis with plants ([https://](https://geosiphon.jimdosite.com/) [geosiphon.jimdosite.com/](https://geosiphon.jimdosite.com/)). Spores are globose, subglobose, ellipsoid, ovoid or sometimes irregular, hyaline to white, yellow to brown. Spores present one (glomoid spores) or two spore walls (acaulosporoid and entrophosporoid spores), which contain several layers of diferent types, including evanescent, laminated, fexible, semi-fexible, permanent, semi-permanent, smooth or coriaceous layers. Acaulosporoid spores germinate through a germ tube emerging from a germinal shield (orb) formed on the upper surface of the inner wall. Some species present Melzer's reaction in the spore wall, absent in the inner wall. Representatives of this class form extraradical mycelium, intraradical hyphae, arbuscules and vesicles that stain pale or not at all in trypan blue. **Sexual morph:** Unknown.

**Life modes and distribution:** Obligate biotrophic symbionts, forming arbuscular mycorrhiza with terrestrial plant species and one species forming symbiosis with cyanobacteria of the genus *Nostoc* but not with plants. The members of *Archaeosporomycetes* show a worldwide distribution, occurring in countries with temperate, tropical, subtropical and arid climates. Its representatives have been found associated with plant roots in diverse habitats, including sand dunes, forests, grasslands, polluted places, gardens, and humid ecosystems (Schüßler and Wolf [2005](#page-154-6); Błaszkowski et al. [2021\)](#page-132-8). Fossil records for *Archaeosporomycetes* are limited to *Archaeosporites rhyniensis* (Harper et al. [2020](#page-139-5)) and *Mycokidstonia sphaerialoides* (Walker et al. [2021](#page-159-6)), both from Devonian (400 mya). The development and establishment of the mycorrhizal symbiosis follow the general pathway of *Glomeromycota*. The exception is represented by *Geosiphon* which forms specialized structures called bladders where cyanobacteria are living and transferring photoassimilates in exchange of nutrients from the fungus (Kluge [2002](#page-143-1)).

**Notes:** The class *Archaeosporomycetes* was introduced by Oehl et al. ([2011b](#page-149-6)), including the order *Archaeosporales* previously erected by Schüßler et al. ([2001\)](#page-154-5). Currently, *Archaeosporomycetes* contains one order (*Archaeosporales*), four families (*Ambisporaceae*, *Archaeosporaceae*, *Geosiphonaceae*, *Polonosporaceae*), four genera (*Ambispora*, *Archaeospora*, *Geosiphon*, *Polonospora*) and 20 species. Representatives of *Archaeosporomycetes* are known to form arbuscular mycorrhizal (AM) symbiosis with plant roots (*Ambisporaceae*, *Archaeosporaceae*, *Polonosporaceae*) or non-AM symbiosis with photosynthetic and nitrogenfxing cyanobacteria (*Geosiphonaceae*). The phylogenetic relationships between *Geosiphonaceae* and the other families in *Archaeosporomycetes* need more investigations to be clarifed (Oehl et al. [2011a](#page-149-5); Błaszkowski et al. [2021](#page-132-8)). Some species like *Ambispora fecundispora*, *A*. *nicolsonii*, *A. reticulata*, *Archaeospora myriocarpa* and *A*. *undulata* need a phylogenetic placement since molecular data are not available yet. Phylogenetic analysis based on environmental DNA (eDNA) metabarcoding indicated clades potentially representing new taxa in *Archaeosporomycetes* (Błaszkowski et al. [2021\)](#page-132-8). Genome and transcriptome assemblies are available for two *Ambispora* species, and for *Geosiphon* whose genome possesses all the characteristics of known genomes of AM fungi (Malar et al. [2021;](#page-146-7) Montoliu-Nerin et al. [2021](#page-148-6)).

**Economic and ecological signifcances, examples:** Even if most studies have been carried out with species of *Glomeromycetes*, it has been shown that some representatives of *Archaeosporomycetes* also have economic and ecological importance. *Archaeospora myriocarpa* had high effectiveness in cassava and cofee plants in greenhouse and feld trials (Sieverding [1990](#page-155-8)). *Archaeospora trappei* promoted herbivory tolerance in *Plantago lanceolata* (Bennett et al. [2009\)](#page-131-13) and *Geosiphon* can increase heavy metal tolerance (Schüßler and Wolf [2005\)](#page-154-6). Cabrales et al. [\(2019\)](#page-132-9) showed that AMF inocula, including those with *Ambispora leptoticha*, improved maize yield in acidic soils. Furthermore, in an experiment conducted in soybean cultivars, *A*. *leptoticha* improved drought stress tolerance and productivity when coinoculated with the nitrogen-fxing bacterium *Bradyrhizobium liaoningense* (Ashwin et al. [2023](#page-130-6)). *Ambispora appendicula*, when used as inoculum, increased tolerance against coffee leaf rust (Vallejos-Torres et al. [2023\)](#page-158-7) and conferred bioprotective efects to cassava plants against the nematodes *Meloidogyne* spp. (Séry et al. [2016\)](#page-154-7).

**Type order:** *Archaeosporales* C. Walker & A. Schüßler, in Schüßler, Schwarzott & Walker, Mycol. Res. 105(12): 1418 (2001).

**Type family:** *Archaeosporaceae* J.B. Morton & D. Redecker, Mycologia 93(1): 182 (2001).

**Type genus:** *Archaeospora* J.B. Morton & D. Redecker, Mycologia 93(1): 183 (2001).

**Type species:** *Archaeospora trappei* (R.N. Ames & Linderman) J.B. Morton & D. Redecker 2001.

*Glomeromycetes* Caval.-Sm., Biol. Rev. 73: 246 (1998), emend. Oehl, G.A. Silva, B.T. Goto & Sieverd, Mycotaxon 116: 365–379 (2011).

Index Fungorum Registration Identifer 90168. Type cited as '*Glomomycetes*'.

**Asexual morph:** acaulosporoid, entrophosporoid, entrosacculoid, gigasporoid, glomoid, otosporoid or scutellosporoid glomerospores. Acaulosporoid spores develop laterally on the neck of a sporiferous saccule, singly in soil or rarely in aggregates. Spores are globose, subglobose, more rarely ovoid or irregular, consisting of three walls: an outer wall with one to five layers (swl1–5), sometimes ornamented with projections, depressions, or double ornamentations; a middle bi-layered wall (mwl1–2); and an inner tri-layered wall (iwl1–3), generally with the outer surface ornamented with granular outgrowths (beaded layer). Entrophosporoid spores formed within the necks of sporiferous saccules, singly in soil or rarely in roots. Spores are globose to subglobose, consisting of an outer wall with two layers (swl1–2), reactive in Melzer's, continuous with the wall of the stalk and the saccule, and an inner three-layered wall (iwl1–3) with no Melzer's reaction. Entrosacculoid spores produced inside sporiferous saccule, singly in soil. Spores are globose to subglobose, consisting of three walls: an outer wall with one to three layers (swl1–3), a middle bi-layered wall (mwl1–2); and an inner bi-layered wall (iwl1–2), generally with the upper surface ornamented with granular outgrowths (beaded layer). Gigasporoid spores produced terminally on sporogenous (bulbous) cells that arise from sporogenous hyphae, featuring germ warts on the inner surface of the mono-walled spore wall. Spores are single in soils, large  $(usually > 200 \mu m)$ , globose to subglobose, sometimes ovoid, with two to three layers (swl1–3). Sporogenous cell composed of two layers continuous with the frst two layers of the spore wall. Glomoid spores produced terminally, subterminally or intercalary from subtending hyphae, either in soils or occasionally within roots, singly, in loose clusters or in compact epigeous and sub-hypogeous glomerocarps, with or without peridium, sometimes arising radially from a central plexus of hyphae. Spores are globose, subglobose, irregular or ellipsoid, consisting of one wall, with one to six layers (swl1–6). Subtending hyphae straight, cylindrical to funnel-shaped, infated or constricted at the spore base. Pore open or most frequently occluded by a conspicuous septum. Otosporoid spores, possessing two multiple-layered walls, formed in the proximity of a terminal or intercalary sporiferous saccule, laterally on its neck. Scutellosporoid spores produced on sporogenous (bulbous) cells, with two to four (multi-layered) spore walls. Germinal shields are hyaline, light yellow to (dark-brown) pigmented, mono- to multiplylobed, with germinal tube initiations positioned on compartments (lobes) of the germinal shields. Dimorphic species in the order *Entrophosporales* produce both entrophosporoid and glomoid types. Acaulosporoid, entrophosporoid, entrosacculoid, glomoid and otosporoid species form mycorrhizae with arbuscles, vesicles, and hyphae that stain blue to dark blue in trypan blue. Gigasporoid and scutellosporoid species produce extraradical auxiliary cells instead of intraradical vesicles. **Sexual morph:** Unknown.

**Life modes and distribution:** The members of the class are worldwide distributed, with occurrences in over 175 countries, encompassing all continents. However, analyzes of distribution patterns revealed that some families in *Glomeromycetes* tend to be concentrated in geographic regions infuenced by climate and soil (Davison et al. [2021](#page-134-5)). *Acaulosporaceae* and *Gigasporaceae*, for instance, occur frequently in tropical zones such as Africa and South America, while *Glomeraceae* is well represented in temperate zones such as Europe, Asia and Oceania (Stürmer et al. [2018](#page-156-6)). Furthermore, *Glomeromycetes* spp. are able to form mycorrhizal symbiosis with a wide range of terrestrial and aquatic hosts, and are largely tolerant to stress conditions, including deserts, saline environments, polluted areas, fooded regimes or even in hypoxic environments, such as eutrophic lakes (Ragupathy et al. [1990;](#page-151-6) Mehrotra [1998;](#page-147-5) Paul et al. [2007](#page-149-7); Baar et al. [2011](#page-130-7); Jobim and Goto [2016;](#page-141-5) El-Gazzar et al. [2020](#page-136-1); Queiroz et al. [2020,](#page-151-7) [2022;](#page-151-8) Chen et al. [2022;](#page-133-5) Gomes et al. [2022](#page-137-1); Malicka et al. [2022](#page-146-8)).

**Notes:** Cavalier-Smith [\(1998\)](#page-133-2) introduced the class *Glomomycetes* (later corrected to *Glomeromycetes* by Schüßler et al. [2001](#page-154-5)) in the *Zygomycota* phylum with *Glomerales* and *Endogonales* as sister clades. Schüßler et al. ([2001\)](#page-154-5) described the phylum *Glomeromycota* and transferred *Glomeromycetes* to the new phylum reorganized in four orders: *Glomerales*, and the newly described *Archaeosporales*, *Diversisporales*, and *Paraglomerales*. Oehl et al. ([2008\)](#page-149-8) introduced the new order *Gigasporales* in *Glomeromycetes*, and more recently Błaszkowski et al. ([2022\)](#page-132-7) erected the order *Entrophosporales* to accommodate a single family, *Entrophosporaceae*, previously classifed in *Glomerales* as *Claroideoglomeraceae* (Schüßler and Walker [2010\)](#page-153-2). Currently, four orders, eleven families, 41 genera, and approximately 300 species are recognized in *Glomeromycetes* (Silva et al. [2022b](#page-155-7); Wijayawardene et al. [2022a\)](#page-160-1), making it the most species-rich and the most diverse class in morphological, molecular, and ecological aspects within *Glomeromycota*. Genome assemblies are available for the genera *Acaulospora*, *Cetraspora*, *Dentiscutata*, *Diversispora*, *Entrophospora*, *Funneliformis*, *Fuscutata*, *Gigaspora*, *Glomus*, *Oehlia*, *Racocetra*, *Rhizoglomus* and *Scutellospora* (Kobayashi et al. [2018](#page-143-2); Sun et al. [2019a](#page-156-7), [b](#page-156-8); Montoliu-Nerin et al. [2021](#page-148-6); Sahraei et al. [2022\)](#page-152-2). Transcriptomes are available for *Acaulospora*, *Diversispora*, *Funneliformis*, *Gigaspora*, *Racocetra*, *Rhizoglomus* and *Scutellospora* (Beaudet et al. [2018](#page-131-14)).

The oldest fossil records for *Glomeromycetes* are hyphae and spores (glomoid) from Wisconsin, dated to the Ordovician period (460 mya) (Redecker et al. [2000](#page-151-9)). However, the most impressive records were obtained from the Lower Devonian (400 mya). The fossils shared morphological features similar to existent taxa, for instance, *Scutellosporites devonicus* (Dotzler et al. [2006\)](#page-135-13) presenting scutellosporoid spores, or *Glomites rhyniensis* (Taylor et al. [1995](#page-156-9)) and *G*. *sporocarpoides* (Karatygin et al. [2006](#page-142-5)), both with glomoid spores.

**Economic and ecological signifcances, examples:** *Glomeromycetes* enhance plant performance by increasing water and nutrient absorption, photosynthetic rate, growth, and tolerance to biotic and abiotic stress, e.g., drought, salinity, pathogens and heavy metals. In addition, this symbiosis promotes soil biogeochemical cycling and enhances aggregate stability (Latef et al. [2016](#page-144-11); Singh et al. [2020;](#page-155-9) Falcão and Silva [2023](#page-136-2)).

The majority of model species used in biochemical, genetic and ecological studies of AM symbiosis are classifed within the class *Glomeromycetes*, belonging mostly to the genera *Rhizoglomus* (with *R. irregulare* as the most studied species)*, Gigaspora* and *Funneliformis.* Notably, *Funneliformis mosseae*, *Rhizoglomus clarum*, *R. irregulare* and *R. intraradices* are the most used components of commercial inoculants (Basiru and Hijri [2022](#page-131-15)), and have shown positive responses in inoculations of several agricultural crops, e.g., banana, bean, cassava, lettuce, maize, olive, potato, rice, soybean, tomato and walnut (Sahodaran et al. [2019](#page-152-3); Santana et al. [2019;](#page-153-3) Singh et al. [2019](#page-155-10); Campo et al. [2020;](#page-133-6) Chenchouni et al. [2020;](#page-133-7) Deja-Sikora et al. [2020](#page-135-14); Huang et al. [2020](#page-140-4); Moreira et al. [2020](#page-148-7); Musyoka et al. [2020](#page-148-8); Cavallari et al. [2021;](#page-133-8) Santander et al. [2021;](#page-153-4) Jumrani et al. [2022](#page-142-6)). Furthermore, *Glomeromycetes* spp. are also benefcial to non-agricultural plants, including medicinal and ornamental plants, and those representing forest plant communities (Ma et al. [2019](#page-146-9); Zhang et al. [2019](#page-161-5); Mirzaie et al. [2020](#page-147-6); Klinsukon et al. [2021](#page-143-3); Medeiros et al. [2021](#page-147-7)). *Glomeromycetes* have been detected in decomposing leaf litter (Bunn et al. [2019](#page-132-10); Lima et al. [2023](#page-145-4)), which is a potential substrate for the recovery of impacted environments (Medeiros et al. [2021\)](#page-147-7). Therefore, they can be successfully used by inoculum producers for crop production, sustainable agriculture and ecological restoration (Basiru and Hijri [2022](#page-131-15); Duell et al. [2022](#page-136-3)).

**Type order:** *Glomerales* J.B. Morton & Benny [as 'Glomales'], Mycotaxon 37: 473 (1990).

**Type family:** *Glomeraceae* Piroz. & Dalpé [as 'Glomaceae'], Symbiosis 7: 19 (1989).

**Type genus:** *Glomus* Tul. & C. Tul., G. bot. ital. 2(1): 63 (1844).

**Type species:** *Glomus macrocarpum* Tul. & C. Tul., Giornale Botanico Italiano 1 (2): 63 (1845).

*Paraglomeromycetes* Oehl, G.A. Silva, B.T. Goto & Sieverd., Mycotaxon 116: 374 (2011).

Index Fungorum Registration Identifer 519687.

Obligate biotrophic symbionts, forming arbuscular mycorrhiza with the majority of terrestrial and aquatic plant species. **Sexual morph**: Unknown. **Asexual morph**: glomoid glomerospores. Spores globose to subglobose, sometimes ovoid or irregular, hyaline or pigmented, small (50–130 µm), formed singly or in loose clusters, in soil or inside roots. Spore structure consists of one wall with two to three layers, or two walls with many layers (e.g., *Paraglomus* spp.). Spore wall layer 1 is usually evanescent, short-lived and completely sloughed off in mature spores. Layer 2 is generally laminate, while layer 3, when present, is fexible, hyaline. Some *Paraglomus* spp. might have ornamentations on the spore wall. Inner spore wall consisting of two to three hyaline and fexible layers. Layers of the outer or inner walls may not exhibit any reaction in Melzer's reagent, or turn orange, yellow to pinkish-yellow. Subtending hyphae are straight or curved, cylindrical to slightly funnel-shaped, or rarely fared or infated, with a pore open or occluded by a straight or curved septum. The wall of the subtending hypha is continuous with the spore wall layers. Germinal shield or orb unknown. Spores germinate directly through the spore

wall. Mycorrhizal structures stain faintly or not at all in trypan blue. Sporocarps unknown.

**Life modes and distribution:** Members of *Paraglomeromycetes* are generally ubiquitous in the world as they were found in alpine, temperate, mediterranean to tropical, subtropical, arid and humid areas (Mello et al. [2013](#page-147-8); Błaszkowski et al. [2017](#page-132-11)). Some studies highlighted the occurrence of *Paraglomeromycetes* in highly polluted areas (Dranguet et al. [2017](#page-136-4); Malicka et al. [2022\)](#page-146-8). Notably, *Paraglomus occultum* is widely distributed (Mello et al. [2013\)](#page-147-8), as well as *P. turpe* which has been detected in several agricultural soils of Europe (Oehl et al. [2016\)](#page-149-9). On the other hand, *Pervetustus simplex* seems to have a restricted distribution, as it has only been found in eight countries. Analogously, *P. bolivianum* has been detected in a prairie, Santa Cruz, Bolivia; in *Caatinga* region, Pernambuco, Brazil (Oehl and Sieverding [2004](#page-149-10); Mello et al. [2013;](#page-147-8) Lima et al. [2022](#page-145-5)) and most recently in southeast Mexico, Oaxaca in the rhizosphere of *Agave angustifolia* and *A. karwinskii* (Sanchez et al. [2018](#page-152-4); Polo-Marcial et al. [2021](#page-150-9)).

**Notes:** Morton and Redecker [\(2001\)](#page-148-9) erected the family *Paraglomeraceae* in the order *Glomales* (=*Glomerales*), *Zygomycota* (Morton and Benny [1990](#page-148-10)) with a single genus, *Paraglomus,* and two species, *P. brasilianum* and *P. occultum*. Originally, the two species were described as *Glomus* spp., relying mainly on nrDNA phylogenies, fatty acid profles, immunological reactions, and faint coloration of the intraradical hyphae in trypan blue. Schüßler et al. ([2001\)](#page-154-5), based on the high divergence of sequences, transferred *Paraglomeraceae* into the new order *Paraglomerales*, *Glomeromycota.* Oehl et al. ([2011a](#page-149-5)), using spore morphology and phylogenies reconstructed from ribosomal sequences, introduced the class *Paraglomeromycetes* to accommodate this order.

The species *Paraglomus albidum*, *P. laccatum* and *P. lacteum* were originally described with a single spore wall, difering from other *Paraglomus* spp. with two walls. Mello et al. [\(2013\)](#page-147-8) suggested that only *P. majewskii* formed a single spore wall, and this species was later transferred to the new genus *Innospora* in *Paraglomeraceae* according to molecular evidence (Błaszkowski et al. [2017\)](#page-132-11). However, since glomoid spores with one wall is a common feature for many taxa in *Glomeromycota*, Błaszkowski et al. [\(2017](#page-132-11)) concluded that *Innospora* does not present any exclusive morphological trait that distinguishes this genus from other glomoid genera. In the same paper, Błaszkowski et al. [\(2017](#page-132-11)) presented a new species with nrDNA sequences highly divergent from those of other members in *Paraglomeraceae*. Consequently, Błaszkowski et al. [\(2017\)](#page-132-11) erected a new family, *Pervetustaceae*, with the new genus and species *Pervetustus* and *P. simplex*, respectively.

Currently, *Paraglomeromycetes* contains one order (*Paraglomerales*), two families (*Paraglomeraceae*, *Pervetustaceae*), three genera (*Paraglomus*, *Innospora*, *Pervetustus*), and twelve species (Mello et al. [2013;](#page-147-8) Błaszkowski et al. [2017;](#page-132-11) Corazon-Guivin et al. [2020\)](#page-134-6). Up to now, the molecular phylogeny is known only for nine species. Environmental sequences deposited in public databases (Mello et al. [2013](#page-147-8); Błaszkowski et al. [2017](#page-132-11)) suggest the existence of many potentially undescribed species, despite the low number of species currently described in *Paraglomeromycetes*, compared to the other classes.

Genome and transcriptome assemblies are available only for two *Paraglomus* species (Beaudet et al. [2018;](#page-131-14) Malar et al. [2021](#page-146-7); Montoliu-Nerin et al. [2021](#page-148-6)).

**Economic and ecological signifcances, examples:** Few studies involving *P. occultum*, showed benefts for agricultural crops under stress conditions. For instance, in orange tree plants under water stress conditions, *P. occultum* promoted growth and tolerance by acting on sucrose metabolism and leaf osmotic regulation (Wu et al. [2017\)](#page-160-7). In tomato plants under salt stress, it increased the expression of stressresponsive genes and the nitrogen balance index (Liang et al. [2022](#page-145-6)). In orange groves, through the glomalin-related soil protein fractions, it contributed to the soil's total nitrogen content (Meng et al. [2020](#page-147-9)). Zhang et al. ([2018a,](#page-161-6) [b\)](#page-161-7) showed that *P. occultum* established common mycelium networks among orange seedlings and, thereby, increased plant growth and proftably modifed the root architecture of the recipient plants. Furthermore, *P. occultum* increased the expression of defense genes against pathogens in plants with citrus canker (Xie et al. [2019\)](#page-160-8).

**Type order:** *Paraglomerales* C. Walker & A. Schüßler, in Schüßler, Schwarzott & Walker, Mycol. Res. 105(12): 1418 (2001).

**Type family:** *Paraglomeraceae* J.B. Morton & D. Redecker [as 'Paraglomaceae'], Mycologia 93(1): 188 (2001).

**Type genus:** *Paraglomus* J.B. Morton & D. Redecker, Mycologia 93(1): 188 (2001).

**Type species:** *Paraglomus occultum* (C. Walker) J.B. Morton & D. Redecker (2001).

*KICKXELLOMYCOTA* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 150 (2018).

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*Thallus* arising from a holdfast on other fungi as a haustorial parasite, or branched, septate, subaerial hyphae; *mycelium* branched or unbranched, regularly septate; septa with median, disciform cavities containing plugs; asexual production by 1- or 2-spored merosporangia, trichospores, or arthrospores; sexual reproduction by zygospores that are globose, biconical, or allantoid and coiled (adopted from Hibbett et al. [2007](#page-139-1) which was placed to subphylum *Kickxellomycotina*).

**Life modes and distribution:** Saprobes or weak non-haustorial mycoparasites in soil and dung; widespread.

**Notes:** Hibbett et al. [\(2007](#page-139-1)) introduced *Kickxellomycotina* to accommodate *Kickxellales* Kreisel ex R. K. Benj.. In the last two decades, several studies have added a considerable number of taxa based on both morphology and phylogeny (e.g. Kurihara et al. [2008](#page-143-4); Chuang et al. [2017](#page-133-9); Ri et al. [2022\)](#page-151-10). The placement of *Kickxellales* in '*Zygomycota*' was confrmed in diferent studies (e.g. Tanabe et al. [2000](#page-156-10); White et al. [2006](#page-159-7)). Tretter et al. (2014) provided a comprehensive background on molecular phylogenetic placements of the taxa of *Kickxellomycotina* (based on SSU, LSU, ITS, MCM7, TSR1, *rpb1*, *rpb2*, *β-tubulin*) and recognized four orders (i.e. *Asellariales*, *Dimargaritales*, *Harpellales* and *Kickxellales*) and four orphan genera (*Barbatospora*, *Orphella*, *Ramicandelaber*, *Spiromyces*). However, the above mentioned studies did not recognize any class in *Kickxellomycotina* but only orders. Tedersoo et al. ([2018\)](#page-157-4) elevated the sub-phylum *Kickxellomycotina* to phylum rank and introduced *Kickxellomycota*. At the same time, Tedersoo et al. [\(2018](#page-157-4)) introduced six new classes, viz., *Asellariomycetes*, *Barbatosporomycetes*, *Dimargaritomycetes*, *Harpellomycetes*, *Kickxellomycetes* (type class) and *Ramicandelaberomycetes*. We believe this approach in Tedersoo et al. ([2018](#page-157-4)) provided more stable condition for the Kingdom of *Fungi*.

**Controversial opinions:** Strassert and Monaghan ([2022a,](#page-156-3) [b](#page-156-4)) regarded *Kickxellomycota* as a subphylum of *Zoopagomycota*, i.e. *Kickxellomycotina*.

*Asellariomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 151 (2018); Fig. [45](#page-108-0)

*Thalli* Branched, flamentous, with harpelloid septa; basal cells for attachment to gut cuticle of a host; sholdfasts elaborate, plugged septum, generative cells disarticulate and function as propagules, do not produce external and deciduous trichospores; no dictyosomes; no zygospore; sexual reproduction absent or extremely uncommon (Benjamin [1979](#page-131-16); Lichtwardt and Lichtwardt [1986\)](#page-145-7).

**Life modes and distribution:** saprobes, have adapted to semi-terrestrial insects (*Collembola*) and isopods dwelling in freshwater, marine, and terrestrial environments.

**Notes:** Manier and Lichtwardt ([1969](#page-146-10)) introduced *Asellariaceae* to accommodate *Asellaria.* Subsequently, Lichtwardt and Manier ([1978](#page-145-8)) upgraded *Asellariaceae* to an ordinal level and introduced *Asellariales.* Tedersoo et al. [\(2018\)](#page-157-4) showed that *Asellariales* has a distinct phylogenetic lineage in the Phylum *Kickxellomycota. Asellariaceae* is the only family in the order *Asellariales*.

**Type order:** *Asellariales* Manier ex Manier & Lichtw., in Lichtwardt & Manier, Mycotaxon 7(3): 442 (1978).

**Type family:** *Asellariaceae* Manier ex Manier & Lichtw., Annls Sci. Nat., Bot. Biol. Vég., sér. 12 9: 526 (1968).

**Type genus:** *Asellaria* R.A. Poiss., Bull. biol. Fr. Belg. 66: 232 (1932).

**Type species:** *Asellaria caulleryi* R.A. Poiss., Bull. biol. Fr. Belg. 66: 233 (1932).

*Barbatosporomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 151 (2018).

Index Fungorum Registration Identifer 554012; Fig. [46](#page-109-0)

*Thallus* branched with a basal cell, bearing trichospores; trichospores cylindrical, without a collar, with multiple fne basal appendages, may bear a cylindrical sleeve or wall at the terminal end, which on dehiscence may reveal appendage-like filaments; zygospores unknown (adopted from Doweld [2014c\)](#page-135-15).

**Notes:** Tretter et al. [\(2013](#page-157-7)) confrmed the phylogenetic placement of *Barbatospora* in *Kickxellomycota.* Doweld ([2014c](#page-135-15)) introduced *Barbatosporaceae* and *Barbatosporales* which currently comprise only one genus, i.e. *Barbatospora.* The genus is also monotypic thus the family is represented by the type species of *Barbatospora.* Tedersoo et al. ([2018\)](#page-157-4) introduced *Barbatosporomycetes* in the phylum *Kickxellomycota.*

**Type order:** *Barbatosporales* Doweld, Index Fungorum 87: 1 (2014).

**Type family:** *Barbatosporaceae* Doweld, Index Fungorum 87: 1 (2014).

**Type genus:** *Barbatospora* M.M. White, Siri & Lichtw., Mycologia 98(2): 335 (2006).

**Type species:** *Barbatospora ambicaudata* M.M. White, Siri & Lichtw., Mycologia 98(2): 335 (2006).

*Dimargaritomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 151 (2018).

Index Fungorum Registration Identifer 554013; Fig. [47](#page-110-0)

*Vegetative hyphae* delicate, hyaline, branched, regularly septate, giving rise to numerous small, slightly swollen appressoria producing restricted, more or less ramified, aseptate haustoria; sporangiophores white or light colored, erect, septate, simple or branched, forming twospored merosporangia directly on the surfaces of terminal

enlargements or on cellular branchlets produced laterally or terminally; sporangiospores hyaline, subglobose, ellipsoid, or bacilliform; spores at maturity remaining dry or immersed in liquid; septa of both the vegetative and fruiting hyphae characteristically with median disciform cavities containing solid, colorless, more or less biconvex plugs bearing small, usually globose enlargements at their poles; zygospores globoid, thick-walled, hyaline, formed from the fusion of similar (adopted from Benjamin [1959](#page-131-17)).

**Life modes and distribution:** Parasites or facultative parasites of other *Mucorales.*

**Notes:** Benjamin ([1959\)](#page-131-17) introduced the family *Dimargaritaceae* to accommodate four genera viz*., Dimargaris*, *Dispira*, *Spinalia* and *Tieghemiomyces*. Later, Benjamin ([1979\)](#page-131-16) upgraded *Dimargaritaceae* to ordinal level and introduced *Dimargaritales.* Tanabe et al. ([2000](#page-156-10)) confrmed the placement of *Dimargaritales* in '*Zygomycota*' using SSU rDNA sequences. Later, James et al. ([2006a\)](#page-141-2) included *Dimargaris bacillispora* in their phylogenetic study of the kingdom *Fungi* and confrmed its placement in '*Zygomycota*'. White et al. ([2006\)](#page-159-7) showed that *Dimargaritales* clustered with *Neozygites* and accommodated in '*Zygomycota*'. However, Tretter et al. [\(2013\)](#page-157-7) provided the most comprehensive approach for *Dimargaritales* and confrmed the placement in subphylum *Kickxellomycotina*. Tedersoo et al. ([2018\)](#page-157-4) elevated *Kickxellomycotina* to a phylum while upgrading *Dimargaritales* to a new class, *Dimargaritomycetes.*

**Type order:** *Dimargaritales* R.K. Benj., in Kendrick, The Whole Fungus 2: 607 (1979).

**Type family:** *Dimargaritaceae* R.K. Benj., Aliso 4(2): 364 (1959).

**Type genus:** *Dimargaris* Tiegh., Annls Sci. Nat., Bot., sér. 6 1: 154 (1875).

**Type species:** *Dimargaris cristalligena* Tiegh., Annls Sci. Nat., Bot., sér. 6 1: 154 (1875).

*Harpellomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 151 (2018).

Index Fungorum Registration Identifer 554014; Fig. [48](#page-111-0)

*Thallus* simple or branched, with basal cell attached to the host; hyphae septate; septa contain a lenticular cavity; sexual reproduction via conical or biconical zygospores; asexual reproduction via exogeneous, lateral, elongate monosporous trichospores; endosymbionts of mostly freshwater arthropods (Benjamin [1979\)](#page-131-16).

**Life modes and distribution:** Reported as parasites of freshwater arthropods larval stages, widespread (Kirk et al. [2008](#page-142-0)).

**Notes:** The order *Harpellales* was introduced by Lichtwardt and Manier ([1978\)](#page-145-8) based on the family *Harpellaceae* L. Léger & Duboscq 1929 which was invalid (see Index Fungorum [2024](#page-141-6)). Hence, Cannon and Kirk [\(2007](#page-133-10)) validated *Harpellaceae* 1929 as *Harpellaceae* L. Léger & Duboscq ex P.M. Kirk & P.F. Cannon 2007.

According to Tretter et al. [\(2013](#page-157-7)), *Harpellales* is a wellestablished order in *Kickxellomycotina* and confrmed its placement close to *Asellariales*. Tedersoo et al. ([2018\)](#page-157-4) upgraded the order *Harpellales* to the class *Harpellomycetes*. Currently, the class comprises one order and two families.

**Type order:** *Harpellales* Lichtw. & Manier, Mycotaxon 7(3): 442 (1978).

**Type family:** *Harpellaceae* L. Léger & Duboscq ex P.M. Kirk & P.F. Cannon, Fungal Families of the World: 152 (2007).

**Type genus:** *Harpella* L. Léger & Duboscq, C. r. hebd. Séanc. Acad. Sci., Paris 188: 951 (1929).

**Type species:** *Harpella melusinae* L. Léger & Duboscq, C. r. hebd. Séanc. Acad. Sci., Paris 188: 951 (1929).

*Kickxellomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 150 (2018).

Index Fungorum Registration Identifer 554010; Fig. [49](#page-112-0)

*Thallus* branched, with septate hyphae giving rise to septate sporangiophores; septa with median disciform cavities containing colorless biconvex or biumbonate plugs that are persistent in 2–3% KOH; asexual reproduction by 1-spored sporangioles formed on pseudophialides that arise from globoid to elongate fertile branchlets termed sporocladia; sexual reproduction by nearly globose zygospores (adopted from Benjamin [1979](#page-131-16)).

**Life modes and distribution:** Saprobes, weak non-haustorial mycoparasites in soil and dung, excrement of cave crickets (Benjamin [1979](#page-131-16); Ri et al. [2022](#page-151-10)).

**Notes:** Tedersoo et al. [\(2018\)](#page-157-4) introduced *Kickxellomycetes* to accommodate the order *Kickxellales* which was widely accepted in previous studies (e.g. White et al. [2006;](#page-159-7) Hibbett et al. [2007](#page-139-1)). Currently, the class comprises one order and one family (Wijayawardene et al. [2022a\)](#page-160-1).

**Type order:** *Kickxellales* Kreisel ex R.K. Benj., in Kendrick, The Whole Fungus 2: 610 (1979).

**Type family:** *Kickxellaceae* Linder, Farlowia 1(1): 56 (1943) [1943–1944].

**Type genus:** *Kickxella* Coem., Bull. Soc. R. Bot. Belg. 1: 156 (1862).

**Type species:** *Kickxella alabastrina* Coem., Bull. Soc. R. Bot. Belg. 1: 156 (1862).

*Ramicandelaberomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 151 (2018).

Index Fungorum Registration Identifer 554015; Fig. [50](#page-113-0)

*Mycelium* composed of colorless, branched, septate hyphae. *Stolons* hyaline, septate, producing rhizoids at the apex. Sporangiophores derived from the distal parts of stolons, septate, at frst prostrated, verticillately branched once or twice by the development of branches, erect in age, often disarticulated at third or fourth septa from rhizoids, often producing fertile terminal heads. Branches verticillately producing branchlets, becoming broad in age, cylindrical or ellipsoidal, irregularly producing shorter branchlets. *Sporocladia* at frst composed of branchlets on branches of sporangiophores, elongate, attenuate distally, often composed of broadened branches of sporangiophores in age. *Sporogenous cells* (pseudophialides) arising from sporocladia and fertile heads, at frst subspherical, in age subspherical or hemispherical, producing sporangiola. *Sporangiola* narrow fusiform, slightly curved, aseptate, hyaline. *Sporangial wall* adnate to the sporangiospore. Zygospores and chlamydospores not observed (extracted from Ogawa et al. 2001).

**Life modes and distribution:** Isolated from soil from Japan and Taiwan (Ogawa et al. 2001; Kurihara et al. [2004](#page-143-5); Chuang et al. [2013](#page-133-11)).

**Notes:** Doweld ([2014d](#page-135-16)) introduced *Ramicandelaberaceae* and *Ramicandelaberales* to accommodate the genus *Ramicandelaber*. The phylogenetic placement of *Ramicandelaber* in *Kickxellomycotina* was confrmed by Chuang et al. ([2013\)](#page-133-11) and Tretter et al. (2014). Tedersoo et al. ([2018\)](#page-157-4) recognized the phylogenetic lineage of *Ramicandelaber* in *Kickxellomycota* and upgraded *Ramicandelaberales* to a class, *Ramicandelaberomycetes*.

Currently, the class comprises one order and one family.

**Type order:** *Ramicandelaberales* Doweld, Index Fungorum 69: 1 (2014).

**Type family:** *Ramicandelaberaceae* Doweld, Index Fungorum 69: 1 (2014).

**Type genus:** *Ramicandelaber* Y. Ogawa, S. Hayashi, Degawa & Yaguchi, Mycoscience 42(2): 193 (2001).

**Type species:** *Ramicandelaber longisporus* Y. Ogawa, S. Hayashi, Degawa & Y. Yaguchi, Mycoscience 42(2): 193 (2001).

*MONOBLEPHAROMYCOTA* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Index Fungorum Registration Identifer 90752.

Fungi with mycelial, rhizomycelial or monocentric thallus; when mycelial, hyphal thallus with a foam appearance due the vacuolated cytoplasm. **Asexual reproduction** by zoospores or autospores; sexual reproduction, when present, oogamous by means of posteriorly unifagellate antherozoids borne in antheridia and nonfagellate female gametes borne in oogonia. The main ultrastructure features of the monoblephs' zoospores are the ribosomes aggregation around the nucleus, numerous lipid droplets predominantly posterior, and many spherical mitochondria. The ribosomes are loosely enclosed by several strands of endoplasmic reticulum, the microtubule root radiates anteriorly from a striated disk that partially encircles the kinetosome, and the rumposome is adjacent to the microbody in microbody-lipid globule complex (MLC) (Powell and Letcher [2014](#page-150-7); Karpov et al. [2017a](#page-142-7); Adl et al. [2019;](#page-129-0) Tedersoo et al. [2018\)](#page-157-4).

**Life modes and distribution:** Saprotrophic especially in plant material, such as submerged fruits and twigs, and algae, in aquatic ecosystems; sometimes in terrestrial ecosystems.

**Notes:** The type genus of *Monoblepharomycota*, *Monoblepharis* Cornu was a well-studied genus, even before molecular techniques were available (e.g. Sparrow [1933](#page-155-11); Marek [1984\)](#page-146-11). Doweld ([2001\)](#page-135-0) introduced the phylum *Monoblepharomycota* to accommodate *Monoblepharidomycetes* J. H. Schafn. and *Hyaloraphidiomycetes* Doweld, which are composed of eucarpic and holocarpic species in the orders *Monoblepharidales* and *Hyaloraphidiales*, respectively. James et al. ([2006a,](#page-141-2) [b](#page-141-1)) recognized the *Monoblepharidales* taxa are phylogenetically related to *Chytridiales*. Dee et al. ([2015\)](#page-135-17) consider *Monoblepharidomycetes s*.*l*. phylogeny have already been showed *Hyaloraphidium* as basal genus, and the mycelial genera as a monophlyletic sister group to the monocentric species, like as *Harpochytrium* and *Oedogoniomyces*.

Currently, *Monoblepharomycota* comprises one subphylum (i.e. *Monoblepharomycotina*) and two classes i.e. *Hyaloraphidiomycetes* and *Monoblepharidomycetes*.

*Hyaloraphidiomycetes* Doweld, Prosyllabus Tracheophytorum, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Index Fungorum Registration Identifer 585035.

Fungi with holocarpic thallus which consists of a cell usually elongate without rhizoidal system. *Autosporos* form within the mother cell and released by rupture of wall cell, the number of spores per cell is 4–8, irregular sometimes. The spores usually contain one nucleus, but up to four nuclei were observed in mature cells; mitochondria are elongated and sometimes even thread-like; large organelles present,



**Fig. 37** *Caulochytrium protostelioides*. **a** Sporangiocarps arising from clump of zoosporangia (×350). **b** Sporangiocarps on agar surface (×600). **c** Twelve zoospores that emerged from the three empty aerial sporangia (×1800). **d** Four zoospores from a single aerial spo-

sometimes associated with lipid globules, mitochondria and cisternae of the endoplasmic reticulum resembles the microbody-lipid globule complex (MLC) of *Chytridiomycetes*; centrioles, basal bodies and fagella were not observed. **Sexual reproduction** Unknown.

**Life modes and distribution:** Saprotrophic in freshwater ecosystems.

**Notes:** The class *Hyaloraphidiomycetes* (including the order *Hyaloraphidiales* and family *Hyaloraphidiaceae*) was established by Doweld ([2001\)](#page-135-0) to accommodate the genus *Hyaloraphidium*. The class comprises a single, monotypic genus (i.e. *H. curvatum*)*,* was previously considered a colorless green alga. Molecular studies with *H*. *curvatum* showed that it is a sister group of the monoblephs.

**Type order:** *Hyaloraphidiales* Doweld, Prosyllabus HyaloraphidialesTracheophytorum, Tentamen Systematics Plantarum Vascularium (Tracheophyta) (Moscou): LXXVII (2001).

**Type family:** *Hyaloraphidiaceae* Doweld, HyaloraphidiaceaeProsyllabus Tracheophytorum, Tentamen Systematics Plantarum Vascularium (Tracheophyta) (Moscou): LXXVII (2001).

**Type genus:** *Hyaloraphidium* Korshikov, Hyaloraphidium in Pascher & Korshikov, Arch. Protistenk. 74: 251 (1931). **Type species:** *Hyaloraphidium curvatum* Korshikov, in Pascher & Korshikov, Arch. Protistenk. 74: 351 (1931).

rangium (×2000). **e** Encystment of four zoospores from an aerial sporangium (×2000). **f** Small aerial sporangium with two zoospores emerging (×2000). **g** Empty aerial sporangium with single exit papilla (×2000)

*Monoblepharidomycetes* J.H. Schaffn, Ohio Naturalist 9(4): 449 (1909) emend. Karpov et al., Fungal Biology 121: 734 (2017).

Index Fungorum Registration Identifer 90244 (Figs. [51](#page-114-0) and [52](#page-115-0)).

*Thallus* mycelial, rhizomycelial, or monocentric, when mycelial with or without constrictions, pseudosepta and catenulations. Monocentric forms may be planktonic, epibiotic, with an extracellular basal foot or holdfast. **Asexual reproduction** by zoospores; sexual reproduction, when present, oogamous by means of posteriorly unifagellate antherozoids borne in antheridia and nonfagellate female gametes borne in oogonia (Karpov et al. [2017a](#page-142-7) modifed). The main ultrastructure features of the monoblephs' zoospores are the ribosomes aggregation around the nucleus, numerous lipid droplets predominantly posterior, and many spherical mitochondria. The ribosomes are lossely enclosed by several strands of endoplasmic reticulum, the microtubule root radiates anteriorly from a striated disk that partially encircles the kinetosome, and the rumposome is adjacent to the microbody in microbody-lipid globule complex (MLC) (Powell and Letcher [2014;](#page-150-7) Adl et al. [2019\)](#page-129-0).

**Life modes and distribution:** Saprobic on vegetable and animal debris, Oomycetes and in association with algae in freshwater ecosystems and organic material in tropical and subtropical soils.

**Notes:** *Monoblepharidomycetes* was established by Schaffner [\(1909](#page-153-5)) and emended by Karpov et al. [\(2017a](#page-142-7)), however they described the class with *Hyaloraphidium,* genus considered here into *Hyaloraphidiomycetes*. Currently,

<span id="page-101-0"></span>



*Monoblepharidomycetes* comprises the order *Monoblepharidales*, with the families *Gonapodyaceae*, *Harpochytriaceae*, *Monoblepharidaceae*, *Oedogoniomycetaceae* and *Telasphaerulaceae*, six saprotrophic genera (*Gonapodya*, *Harpochytrium*, *Monoblepharella*, *Monoblepharis*, *Oedogoniomyces* and *Telasphaerula*) and around 28 species (Powell and Letcher [2014](#page-150-7); Karpov et al. [2017a\)](#page-142-7).

**Type order:** *Monoblepharidales* Sparrow, Mycologia 34(1): 115 (1942).

**Type family:** *Monoblepharidaceae* Warm., Haandb. syst. Bot.,Edn 2 (Kjøbenhavn): 57 (1884).

**Type genus:** *Monoblepharis* Cornu, Bull. Soc. bot. Fr. 18: 59 (1871).

**Type species:** *Monoblepharis sphaerica* Cornu, Bull. Soc. bot. Fr. 18: 59 (1872) [1871].

*MORTIERELLOMYCOTA* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 152 (2018).

Index Fungorum Registration Identifer 554018.

*Thallus* with dichotomously branching, anastomosing hyphae, bearing stylospores; *Sporangiophores* initially coenocytic, irregularily septated when mature; asexual reproduction via sporangia and sporangioles; sporangia spherical, multi-spored; no columella; sporangioles terminal, borne on erecting hyphae; *Spores* ellipsoid or globose or irregular, smooth or ornamented; zygospores naked; mostly saprotrophs in soil (from Tedersoo et al. ([2018\)](#page-157-4) which was based on Hofmann et al. [\(2011](#page-140-5)) and Doweld ([2014f](#page-135-18))).

**Life modes and distribution:** Saprobic, plant-mutualistic or entomopathogenic, widespread.

**Notes:** The genus *Mortierella* Coem. and other morphologically resemble genera were well studied and higher ranks have been introduced based on their unique morphological characteristics (see Cavalier-Smith [1998](#page-133-2); Doweld [2014f](#page-135-18)). Hofmann et al. ([2011\)](#page-140-5) introduced the subphylum *Mortierellomycotina* based on phylogenetic analyses and morphological characteristics. However, Hoffman et al.  $(2011)$  accommodated *Mortierellomycotina* in '*Zygomycota*'. Tedersoo et al. ([2018\)](#page-157-4) introduced the phylum *Mortierellomycota* to



**Fig. 39** Morphology of *Lobulomyces angularis* JEL045. **a** Young thallus with branching of rhizoidal axis. **b**, **c** Mature thallus discharging zoospores (photo credit to Dr. Joyce E. Longcore)

<span id="page-102-0"></span>accommodate *Mortierella* and other genera included in *Mortierellomycotina.* Currently, the phylum comprises one subphylum, one class, one order and one family.

*Mortierellomycetes* Doweld, Index Fungorum 46: 1 (2013).

Index Fungorum Registration Identifer 550332; Figs. [53,](#page-116-0) [54](#page-117-0)

**Sexual morph:** Homo- and heterothallic formation of zygospores on suspensors lacking appendages, thick-walled, naked or dimpled (Takashima et al. 2016). **Asexual morph**: Rapid mycelial growth, often forming conspicuous rosettelike/zonate pattern on the plates. *Thallus* consisting of coenocytic, anastomosing, dichotomously branching hyphae, sometimes with rhizoids, chlamydospores or stylospores (aerial, stalked, spiny chlamydospores). *Sporangiophores* with infated base and attenuate apex, erect, branched or unbranched, initially not septated, potentially irregularily septated when fully stalked. *Sporangia* spherical, multispored; columella lacking or rudimentary. *Sporangiospores* globose to ellipsoid or irregular, either laevigate or ornamented (Hofmann et al. [2011;](#page-140-5) Dowel [2014f\)](#page-135-18). *Sporangiola* terminal, borne on erecting hyphae, ramifcations primarily horizontally expanding. *Sporocarps* containing sporangia are observed exclusively in the genus *Modicella* (Smith et al. [2013](#page-155-12)).

**Life modes and distribution:** Saprobic, plant-mutualistic or entomopathogenic in soil or organic substrates.

**Notes:** *Mortierellomycetes* belong to the phylum *Mortierellomycota*, the latter corresponding to the subphylum *Mortierellomycotina* (Tedersoo et al. [2018\)](#page-157-4). The family *Mortierellaceae* was reorganized based on a phylogenomic approach (Vandepol et al. [2020](#page-158-8)), resulting in 15 monophyletic genera: *Actinomortierella*, *Benniella*, *Dissophora*, *Entomortierella*, *Gamsiella*, *Gryganskiella*, *Haplosporangium*,

## *Linnemannia*, *Lobosporangium*, *Lunasporangiospora*, *Modicella*, *Mortierella*, *Necromortierella*, *Podila* and *Tyroliella.*

Members of *Mortierellomycetes* exhibit a global distribution across various climates (tropical, subtropical temperate, arctic-alpine) (Gams [1977](#page-137-2); Domsch et al. [1980](#page-135-19)), and thrive in snow-covered soil (Linnemann [1941;](#page-145-9) Wagner et al. [2013](#page-158-9); Telagathoti et al. [2021\)](#page-157-8). While most taxa are soil saprobes, some have been reported as plant mutualistic or entomopathogenic fungi. Additionally, many species are associated with endohyphal or epihyphal bacteria (Takashima et al. [2020](#page-156-11); Telagathoti et al. [2021](#page-157-8)).

**Type order:** *Mortierellales* Caval.-Sm., Biological Reviews of the Cambridge Philosophical Society 73: 246 (1998).

**Type family:** *Mortierellaceae* Luerss., Handbuch der systematischen Botanik mit besonderer Berücksichtigung der Arzneipfanzen 1: 63 (1877).

**Type genus:** *Mortierella* Coem., Bull. Acad. R. Sci. Belg., Cl. Sci., sér. 2 15: 536 (1863).

**Type species:** *Mortierella polycephala* Coem., Bull. Acad. R. Sci. Belg., Cl. Sci., sér. 2 16: 536 (1863).

*MUCOROMYCOTA* Doweld, *Prosyllabus Tracheophytorum*, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Index Fungorum Registration Identifer 90756.

Several members of the *Mucoromycota* produce visible colonies and conspicuous spores. *Hyphae* commonly diferentiates into rhizoids, chlamydospores, and stolons, with fxation, resistance, and dissemination functions, respectively. The *Mucoromycota* produce their sporangia by cleaving multinucleate cytoplasm. *Sporangiophores* terminate in a varied-shaped columella that persists and extends into sporangium, sporangiola, or in a varied-shaped vesicle where merosporangia or sporangiola arise from. Each sporangium has a wide range of sporangiospores, with some smaller ones

<span id="page-103-0"></span>**Fig. 40** Taxa belonging to diferent classes of *Chytridiomycota*. **a** *Batrachochytrium dendrobatidis* (JEL197, type) in *Rhizophydiomycetes*. **b** *Thoreauomyces humboldti* (JEL095, type) in *Spizellomycetes.* **c**–**e** *Neokarlingia chitinophila* (JEL510) in *Polychytriomycetes*. **c** Zoosporangium. **d** Zoosporangium with multiple papillae. **e** Resting spores. (photo credit to Dr. Joyce E. Longcore)



(called sporangiola) having only one or a few spores, and others (called merosporangia) having 5–20 merosporangiospores (Benny et al. [2014](#page-131-18); Spatafora et al. [2016;](#page-155-6) Kendrick [2017\)](#page-142-8). Sexual stages are represented by zygospores with apposed or opposed suspensors in this phylum. Hyphae of compatible mating types give arise zygophores, which fuse haploid nuclei. Sporocarps with zygospores are formed in few species.

**Life modes and distribution:** Saprobes, ectomycorrhizal, endophytes, and facultative (non-haustorial) parasites of fungi, plants and animals, causing mucormycosis mostly in immunocompromised patients.

**Note:** *Mucoromycota* was frst described by Doweld [\(2001](#page-135-0)), but it was previously introduced as "nom. nov," based on *Melanomycotina* (Cavalier-Smith [1998](#page-133-2)). Fungi of this phylum have been formerly included in phylum '*Zygomycota*' (identity is invalid now). The information was derived from molecular and phylogenetic evidence on monophyly and divergence times. Zhao et al. [\(2023\)](#page-162-1) updated the *Mucoromycota* classifcation framework. In particular, *Mucoromycota* fungi are characterized by their reproductive structures. Species that produce stolons anchor their hyphae to the substrate with rhizoids, which penetrate the surface of the host and grow in saprotrophic way. During maturity, most species have a fragile sporangium wall that is easily disrupted, allowing sporangiospores to escape. Some species have an apophysis below the columella, which is an expansion of the sporangiophore. For sexual reproduction, trisporic acid initiates the conversion of vegetative hyphae to zygophores (Benny et al. [2014\)](#page-131-18) that enlarge after making contact by expanding the cell wall forming gametangia, which are adherent cells that contain the haploid nuclei. The zygospore is formed after plasmogamy and karyogamy in a zygosporangium, which is a thick-walled, smooth or with warty projections structure. It is most common for a zygosporangium to germinate into a germ sporangium, germ sporangiolum or germ merosporangium in which spores are produced (Alexopoulos et al. [1996](#page-130-8); Benny [2012\)](#page-131-19). Some specimens of *Mucoromycota* can form azygosporangia, which are zygosporangia-like structures formed by parthenogenesis and not by sexual reproduction (Wagner et al. [2020](#page-158-10)).

## *Endogonomycetes* Doweld, Index Fungorum 57: 1 (2014). Index Fungorum Registration Identifer 550357; Fig. [55](#page-118-0)



<span id="page-104-0"></span>**Fig. 41** Representative characters of *Capillidiaceae*. **a** *Capillidium adiaeretum* (capilliconidia). **b** *Capillidium adiaeretum* (primary conidiophore). **c** *Capillidium rhysosporum* (primary conidiophore). **d**

Coenocytic, flamentous, underground sporocarps. Usually hypogeous or rarely epigeous, saprobic, and free-living, some forming ectomycorrhizal associations. In addition to coenocytic somatic hyphae, there may be septa that include micropores. Reproduction by globose or ovoid zygospores with unequal or nearly equal suspensors that are formed in sporocarps. Sexual and somatic hyphae are similar (Benny [2005](#page-131-20); Desirò et al. [2017\)](#page-135-20).

**Life modes and distribution:** Saprobes and ectomycorrhizal, widespread.

**Note:** *Endogonomycetes* (*Endogonales*) may have been one of the earliest mycorrhizal partners associated with land plants (Chang et al. [2019](#page-133-12)). There is a lack of collection of these fungi, and molecular data are needed to determine phylogenetic afliations for many taxa. Mostly, *Endogonales* lineages cannot be maintained *in vitro*, similarly to *Glomeromycotina*, obligately biotrophs mycorrhizal fungi (Spatafora et al. [2016](#page-155-6); Desirò et al. [2017\)](#page-135-20). The isolation and maintenance of a few species have, however, been successful

*Capillidium heterosporum* (secondary conidia). **e** *Capillidium rugosum* (zygosporus). **f** *Capillidium rhysosporum* (primary conidia). Bars: **a**–**f**=20 μm

in the laboratory with considerable effort (Field et al. [2015](#page-136-5); Yamamoto et al. [2017](#page-161-8)). Desirò et al. ([2017\)](#page-135-20) developed a multigene tree of *Endogonales* based on sporophores collected from four continents over the past three decades and found two deeply divergent phylogenetic clades within *Endogonales* with signifcant unexplained diversity. Typical mycorrhizal lifestyle signatures are found in the genomes of *Endogonaceae* and suggest that *Endogonales* might have colonized land through mycorrhizal associations with plants during their early evolution (Chang et al. [2019\)](#page-133-12). Aside from ectomycorrhizal species, *Endogonomycetes* (*Endogonales*) also include saprobic species (Bidartondo et al. [2011](#page-132-12)). According to Tedersoo and Smith (2013), ectomycorrhizae evolved twice within *Endogonales*. In the *Endogonomycetes*, zygospores are produced in sporocarps, and (asexually) chlamydospores can also be formed (Bidartondo et al. [2011](#page-132-12); Desirò et al. [2014;](#page-135-21) Dickie et al. [2015](#page-135-22); Chang et al. [2019](#page-133-12)).

**Type order:** *Endogonales* Jacz. & P.A. Jacz., Opredelitel' Gribov, I Ficomiteti (Leningrad): 8 (1931).



<span id="page-105-0"></span>**Fig. 42** *Entorrhiza casparyana*. **a–g** Intracellular more or less coiled hyphae and teliospores in diferent developing stages (host cells not drawn). **h–k** Germinating teliospores in diferent develop-

ing stages. Note the sigmoid shape of the propagules. Scale bar **a**–**g,**   $\mathbf{i}-\mathbf{k} = 20 \, \mu \text{m}$ ,  $\mathbf{h} = 10 \, \mu \text{m}$  in (h) (redrawn from Bauer et al. [2015](#page-131-12))

**Type family:** *Endogonaceae* Paol., Syll. fung. (Abellini) 8: 905 (1889).

**Type genus:** *Endogone* Link, Mag. Gesell. naturf.Freunde, Berlin 3(1–2): 33 (1809).

**Type species:** *Endogone pisiformis* Link, Mag. Gesell. naturf. Freunde, Berlin 3(1–2): 33 (1809).

*Mucoromycetes* Doweld, *Prosyllabus Tracheophytorum*, Tentamen Systematis Plantarum Vascularium (Tracheophyta) (Moscow): LXXVII (2001).

Index Fungorum Registration Identifer 90755; Fig. [56](#page-119-0)

*Mycelium* is abundant, rapidly growing, and anamorphic structures are normally formed in large quantities in it. Mycelium consists of non-septate or irregularly septated cells (Walther et al. [2019](#page-159-8); Wagner et al. [2020](#page-158-10)). *Sporangiospores* in anamorphic form are produced in multispored sporangia, as well as in one to few-spored sporangiola and merosporangia. The sporangium can be deliquescent, evanescent or persistent-walled, and the release of spores occur actively (less common) or passively. Most species do not produce chlamydospores, arthrospores, or yeast cells, (Von Arx [1982\)](#page-158-11). *Columella* with diferent shapes characterizes sporangia and some sporangiola, but merosporangia and some sporangiola are also formed on a varied-shaped vesicle. *Mucoromycetes* have well-developed columella or vesicle as synapomorphies. *Zygospore* (meiospore) is formed when homothallic or heterothallic species conjugate with each other (Hoffmann et al.  $2013$ ). In addition to the specifc ornamentation of the exospores (warty, rough, smooth), zygospores also display protective appendages such as antler-like or fnger-like on their support cells (suspensors) (Zycha et al. [1969](#page-162-2)). Some *Mucoromycetes* species exhibit dimorphism, capable of switching from flamentous, multicellular to yeast-like states (Bartnicki-Garcia and Nickerson [1962\)](#page-131-21). It has been revealed through the application of molecular phylogenetics that some of the above-mentioned morphological characters are not taxonomically signifcant (Hurdeal et al. [2021\)](#page-140-7).

**Life mode and distribution:** *Mucoromycetes* (*Mucorales*) are cosmopolitan fungi. In the fungal tree of life, this class belongs to the early-diverging fungal groups (Spatafora et al. [2016\)](#page-155-6). However, there is no information on the ecological role or geographical distribution of most of these species (Voigt



**Fig. 43** Types of symbiotic association in *Glomeromycota*. AMF form mycorrhiza with roots of terrestrial and aquatic plants. *Geosiphon pyriforme*, occurring in humid soil environments, is the unique

<span id="page-106-0"></span>et al. [2021\)](#page-158-3). Species of this class can be isolated from varied substrates, including soil, fruits, leaves, grains and stored foods, animal excrement and insects (Nguyen et al. [2021](#page-148-11); Voigt et al. [2021](#page-158-3)) (Table [22](#page-120-0)). In general, specimens of *Mucoromycetes* do not have the ability to degrade complex sugars, such as cellulose and lignin, but they can easily degrade soluble sugars, such as glucose, being referred to by some authors as "sugar fungi" (Benny [2012\)](#page-131-19). However, some specimens are capable of degrading cellulose, pectin, starch, and xylan (Pawłowska et al. [2019\)](#page-149-11). *Mucoromycetes* tend to be, among fungi, pioneers in the colonization of diferent substrates. Therefore, rapid sporulation and accelerated mycelial growth, even in nutrient-poor culture media, are striking characteristics of most taxa, especially *Mucorales* species (Benny [2008](#page-131-22); Benny et al. [2016](#page-131-23)). Soil and herbivores and small rodents dung are the best substrates for the isolation of these fungi.

fungus of the *Archaeosporomycetes* that forms an association with photosynthetically and nitrogen-fxing cyanobacteria of the genus *Nostoc*

*Pilobolus* is the the unique obligatory coprophilous genus, and its species present mechanisms of active expulsion of the sporangium with positive phototropism, adhesive sporangia, sporangiospores that survive digestion by herbivores, and the ability to grow in relatively high pH conditions (Dix and Webster [1995\)](#page-135-23). Some species of *Cunninghamella*, *Lichtheimia*, *Mucor* and *Rhizopus*, *Rhizomucor*, *Syncephalastrum* have been reported as pathogens of human (Table [23](#page-120-1)).

**Notes:** Among the early diverging fungi, the *Mucoromycetes* (*Mucorales*) are a well-studied group of fungi. A mucoralean fungus reproduces asexually and is classifed into diferent genera and families based on its vegetative and anamorphous structures such as spores or sporangia (Von Arx [1982\)](#page-158-11). In most keys to families and genera, teleomorphs (zygospores) have been neglected or only incidentally used.



<span id="page-107-0"></span>**Fig. 44** Microscopic morphology of taxa in *Glomeromycota*. **a** Spore wall of *Ambispora appendicula* (*Ambisporaceae*—*Archaeosporales*). **b** Detail of sporiferous saccule and spore in *Acaulospora tuberculata* (*Acaulosporaceae—Diversisporales*). **c** Spore with three walls in *Acaulospora* sp. (*Acaulosporaceae—Diversisporales*). **d** Detail of subtending hyphae in *Diversispora varaderana* (*Diversisporaceae— Diversisporales*). **e** Clusters of spores produced by *Dominikia gan-*

Over 100 years ago, heterothallic and homothallic modes of sexuality were discovered in *Mucoromycetes* (Blakeslee [1904\)](#page-132-13). The formation of sexual reproductive structures is mediated by volatiles such as trisporoids and beta-carotene (Gooday [1968\)](#page-137-3). We are unsure whether zygospores play a greater role in reproduction and distribution than asexual spores. Germination has only been described and illustrated for a few species under laboratory conditions (Michailides and Spotts [1988;](#page-147-10) Yu and Ko [1997](#page-161-9); Wagner et al. [2020](#page-158-10)). *Mucoromycetes* are traditionally classifed using physiology

*suensis* (*Glomeraceae*—*Glomerales*). **f** Detail of Melzer reaction in spores of *Epigeocarpum crypticum* (*Glomeraceae*—*Glomerales*). **g** Spore with suspensor-like cell and germinal shield in *Fuscutata aurea* (*Dentiscutataceae*—*Gigasporales*). **h** Suspensor cell in *Gigaspora* sp. (*Gigasporaceae*—*Gigasporales*). **i** Large glomerocarps produced by *Rhizoglomus maiae* (*Glomeraceae*—*Glomerales*). Scale bars **a**, **c**, **d**, **e**=10 µm; **b**, **g**, **h**=20 µm; **i**=250 µm

and morphology but species, genera, and families can be distinguished by only a few distinguishable morphological characters in it. It is difficult to identify the family structure of the *Mucorales*, but molecular data and high-resolution microscopy have enabled the discovery of new, potentially phylogenetic informative characters. The most recent signifcant changes were done by Zhao et al. ([2023](#page-162-1)).

**Industrial applications:** *Mucoromycetes* are used in a wide range of biological transformations, including the production
**Fig. 45** *Asellaria gramenei, A. saezii* and *A. ligiae*. The species of *Asellaria* are here represented to easily compare shapes and sizes. Indicated here are: bc, basal cell; bcb, basal cell branches; at, arthrospores; at' arthrospores not disarticulated from the thallus; tb, thallial branches. Scale bar =  $50 \mu m$  for all the fgures (redrawn from Valle [2006\)](#page-158-0)



of feed and food additives, pharmaceutical ingredients (including lycopene), and chitosan, an exclusive *Mucoromycetes* (*Mucorales*) product (Shahidi et al. [1999](#page-154-0); Gładkowski et al. [2011\)](#page-137-0). Some species have been traditionally used in the production of fermented foods in Asia, such as tofu and tempeh (Hesseltine [1965,](#page-139-0) [1983;](#page-139-1) Nout and Kiers [2005](#page-149-0)). Several *Mucoromycetes* species have been reported as plant pathogens (especially infecting fruits and vegetables) (Martin [1964](#page-146-0); Ray and Ravi [2005](#page-151-0)), facultative parasites of other fungi (Hoffmann et al. [2013](#page-140-0)), and biotrophic or necrotrophic parasites (not well studied) (Zycha et al. [1969\)](#page-162-0) (Table [24](#page-121-0)).

**Type order:** *Mucorales* Dumort. [as 'Mucorarieae'], Analyse des familles des plantes: avec l'indication des principaux genres qui s'y rattachent: 73 (1829).

A. ligiae

**Type family:** *Mucoraceae* Fr. [as 'Mucoroidei'], Syst. mycol. (Lundae) 1: xlix (1821).

**Type genus:** *Mucor* Fresen., Beitr. Mykol. 1: 7 (1850). **Type species:** *Mucor mucedo* Fresen., Beitr. Mykol. 1: 7 (1850).

*Umbelopsidomycetes* Tedersoo, Sánchez-Ramírez, Kõljalg, Bahram, Döring, Schigel, T. May, M. Ryberg & Abarenkov, Fungal Diversity 90: 152 (2018).

Index Fungorum Registration Identifer 554017.

A. saezii

**Fig. 46** *Barbatospora ambicaudata* **a** Detaching with a cylindrical wall that resembles a cap at the terminal end. **b** Sporulating branchlets removed from a simuliid larva. **c** Thallus holdfast affixed to a larva's hindgut cuticle. **d**–**g** A reconstruction of a sequence of development in released trichospores, following dehiscence of the cap-like structure at the terminal end, displaying numerous appendages basally and appendage-like flaments distally (modifed drawing from White et al. [2006\)](#page-159-0)



*Colonies* velutinous, reddish, brownish, grey or ochraceous, slow growing in culture media. *Sporangiophores* arising from vesicles on substrates, cymosely or umbellately branched (Wang et al. [2013\)](#page-159-1). Septa are near the branching and distant from the sporangium. **Asexual reproduction** occurs through elongate or globose, uni- and/or multispored ochraceous, reddish, or hyaline sporangium, and chlamydospores. *Columellae* are small or absent, and sporangiospores are varied-shaped, angular or not, hyaline or pigmented, in some species bearing appendages (Benny [2005](#page-131-0); Wang et al. [2013\)](#page-159-1). *Chlamydospores* are abundant and full of lipids in culture. Sexual reproduction are unknown (Spatafora et al. [2016](#page-155-0)).

**Life mode and distribution:** The majority of species are saprobes in soil, debris, dung and sugar-rich plant parts (Spatafora et al. [2016\)](#page-155-0). Although *Umbelopsidomycetes* are cosmopolitan fungi, there is no information on the ecological role or geographical distribution of the majority of these fungi.

**Notes:** *Umbelopsidomycetes* was introduced by Tedersoo et al. [\(2018](#page-157-0)) based on divergence time. Previously, species of *Umbelopsidomycetes* (*Umbelopsidales*) were classifed as *Mucoromycetes* (*Mucorales*) and *Mortierellomycetes*

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(*Mortierellales*). Based on molecular data, Meyer and Gams ([2003](#page-147-0)) delineated *Umbelopsis* from *Mortierella* and introduced *Umbelopsidaceae* as a new family. Following data from genome-scale analyses, this group appears to be distinct from *Mucorales* on a phylogenetic level (Spatafora et al. [2016\)](#page-155-0). Some *Umbelopsidomycetes* (*Umbelopsis*) are heat-resistant (Bollen and van der Pol-Luiten [1975\)](#page-132-0), but they are not able to grow at extremely high temperatures.

**Type order:** *Umbelopsidales* Spatafora, Stajich & Bonito, in Spatafora, Chang, Benny, Lazarus, Smith, Berbee, Bonito, Corradi, Grigoriev, Gryganskyi, James, O'Donnell, Roberson, Taylor, Uehling, Vilgalys & White, Mycologia 108(5): 1035 (2016).

**Type family:** *Umbelopsidaceae* W. Gams & W. Mey., in Meyer & Gams, Mycol. Res. 107(3): 348 (2003).

**Type genus:** *Umbelopsis* Amos & H.L. Barnett, Mycologia 58(5): 807 (1966).

**Type species:** *Umbelopsis versiformis* Amos & H.L. Barnett, Mycologia 58(5): 807 (1966).

*NEOCALLIMASTIGOMYCOTA* M.J. Powell, in Hibbett et al., Mycol. Res. 111(5): 516 (2007).

Index Fungorum Registration Identifer 501279; Fig. [57](#page-122-0)



**Fig. 47** *Dimargaris xerosporica*. **a** Habit sketches of sporophores.×30. **b** Typical sporophore showing characteristic cymose branching pattern. The lowermost fertile head is the oldest; the one at the tip of the apparent main axis (upper right) is the yourgest. $\times$ 145. **c** 

Obligate anaerobic fungi, present in the alimentary tract of foregut, pseudoforgut, and hindgut fermenting mammalian and few reptilian and avian herbivores. Lifecycle involves the production of and release of motile fagellated spores (zoospores) from sporangia. These zoospores encyst, germinate and develop into a thallus structure that anchors the formation of new sporangia. Produces posteriorly monofagellated or polyfagellated spores. Thallus development pattern are monocentric or polycentric, rhizoidal growth is either flamentous or bulbous. Lack mitochondria but contains hydrogenosomes of mitochondrial origin. Lack respiratory capacity and produce energy solely by fermentation. Produces cellulosomal structures bearing plant biomass degrading enzymes.

**Life modes and distribution:** Anaerobic gut fungi of herbivores, widespread.

Haustorium.  $\times$  1080. **d**–**f** Three successive stages in the development af the sporiferous branchlets and merosporangia.×1360. **g** Three mature merosporangia prior to disappearance of the sporangia wall and four sporangiospores. $\times$ 1360 (redrawn from Benjamin [1965\)](#page-131-1)

**Notes:** The members of *Neocallimastigomycota* have mainly been reported from 'the rumen and alimentary tract of larger mammalian and some reptilian, marsupial and avian herbivores' (Hanafy et al. [2022](#page-138-0)). The phylogenetic placement of these taxa has been confrmed in early divergent fungi by Li et al. [\(1993\)](#page-145-0) who introduced the Order *Neocallimastigales* J.L. Li et al. (in *Chytridiomycota*) to accommodate *Neocallimastigaceae* I.B. Heath.

In their phylogenetic analyses, James et al. [\(2006b\)](#page-141-0) showed that the *Neocallimastix* strain resided in *Chytridiomycota* along with *Chytridiales*, *Monoblepharidales*, and some *Spizellomycetales* and further named this clade as 'euchytrids'. White et al. [\(2006\)](#page-159-0) also agreed on the placement of *Neocallimastix* in *Chytridiomycota* in their analyses. However, both analyses by James et al. ([2006b](#page-141-0)) and White et al. ([2006\)](#page-159-0) used only one strain to represent *Neocallimastigales.* James et al. [\(2006b](#page-141-0)) provided a detailed analysis



**Fig. 48** Sporulating thallus of *Harpella melusinae* and one released spore (from Reichle and Lichtwardt [1972](#page-151-1))

of *Chytridiomycota* and recognized *Neocallimastigales* as a well separated, order in the phylum. Hibbett et al. ([2007](#page-139-2)), thus, introduced the class *Neocallimastigomycetes* and the phylum *Neocallimastigomycota* to accommodate these taxa which how a distinct phylogenetic lineage in the kingdom of *Fungi*. Hibbett et al. [\(2007](#page-139-2)) was broadly accepted by subsequent studies (e.g. Gruninger et al. [2014](#page-137-1); Powell and Letcher [2014](#page-150-0); Hanafy et al. [2022](#page-138-0)). However, Spatafora et al. ([2016\)](#page-155-0) recognized *Neocallimastigomycetes* as a distinct class in *Chytridiomycota*.

Tedersoo et al. ([2018](#page-157-0)), Wijayawardene et al. ([2020a\)](#page-160-0) and Voigt et al. [\(2021\)](#page-158-1) have recently recognized *Neocallimastigomycota* as a distinct phylum in their phylogenetic analyses. Currently, *Neocallimastigomycota* comprises one subphylum (*Neocallimastigomycotina*) and one class (*Neocallimastigomycetes*).

*Neocallimastigomycetes* M. J. Powell, Mycol. Res. 111:516. 2007.

Index Fungorum Registration Identifer 501280; Fig. [55](#page-118-0) Description and notes are similar to the phylum description.

**Type order:** *Neocallimastigales* J.L. Li, I.B. Heath & L. Packer [as 'Neocallimasticales'], Can. J. Bot. 71: 403 (1993).

**Type family:** *Neocallimastigaceae* I.B. Heath [as 'Neocallimasticaceae'], in Heath, Bauchop & Skipp, Can. J. Bot. 61(1): 304 (1983).

**Type genus:** *Neocallimastix* Vávra & Joyon ex I.B. Heath, in Heath, Bauchop & Skipp, Can. J. Bot. 61(1): 304 (1983). **Type species:** *Neocallimastix frontalis* (R.A. Braune) Vávra & Joyon ex I.B. Heath, in Heath, Bauchop & Skipp, Can. J. Bot. 61(1): 306 (1983).

*OLPIDIOMYCOTA* Doweld, Index Fungorum 42: 1 (2013). Index Fungorum Registration Identifer 550327.

**Asexual morph:** Sporangium single, endobiotic, with cell walls; zoospores posteriorly unifagellate, with a single globule, cone-shaped striated rhizoplast fused to both the functional and vestigial kinetosomes, gamma-like particles and rough endoplasmic reticulum; nucleus associated with the basal body, no nuclear cap. **Sexual morph:** Unknown.

**Life modes and distribution:** The members of *Olpidiomycetes* distribute in a broad range of ecosystems (worldwide) and are mainly reported as plant pathogens and vectors of plant pathogenic viruses. Phytopathogenic members of the class are root parasites of important crop plants (e.g., *Olpidium*, *Cibdelia*, *Leiolpidium*), while some are reported as pathogenic to algae and rotifers (e.g., *Olpidium*). Some members are saprobes (e.g., *Chytridhaema*) that are important in decomposition and nutrient cycling.

**Notes**: Doweld [\(2013a\)](#page-135-0) introduced the phylum *Olpidiomycota* to accommodate *Olpidium* (A. Braun) Rabenh. At the same time, Doweld et al.  $(2013a)$  $(2013a)$  introduced the subphylum *Olpidiomycotina* Doweld. Before the introduction of the phylum, the phylogenetic placement of *Olpidium* taxa in basal fungi has been confrmed by James et al. [\(2006a](#page-141-1)[,b](#page-141-0)), White et al. [\(2006\)](#page-159-0) and Sekimoto et al. ([2011\)](#page-154-1) but regarded as uncertain in '*Zygomycota*'/ or '*Chytridiomycota*'. Hibbett et al. ([2007](#page-139-2)), however, did not include *Olpidium* in any higher position but regarded that traditionally the genus has been placed in *Chytridiomycota*. Spatafora et al. [\(2016\)](#page-155-0) did not include any *Olpidium* taxa in their phylogenetic analyses but mentioned the placement accepted by James et al.  $(2006a,b)$  $(2006a,b)$  $(2006a,b)$  and Sekimoto et al.  $(2011)$  $(2011)$ . Tedersoo et al.  $(2018)$  $(2018)$  $(2018)$ , however, clearly showed the placement of *Olpidium s. str.* as a distinct lineage in the kingdom *Fungi*, sister to *Basidiobolomycota*. Furthermore, Tedersoo et al. ([2018](#page-157-0)) introduced the subkingdom *Olpidiomyceta*, which only comprises one phylum.

### *Olpidiomycetes* Doweld, Index Fungorum 42: 1 (2013).

Index Fungorum Registration Identifer 550325; Fig. [58](#page-123-0) The description is similar to the description of the phylum.

**Fig. 49** *Kickxella alabastrina* Coemans. **a** Habit sketch showing general characteristics of the fruiting structures. $\times 30$ . **b** Once branched sporangiophore bearing distal whorls of sporocladia.×180. **c**, **d** Early stages in the development of the sporocladia showing their simultaneous origin in a whorl around the slightly swollen apex of the sporangiophore. $\times 600$ . **e–g** Successive stages in the development of the sporocladium and pseudophialides prior to sporangiole formation. $\times 600$ . **h** Mature sporoclaclium showing disposition of pseudophialides and sporangiola on upper surface.×600. **i** Mature sporocladium after sporangiola have fallen away; note furcate apical cell.×600. **j**–**m** Four pseudophialides showing successive stages in the aerogenous development of the unispored sporangiole.×1360. **n** Two mature sporangiola. × 1360 (redrawn from Benjamin [1958](#page-131-2))



**Notes**: *Olpidiomycetes* was introduced by Doweld ([2013a](#page-135-0)). *Olpidiomycetes* encompasses a single order *Olpidiales*. *Olpidiales* was introduced by Cavalier-Smith ([2013](#page-133-0)) but it was invalid as per Article 42. Later, Doweld  $(2013a)$  $(2013a)$  effectively published the order and introduced a new class *Olpidiomycetes* to accommodate the order. While proposing the highlevel classifcation of the fungi and a tool for evolutionary ecological analyses. Recently Chang et al. ([2021\)](#page-133-1) confrmed genome-scale phylogeny of *Olpidium* as the closest living zoosporic fungus to the non-fagellated, terrestrial fungi (whole-genome sequence of *Olpidium bornovanus* S191: JAEFCI000000000**)**. Previously, Sekimoto et al. ([2011\)](#page-154-1) placed *Olpidium* under *Zygomycota* through molecular phylogenetic analyses of *tef2*, *rpb1*, *rpb2* and *ACT*.

**Type order:** *Olpidiales* Caval.-Sm., Eur. J. Protist. 49(2): 157 (2012) [2013].

**Type family:** *Olpidiaceae* J. Schröt. [as 'Olpidiacei'], in Cohn, Krypt.-Fl. Schlesien (Breslau) 3.1(1–8): 180 (1886) [1889].

**Type genus:** *Olpidium* (A. Braun) Rabenh., Flora Europaea algarum aquae dulcis et submarinae 3: 282 (1868) [1867]. **Type species: not indicated**

(*Olpidium endogenum* (A. Braun) J. Schröt. 1886 (Designated by Clements & Shear, Gen. fung., Edn 2 (Minneapolis): 234. 1931 *fde* Index Fungorum 2023).

*ROZELLOMYCOTA* Doweld, Index Fungorum 43: 1 (2013) emend. Tokarev & Nassonova.

**Fig. 50** *Ramicandelaber longisporus* PCNB-1157 (IFO 33179). **a** Habits. **b**–**f** Upper parts of sporangiophores showing the development of sporocladia (branchlets on branches of sporangiophores). Sporocladia produce subspherical pseudophialides. Scale bar  $a$ – $f$  = 10  $\mu$ m (redrawn from Ogawa et al. 2001)



Index Fungorum Registration Identifer 550328; Fig. [59](#page-124-0) Intracellular obligate parasites parasitize host via phagocytosis of host organelles (in *Rozella*) or via osmotrophy (in all other known forms), penetration of the host via a polar tube. *Cells* amoeboid, multinucleate or uninucleate, covered by a single membrane, unwalled; pseudopodia-like extensions interdigitated between and partially extended around host organelles; microtubules underlie the plasma membrane in a random pattern and extend into cytoplasmic extensions. The most basal lineage of the phylum possesses bilaterally symmetrical posteriorly unifagellate zoospores while other groups are non-motile*. Resting spores* thick-walled. With the exception of *Rozella*, *Mitosporidium,* and *Paramicrosporidium*, the mitochondria are reduced to mitosomes lacking the genetic apparatus (modifed after Doweld [2013c](#page-135-1)).

**NB:** The phylum *Rozellomycota* sensu Wijayawardene et al. [\(2020a\)](#page-160-0) in its current state lacks some important subdivisions (classes), which would incorporate a number of basal forms. This should be noted when dealing with its taxonomic framework, limited to two classes *Rudimicrosporea* and *Microsporidea,* which only cover the subphylum *Microsporidia,* leaving out a series of important taxa either as genera *incertae sedis* (*Mitosporidium*, *Morellospora*, *Nucleophaga*, *Paramicrosporidium*, *Rozella*) or as orders *incertae sedis* (*Chytridiopsida*).

**Life modes and distribution:** cosmopolitan obligate intracellular parasites of *Chytridiomycota*, *Blastocladiomycota*, *Monoblepharomycota*, and *Oomycota* (Karpov et al. [2014c](#page-142-0)), *Bacillariophyceae* (Chambouvet et al. [2019](#page-133-2)),



**Notes:** Karpov et al. [\(2014a](#page-142-0)) introduced the superphylum *Opisthosporidia*, which accommodated the phyla *Cryptomycota/Rozellida, Microsporidia,* and the newly established *Aphelida*, the latter based upon the class *Aphelidea.* This taxonomic solution refects the deep branching of this monophyletic group within *Holomycota* as a sister group to the kingdom *Fungi.* Further studies did not support the monophyly of Opisthosporidia (Torruella et al. [2018](#page-157-0); Galindo et al. [2022](#page-137-2)), and Tedersoo et al. [\(2018](#page-157-0)), further supported by Wijayawardene et al. ([2018b](#page-160-1), [2020a](#page-160-0), [2022a](#page-160-2)), placed *Cryptomycota/Rozellida* and *Microsporidia* as basal *Fungi*, bounded by the phylum *Rozellomycota* (Vávra and Lukeš [2013\)](#page-158-2)*.* In the frames of the current compendium, which represents "a mycologist's view" of taxonomy, we are called upon using this affiliation, though we strongly believe it being debatable and requires further consideration. Meanwhile, the original diagnosis by Doweld ([2013c\)](#page-135-1) only described the *Rozella* part of the phylum, while nowadays a number of organisms included in the phylum has increased dramatically, which has provoked a need to modify the diagnosis. Since the new members of the phylum all parasitize host cells by importing nutrients from its cytoplasm and organelles (Vavra and Lukes [2013](#page-158-2); Haag et al. [2014](#page-138-1); Corsaro et al. [2014;](#page-134-0) Timofeev et al. [2020\)](#page-157-1), not phagocytozing it, this part of diagnosis is also modifed.

**Fig. 51** *Monoblepharis macrandra.* Nomarski interference micrographs. **a** Solitary oogonium showing basal cross wall and the receptive papilla and lipid droplets of the egg. **b** Terminal antheridium cut off by basal cross wall prior to cleavage of sperms. **c** Oogonium with hypogynous antheridium. **d** Exogenous oospore showing bullations, thick wall, and attachment to mouth of empty oogonium. **e** Zoosporangium showing basal cross wall and cleavage planes of zoospores. Scale bar **a**–**f**=10 µm (redrawn from Marek [1984\)](#page-146-1)





*Rudimicrosporea* Sprague, Comp. Pathobiol. (New York, London): 1–31 [\(1977](#page-155-1)) emend. Tokarev & Nassonova.

Synonym: *Metchnikovellidea* Weiser, Vestnik Cesk. Spol. Zool. 41: 312 (1977).

Index Fungorum Registration Identifer 90611.

Minute, intracellular protozoan parasites. They are exclusively hyperparasites of gregarines, which are all (with one possible exception) in turn parasites of annelids, mostly marine. The spores have the essential characters of microsporidian spores, but difer sharply in having a highly simplifed or rudimentary extrusion apparatus missing a polar sac/ anchoring disk, a polaroplast and a posterior vacuole. The polar flament (tube) represented by a manubrium (instead of a long, coiled flament) that extends posteriorly from a polar sac and terminates in a tubulo-vesicular network. The sporoplasm, consisting of nucleus and cytoplasm, occupies the rest of the intrasporal cavity. The spore wall consists of the usual three layers (exospore, endospore, plasmalemma), but endospore is rather thin, almost undeveloped. The spores, having the simple extrusion apparatus, are roughly spherical or oval and quite small. In the vegetative sequence uninucleate and plasmodial forms have been reported. A dimorphism is exhibited in the occurrence of two sporulation (sporogonial) sequences. In one sequence the plasmodium divides and produces numerous sporoblasts which transform into free spores resided either directly in the cytoplasm or within a parasitophorous vacuole. In the other sequence small plasmodia become encysted and produce a small, often constant number of relatively small, uninucleate spores within a cystic wall (spore sac) (modifed after Sprague et al. [1977](#page-155-1)).

**Life mode and distribution:** cosmopolitan obligate intracellular parasites of *Apicomplexa,* namely *Gregarinasina,* which predominately parasitize marine annelids (Nassonova et al. [2016](#page-148-0)).

**Notes:** The last paragraph of the diagnosis is omitted as it gives a synopsis of *Rudimicrosporea* as of 1977, mentioning



**Fig. 52** Light microscopic images of *Monoblepharidomycetes*: **a**–**c** *Harpochytrium* sp. (after: Powell and Letcher [2014](#page-150-0), modifed), **d**–**g** *Telasphaerula gracilis* (after: J.E. Longcore in: Karpov et al. [2017a,](#page-142-1) modifed). **a** Germination of zoospore cysts (cy) of *Harpochytrium* sp. in an uniaxial thallus with basal holdfast (arrow). **b** *Harpochytrium* sp. thallus with highly vacuolated, foamy appearing

three genera with the respective number of species known by that time, and their geographical distribution. Since the taxonomic composition has been extended since then, including erection of one more family (Larsson [2000\)](#page-144-0), the beginning of the diagnosis is also be the subject of change. The term *sporocyst* (as well as the term *cyst*) is not currently in use, and it is replaced by the term *spore sac* (Larsson [2014\)](#page-144-1).

**Type order:** *Metchnikovellida* Vivier, Protistologica 11(2): 345–361 (1975).

**Type family:** *Metchnikovellidae* Caullery & Mesnil (1914) emend. Larsson [2014.](#page-144-1) Microsporidia: Pathogens of Opportunity. Ames, Iowa. 605–634. (2014).

**Type genus:** *Metchnikovella* Caullery & Mesnil C. R. Hebd. Seanc. Acad. Sci. Paris 125: 789 (1897).

**Type species:** *Metchnikovella spionis* Caullery & Mesnil C. R. Hebd. Seanc. Acad. Sci. Paris 125: 789 (1897).

*Microsporidea* Corliss & Levine, J. Protoozool. 10(Suppl.): 26–27 (1963) emend. Tokarev & Nassonova.

Index Fungorum Registration Identifer 90587.

Diagnosis: Obligate intracellular parasites with developmental sequences ended by formation of unicellular spores, containing uni- or binucleate germ (sporoplasm) and extrusion apparatus, unique in its structure, with the polar tube as its main part. The extrusion apparatus well developed.

cytoplasm (arrow). **c** *Harpochytrium* sp. cleaving zoospores beginning at apex (arrow) of thallus. **d** polycentric thallus of *Telasphaerula gracilis* consisting of swellings connected by fne isodiametric tubes. **e** zoospore recently emerged from adjacent zoosporangium (arrow). **f**, **g** zoospores with typical shape for *Monoblepharidales*. Scale bars: **a**–**g** 10 µm

Flagella, centrioles, and lysosomes absent, mitochondria reduced to mitosomes, ribosomal RNA shortened highly contracted, providing giving the appearance of prokaryotictype ribosomes molecules. Prespore developmental stages with uniform nuclei, either single or paired (diplokaryon). Nuclei haploid or diploid. Sexual process and intermediate host involvement possible. Infection performed via extrusion of polar tube which injects the germ into the host cell. Development either in direct contact with hyaloplasm and host cell organelles or within vacuole of host or parasite origin (modifed after Issi et al. 2008.)

**Life mode and distribution:** cosmopolitan obligate intracellular parasites of *Ciliata, Paramyxea,* and *Metazoa* (Vavra and Lukes 2013; Stentiford et al. [2017](#page-156-0)).

**Notes:** The original diagnosis for the class *Microsporidea* is as following: "spore of unicellular origin; single sporoplasm; single valve; one or two long, tubular polar flaments through which sporoplasm escapes; cytozoic in invertebrates, especially arthropods, and lower (rarely higher) vertebrates" (Corliss and Levine 1963). It refected the fact *Microsporidia* and *Myxozoa (Myxosporidia)* were referred to as the two respective classes of the subphylum *Cnidospora* of the protozoan phylum *Sporozoa*. Modern taxonomy, strongly supported by molecular phylogeny data, showed

**Fig. 53** *Mortierella polycephala* (CBS 227.35, isolated by G. Linnemann). **a** Sporangia single, with irregular or with racemose branching. **b** Stalked chlamydospores (stylospores). **c** Smooth-walled chlamydospores among substrate hyphae, hyphae with oil drops and septa, tangle of hyphae. **d** Colony morphology on MEA and **e** on LcA after 1 week at 20 °C. Scale bars: **a**=50 μm, **b**–**c**=10 μm,  $d$ **-e**=1 cm



polyphyletic origin of this group and made this afliation obsolete. We therefore propose a redefnition of the class *Microsporidea*, based upon a more recent diagnosis of Issi and Voronin (2007) which was proposed in the frames of contemporary views of protistology (i.e., apart from *Fungi*). Those authors, however, preferred only to include one distinctive feature (well-developed extrusion apparatus), further indicating the diagnosis of the class corresponding to that of the phylum. Here, as the phylum *Microsporidia* is lowered to the subphylum rank in accordance with Wijayawardene et al. [\(2020a](#page-160-0)), the defnition of the class *Microsporidea* lacks its important attributes and we introduce the main features of the class corresponding to those of the phylum, in the class diagnosis itself.

**Type order:** *Nosematida* Labbe (1899) emend. Tokarev & Issi in Wijayawardene et al., Mycosphere 11(1): 1060–1456 (2020).

**Type family:** *Nosematidae* Labbe (1899) emend. Tokarev, Huang, Solter, Malysh, Becnel & Vossbrinck, J. Invertebr. Pathol. 169:107279 (2020).

**Type genus:** *Nosema* Naegeli, Bot. Ztg. 15: 760 (1857). **Type species:** *Nosema bombycis* Naegeli, Bot. Ztg. 15: 760 (1857).

*SANCHYTRIOMYCOTA* Galindo, López-García, Torruella, Karpov & Moreira, Nat. Commun. 12 (no. 4973): 9 (2021).

Index Fungorum Registration Identifer 558519; Fig. [60](#page-125-0)

Monocentric thallus, epibiotic; usually amoeboid zoospores with flopodia and immobile posterior pseudocilium **Fig. 54** Morphological and culture characteristics of *Mortierellomycetes* species. **a** Colony of *Mortierella alpina* on PDA plate. **b** Sporangiophore of *Entomortierella* sp. **c** Spores of *Entomortierella* sp. **d** Chain of gemmae of *Entomortierella* sp. stained in lactophenol cottonblue. **e** Cluster of gemmae of *Entomortierella* sp. Scale bars: **b**–**e** 50 µm



with longest-known kinetosome in fungi  $(1-2 \mu m)$ ; centrosome in sporangium of two centrioles composed by nine microtubular singlets (adapted from Galindo et al. [2021](#page-137-3)).

**Life mode and distribution:** Obligate parasites of freshwater algae, cosmopolitan.

**Notes:** Galindo et al. ([2021\)](#page-137-3) introduced this phylum to accommodate two obligate parasites of freshwater algae (which are called as 'Sanchytrids') viz*., Sanchytrium* Karpov & Aleoshin 2017 (type: *Sanchytrium tribonematis* Karpov & Aleoshin 2017 *fde* Karpov et al. [2017a](#page-142-1), [b](#page-142-2)) and *Amoeboradix* Karpov et al. [2017a,](#page-142-1) [b\)](#page-142-2) (type species: *Amoeboradix gromovii* Karpov et al. [2017a,](#page-142-1) [b](#page-142-2) *fde* Karpov et al. [2018](#page-142-3)). However, *Amoeboradix* 2017 and *Amoeboradix gromovii* 2017 are

invalid (Art. F.5.1, Shenzhen) and thus validated in Index Fungorum [\(2024](#page-141-2)).

Ribosomal DNA molecular phylogenies placed *Sanchytrium* in *Monoblepharidomycetes* (Karpov et al. [2017a\)](#page-142-1), and later among non-zoosporic fungi (Karpov et al. [2018\)](#page-142-3). Based on these data Tedersoo et al. ([2018](#page-157-0)) introduced *Sanchytriales* Tedersoo et al. and *Sanchytriomycetes* Tedersoo et al. to accommodate *Sanchytrium* (i.e., *Sanchytriaceae* Karpov & Aleoshin) and treated as a distinct clade in the phylum *Monoblepharomycota*. According to the multigene phylogeny (Galindo et al. [2021](#page-137-3)) both the *Sanchytrium* and *Amoeboradix* give a well-supported monophyletic lineage sister to the *Blastocladiomycota*.

Below we introduce new subphylum *Sanchytriomycotina*. Hence, the phylum consists of one subphylum and one class. <span id="page-118-0"></span>**Fig. 55** *Endogone* spp. and diferent stages of the life cycle. *Endogone pisiformis*: **a**–**e** Successive stages in the formation of zygospores. **f** Mature spores with gametes attached. **g** Peculiarly diferentiated hypha forming the superficial tomentum of a young individual. *Endogone multiplex***: h** Mature spore showing hyphal envelope. **i** Mature spore with discrete gamete attachments, treated with KOH. **j** Larger and smaller spore-groups surrounded by envelope with incorporated humus-material. *Endogone tuberculosa***: k** Section of sporocarp showing disposition of sporogenous areas, (light), with surrounding and penetrating earthy envelope (dark): magnifed about seven times as reduced. **l** Portion of gleba showing spores and pseudoparenchymtous hyphal tissue. **m** Spore showing envelope and origin from gametes. **n** Fully matured spore with greatly thickened exospore. **o**, **p** Spores of more elongate type. *Endogone incrassate***: q** Section of a portion of sporocarp, showing peridium and gleba with fully mature and more immature spores. **r** Spore showing characteristic fatty contents and probably sexual origin. **s** Fully mature spore with greatly thickened exospore and compacted surrounding hyphae (Redrawn from Thaxter [1922](#page-157-2))



#### *Sanchytriomycotina* Karpov *subphylum nov.*

Index Fungorum Identifer IF 558746.

The description is similar to the description of the phylum.

*Sanchytriomycetes* (Tedersoo et al.) Galindo, López-García, Torruella, Karpov & Moreira, Nat. Commun. 12 (no. 4973): 9 (2021).

Index Fungorum Registration Identifer 554005; Fig. [60](#page-125-0) The description is similar to the description of the subphylum.

**Type order:** *Sanchytriales* (Tedersoo et al.) Galindo, López-García, Torruella, Karpov & Moreira, Nat. Commun. 12 (no. 4973): 9 (2021).



**Fig. 56** *Mucor septatiphorus.* **a** Colony surface on PDA at 25 °C. **b** Colony reverse on PDA at 25 °C. **c** Sporangiophore with sporangium. **d**, **e** Branched sporangiophore with columellae. **f** Branched sporan-

**Type family:** *Sanchytriaceae* Karpov & Aleoshin, in Karpov, Mamanazarova, Popova, Aleoshin, James, Mamkaeva, Tcvetkova, Vishnyakov & Longcore, Fungal Biol. 121(8): 735 (2017).

**Type genus:** *Sanchytrium* Karpov & Aleoshin, in Karpov, Mamanazarova, Popova, Aleoshin, James, Mamkaeva, Tcvetkova, Vishnyakov & Longcore, Fungal Biol. 121(8): 735 (2017).

giophore with terminal columella and lateral sporangium. **g** Sporangiophore with columella. **h** Rhizoids. **i** Sporangiospores. Scale bars  $a - i = 20 \mu m$ 

**Type species:** *Sanchytrium tribonematis* Karpov & Aleoshin, in Karpov, Mamanazarova, Popova, Aleoshin, James, Mamkaeva, Tcvetkova, Vishnyakov & Longcore, Fungal Biol. 121(8): 735 (2017).

*ZOOPAGOMYCOTA* Gryganskyi, M.E. Sm., Spatafora & Stajich, in Spatafora, Chang, Benny, Lazarus, Smith, Berbee, Bonito, Corradi, Grigoriev, Gryganskyi, James, O'Donnell,





**Table 23** Medicinally/clinical signifcances of *Mucoromycetes*, examples:



Roberson, Taylor, Uehling, Vilgalys & White, Mycologia 108(5): 1035 (2016).

Index Fungorum Registration Identifer 816300.

*Haustoria* are produced by some of the animal pathogens and mycoparasites. *Zoopagomycota* hyphae may be compartmentalized by septa that may be complete or uniperforate; in the latter, bifurcate septa contain electron opaque lenticular plugs. *Zygospore* formation typically involves modifed hyphal tips, thallus cells, or hyphal bodies (yeastlike cells) that function as gametangia. The hyphal diameter is characteristically narrow in thalli that are branched or unbranched; sometimes specialized haustoria are produced in association with hosts. Only a handful of species have been successfully maintained in axenic culture. Sexual reproduction, where known, is by gametangial conjugation, forming globose zygospores on apposed diferentiated or undiferentiated suspensor cells. **Asexual reproduction** is by arthrospores, chlamydospores, conidia, or multispored merosporangia that may be simple or branched (extracted from Spatafora et al. [2016](#page-155-0)).

**Life mode and distribution:** Associations with animals (e.g. pathogens, commensals, mutualists) and mycoparasites.

**Notes:** Spatafora et al. ([2016](#page-155-0)) introduced the phylum *Zoopagomycota* after molecular evidence showed it formed a paraphyletic clade with *Mucoromycota*. The phylogenetic placement was confrmed by Wang et al. [\(2023c\)](#page-159-4). Since Hibbett et al. ([2007](#page-139-2)), *Zygomycota* was formally abandoned and the subphylum *Zoopagomycotina* Benny was proposed; the subphylum was originally identifed as belonging to the "zygomycetes" (James et al. [2006b](#page-141-0); White et al. [2006](#page-159-0); Sekimoto et al. [2011](#page-154-1); Ebersberger et al. [2012;](#page-136-0) Chang et al. [2015](#page-133-4)), a group created to accommodate basal fungi exclusively characterized by associations with animals and fungi. The class *Zoopagomycetes* was introduced by Dowell (2014 k)

<span id="page-121-0"></span>



and comprises *Zoopagales.* The order's monophyletic nature was confirmed by Davis et al. ([2019\)](#page-134-5) however the authors could not confrm their relationships with other subphyla due to the scarcity of available taxa.

#### *Zoopagomycetes* Doweld, Index Fungorum 60: 1. 2014.

Index Fungorum Registration Identifer 550358; Figs. [61,](#page-126-0) [62](#page-127-0)

*Thallus* simple, branched or none, or hyphal system extensively branched. Ectoparasites forming haustoria inside the host. Arthrospores, chlamydospores or uni- or multispored sporangiola (asexual reproduction); sporangiospores produced in simple or branched merosporangia. *Zygospores* globose, produced on diferentiated, straight or coiled suspensors. Endo- or ectoparasites, predaceous.

**Notes:** Currently, the class comprises one order *Zoopagales* (families *Cochlonemataceae*, *Helicocephalidaceae*, *Piptocephalidaceae*, *Sigmoideomycetaceae* and *Zoopagaceae*).

**Type order:** *Zoopagales* Bessey ex R.K. Benj., in Kendrick, The Whole Fungus 2: 590 (1979).

**Type family:** *Zoopagaceae* Drechsler ex Drechsler emend. Duddington, Mycologia 27(1): 30 (1935).

**Type genus:** *Zoopage* Drechsler Mycologia 27(1): 30 (1935).

**Type species:** *Zoopage phanera* Drechsler, Mycologia 27(1): 30 (1935).

# **Placements of fossil fungi in higher ranks**

*Fungi* are eukaryotic, uni- or multi-cellular, parasitic/saprobic, omnipresent organisms. Fossil fungi have been recorded in the form of dispersed spores, mycelia, sporocarps and mycorrhiza and are commonly observed in the palynological preparations or in thin sections of sedimentary rocks. These have been sporadically reported since long (Williamson [1878](#page-160-3), [1880;](#page-160-4) Kidston and Lang [1921](#page-142-4); Edwards [1922\)](#page-136-4) but their study received major attention with the amplifcation of palynological studies since the 1950s. Although fungal remains are encountered in the sediments of all ages, their fossil records indicate that *Fungi* got well established during the Cretaceous and became conspicuously abundant in the Tertiary Period (Jain [1974](#page-141-3); Jansonius [1976;](#page-141-4) Jain and Kar [1979;](#page-141-5) Ramanujam [1982;](#page-151-5) Kalgutkar and Jansonius [2000](#page-142-5); Tripathi [2009](#page-157-4); Saxena and Tripathi [2011;](#page-153-1) Saxena et al. [2021](#page-153-2)), suggesting that their proliferation is linked with diversifcation of angiosperms. Being fragmentary, fossil fungi generally lack characteristic features that are diagnostic of extant taxa, hampering their classifcation with extant fungi. However, a good number of fossil fungal taxa (other than spores) can be related to their extant counterpart (up to order/ family or even genus level). Here, only such legitimate genera, which could be related to extant taxa, have been considered, see below. Holotypes of type species of 27 genera are illustrated (Fig. [61\)](#page-126-0).

### 1. Phylum: *Ascomycota* Caval.-Sm.

*Ascomycota* taxa are known as the sac fungi, is the largest and most diversifed phylum of *Fungi* and is characterised by having septate hyphae with simple septal pores. The most important feature of this fungal group is the 'ascus' (pl. asci), a microscopic sexual structure in which non-motile spores, called ascospores, are formed. Asci are often housed in a fruiting body or ascocarp. However, a large number of species of *Ascomycota* are asexual, i.e. they do not have a sexual cycle and thus do not form asci or ascospores. As evident from the study of fossil fungi, the majority of dispersed fungal spores found in palynological preparations are

<span id="page-122-0"></span>**Fig. 57** Taxa of *Neocallimastigaceae.* **a**, **b** Zoospores and sporangia of *Neocallimastix frontalis*: (**a** Phase contrast microscopic illustration showing the *Neocallimastix frontalis* polyfagellated zoospores. **b** Phase contrast microscopic illustration showing the *Neocallimastix frontalis* sporangia. *Neocallimastix frontalis* produces globose shaped sporangia. Sporangiophores in *Neocallimastix frontalis* are short and often display an egg-cup like morphology). **c** *Anaeromyces contortus* Phase contrast microscopy showing hyphal structure. The genus *Anaeromyces* is characterized by displaying multiple constrictions, usually occurring at regular intervals resulting in a distinctive and taxonomically informative sausage-shaped morphology. **d** *Caecomyces communes*. Phase contrast microscopy showing hyphen structure. The genus *Caecomyces* does not produce atypical flamentous hyphae. Rather, *Caecomyces* spp. produces bulbous rhizoidal growth pattern. Only the genera *Caecomyces* and *Cyllamyces* display this characteristic bulbous rhizoidal growth pattern



produced by the *Ascomycota*. Ascospores are produced, usually in groups of four or eight, inside a sac (ascus) by the socalled 'perfect (sexual) stage' of these fungi. The 'imperfect (asexual) stage' may produce several types of asexual spores, named as conidiospores (or conidia).

The following fossil genera exhibit affinity with *Ascomycota*, but their resemblance with taxa of lower ranks could not be established: *Adendorfa* G.Woroviec et al., *Appianoporites* S.Y. Sm. et al., *Asteromites* Poinar, *Aureofungus* Hibbett et al., *Cashhickia* T.N. Taylor et al., *Casparyotorula* Rikkinen et al., *Cephalothecoidomyces* G. Worobiec et al., *Chlorolichenomycites* Honegger et al., *Entropezites* Poinar & R. Buckley, *Eomelanomyces* Beimforde et al., *Galloea* Alstrup & Søchting, *Honeggeriella* Matsunaga et al., *Jimwhitea* M. Krings & T.N. Taylor, *Palaeoanellus* A.R. Schmidt et al. and *Spheciophila* Poinar. Fossil ascomycetus fungi are represented by the following eight classes: *Dothideomycetes* O.E. Erikss. & Winka, *Eurotiomycetes* O.E. Erikss. & Winka, *Laboulbeniomycetes* Engl., *Lecanoromycetes* O.E. Erikss. & Winka, *Leotiomycetes* O.E. Erikss. & Winka, *Pezizomycetes* O.E. Erikss. & Winka, *Sordariomycetes* O.E. Erikss. & Winka and *Taphrinomycetes* O.E. Erikss. & Winka, as given below*.*

#### 1.1. Class: *Dothideomycetes* O.E. Erikss. & Winka

Class *Dothideomycetes* constitutes one of the most abundant classes of fossil *Ascomycota*. This class is represented by the following genera: *Alternaria* Nees, *Appendicisporonites* R.K. Saxena & S. Khare [1992](#page-153-3) (Fig. [63](#page-127-1)-a), *Asterina* Lév., *Asterinites* Doub. & D. Pons, *Asterothyrites* Cookson, *Brefeldiellites* Dilcher, *Caldesites* Puri, *Callimothallus* Dilcher [1965](#page-135-2) (Fig. [63-](#page-127-1)b), *Cribrites* R.T. Lange, *Cryptodidymosphaerites* Currah et al., *Cucurbitariaceites* R.K. Kar et al. [1972](#page-142-6) (Fig. [63](#page-127-1)-c), *Dictyosporiuminites* Debi Mukh., *Dictyotopileos* Dilcher, *Epicoccum* Link, *Euthythyrites* Cookson, *Haplopeltis* Theiss., *Hysterites* Gӧpp., *Kalviwadithyrites* M.R. Rao in R.K. Saxena [2012](#page-153-4) (Fig. [63-](#page-127-1)d), *Koshalia* S. Sarkar & V. Prasad [2003](#page-153-5) (Fig. [63](#page-127-1)-e), *Leptosphaerites* Richon, *Mariusia* D. Pons & Boureau, *Melanosporites* Pampal., *Microthallites* Dilcher, *Microthyriacites* Cookson, *Microthyrites* Pampal., *Molinea* Doub. & D. Pons, *Palaeoasterina* S. Mitra et al. *Palaeocercospora* S. Mitra & M. Banerjee [2000](#page-148-2)



<span id="page-123-0"></span>**Fig. 58** *Olpidium brassicae* (=*Chytridium brassicae*) (Redrawn from Lay et al. ([2018\)](#page-144-3) who accepted the name *Olpidium brassicae* and they used Woronin ([1878\)](#page-160-5) to illustrate the taxon). **a**, **b**, **c** the zoosporangia of *Chytridium brassicae* in various developmental stages. **d** zoospores

of *C. brassicae*. **e**, **f**, **g** showed resting spores or spores of *C. brassicae*. **h** a cross section of a completely healthy root of a young cabbage seedling. **i**, **j**, **k** and **l** showed the root cells of cabbage

(Fig. [63-](#page-127-1)f), *Palaeoleptosphaeria* Barlinge & Paradkar 1982 (Fig. [63](#page-127-1)-g), *Palmellathyrites* Locq. et al., *Parmathyrites* K.P. Jain & R.C. Gupta [1970](#page-141-6) (Fig. [63](#page-127-1)-h), *Pelicothallos* Dilcher, *Perisporiacites* Félix *Phragmothyrites* W.N. Edwards [1922](#page-136-4) (Fig. [63-](#page-127-1)i), *Pleosporites* Y. Suzuki and *Torula* Pers., *Plochmopeltinites* Cookson, *Polyhyphaethyrites* R. Srivast. & R.K. Kar 2004 (Fig. [63-](#page-127-1)j), *Ratnagiriathyrites* R.K. Saxena & N.K. Misra [1990](#page-153-6) (Fig. [63-](#page-127-1)k), *Rhytidhysteriumites* Debi Mukh., *Spinosporonites* R.K. Saxena & S. Khare, *Sporidesmium* Link, *Stomiopeltites* Alvin & M.D. Muir, *Trichopeltina* Theiss., *Trichopeltinites* Cookson, *Trichothyrites* F. Rosend., *Ussurithyrites* Krassilov and *Vizella* Sacc.

### 1.2. Class: *Eurotiomycetes* O.E. Erikss. & Winka

This class is represented by the following genera: *Chaenothecopsis* Vain*.*, *Coleocarpon* Stubblef. et al., *Cryptocolax* R.A. Scott, *Memnonillasporonites* Debi Mukh., *Roannaisia* T.N. Taylor et al., *Sporocarpon* Will. and *Traquairia* Carruth.

### 1.3. Class: *Laboulbeniomycetes* Engl.

This class is represented by two genera, viz., Columnomyces R.K. Benj. and *Stigmatomyces* H. Karst.

### 1.4. Class: *Lecanoromycetes* Erikss. & Winka

This class is represented by *Sphaerophorus* Pers.

# 1.5. Class: *Leotiomycetes* O.E. Erikss. & Winka

This class is represented by the following genera: *Erisiphites* Pampal., *Lithouncinula* N. Sharma, et al., *Palaeosclerotium* G.W. Rothwell, *Perisporites* Pampal., *Protoerysiphe* N. Sharma et al. and *Trimmatostroma* Corda.

# 1.6. Class: *Pezizomycetes* O.E. Erikss. & Winka:

This class is represented by the following genera: *Ascodesmisites* Trivedi et al., *Paleomorchella* Poinar and *Pezizites* Göpp. & Berendt.

#### 1.7. Class: *Sordariomycetes* O.E. Erikss. & Winka:



**Fig. 59** The representatives of *Rozellomycota*. **A** *Rozella allomycis* parasitizing the blastocladiomycete *Allomyces* sp. (light microscopic photo by Timothy James CC BY-SA 3.0, [htps://commons.wikimedia.](htps://commons.wikimedia.org/w/index.php?curid=8727738) [org/w/index.php?curid=8727738\)](htps://commons.wikimedia.org/w/index.php?curid=8727738). **B** *Chytridiopsis trichopterae* from the caddisfy *Polycentropus favomaculatus*, mature spores in parasitophorous vacuole (PV), arrowheads indicate remnants of plasmodium (transmission electron microscopic photo reprinted from Larsson (1993) under license 5622170021648 granted by John Wiley & Sons). **C** *Metchnikovella incurvata*, a hyperparasite of gregarines *Polyrhabdina pygospionis* from the *polychaete Pygospio* elegans.

<span id="page-124-0"></span>Class *Sordariomycetes* constitutes one of the most abundant classes of fossil *Ascomycota*. This class is represented by the following genera: *Acremonium* Link, *Botryodiplodia* Sacc., *Brachysporisporites* Lange & Smith [1971](#page-144-4) (Fig. [63-](#page-127-1)l) and *Brachysporium* Sacc., *Chaethomites* Pampal., *Chaetosphaeria* Tul. & C. Tul., *Clasterosporium* Schwein., *Diploneurospora* K.P. Jain & R.C. Gupta [1970](#page-141-6) (Fig. [63](#page-127-1)-m), *Hypoxylonites* Elsik [1990](#page-136-5) (Fig. [63-](#page-127-1)n), *Melanospora* Corda, *Meliola* Fr., *Meliolinites* Selkirk [1975](#page-154-4) (Fig. [63](#page-127-1)-o) and *Meliostroma* R. Kar et al., *Palaeoamphisphaerella* Ramanujam & Srisailam (Fig. [63-](#page-127-1)p), *Palaeocirrenalia* Ramanujam & Srisailam [1980](#page-151-6) (Fig. [63](#page-127-1)-q), *Palaeoclaviceps* Poinar et al., *Palaeocolletotrichum* S. Mitra & M. Banerjee [2000](#page-148-2) (Fig. [63](#page-127-1) r), *Palaeosordaria* Sahni & H.S. Rao [1943](#page-152-1) (Fig. [63-](#page-127-1)s), *Paleoophiocordyceps* G.H. Sung et al., *Paleoserenomyces* Currah et al., *Petrosphaeria* Stopes & H. Fujii, *Spataporthe* Bronson et al., *Sphaerites* Unger, *Valsarites* Puri and *Zopfella* G. Winter.

1.8. Class: *Taphrinomycetes* O.E. Erikss. & Winka

Infected gregarine cell flled with the spore sacs (cyst) and free spores (fsp) (light microscopic photo reprinted from Galindo et al. 2018 CC B Y 4 .0). **d** Mature spore of *Metchnikovella incurvate* (transmission electron microscopic photo reproduced with a kind permission of Yuliya Y. Sokolova, Institute of Cytology RAS, St. Petersburg, Russia). **e** Mature spore of *Crepidulospora beklemishevi* from the mosquito *Anopheles beklemishevi* (transmission electron microscopic photo reproduced w ith a k ind p ermission o f Anastasia V. S imakova, Tomsk S tate University, Tomsk, Russia). Scale bars:  $\mathbf{b} = 1 \, \mu \text{m}$ ;  $c = 5 \mu m$ ; **d** = 0.5  $\mu m$ .

This class is represented by a single genus, viz., *Protomyces* Unger.

### 2. Phylum: *Basidiomycota* R.T. Moore

Basidiomycota are flamentous fungi composed of hyphae (except for basidiomycota-yeast) and reproduce sexually via the formation of specialized club-shaped end cells called basidia. They normally bear external meiospores (usually four). These specialized spores are called basidiospores. However, some *Basidiomycota* are obligate asexual reproducers. *Basidiomycota* that reproduces asexually can typically be recognized as members of this division by gross similarity to others, by the formation of a distinctive anatomical feature (the clamp connection) and cell wall components. The following fossil genera exhibit affinity with *Basidiomycota*, but their resemblance with taxa of lower ranks could not be established: *Lycoperdites* Poinar and *Mycetophagites* Poinar & Buckley. Fossil basidiomycetus fungi are represented by the following four classes: *Agaricomycetes* Doweld, *Agaricostilbomycetes* R. Bauer et al.,

<span id="page-125-0"></span>**Fig. 60** Light microscopic images of *Sanchytrium tribonematis.* **a**, **b** amoeboid zoospores with flopodia (f) and posterior pseudocilium (pc). **c** sporangia (sp) on the *Tribonema* flament. **d** rhizoid (arrow) of sporangium. **e** zoospores (zo) releasing sporangium through the discharge pore (arrow). Scale bars: **a**, **b**=3.5  $\mu$ m, **c**–**e**=10  $\mu$ m



*Pucciniomycetes* R. Bauer et al. and *Ustilaginomycetes* R. Bauer et al.

## 2.1. Class: *Agaricomycetes* Doweld

This class is represented by the following genera: *Archaeomarasmius* Hibbett et al., *Coprinites* Poinar & Singer, *Cyathus* Haller, *Eopolyporoides* Rigby in G. Playford et al., *Fomes* (Fr.) Fr., *Ganodermites* A. Fleischm. et al., *Geastrum* Pers., *Gerontomyces* Poinar, *Gondwanagaricites* Heads et al., *Hydnum* L., *Nidula* V.S. White, *Lithopolyporales* R. Kar et al. [2003](#page-142-7) (Fig. [63](#page-127-1)-t), *Palaeoagaricites* Poinar & Buckley, *Palaeoclavaria* Poinar & A.E. Br., *Palaeogaster* Poinar et al., *Phellinites* Singer & S. Archang., *Protomycena* Hibbett et al., *Pseudopolyporus* Hollick, *Scleroderma* Pers. and *Trametites* Mesch.

### 2.2. Class: *Agaricostilbomycetes* R. Bauer et al.

This class is represented by a single genus, viz., *Stilbum* Tode.

## 2.3. Class: *Pucciniomycetes* R. Bauer et al.

This class is represented by the following genera: *Aecidites* Debey & Ettingsh., *Aeciosporonites* Debi Mukh. *Hapalophragmites* Ramanujam & Ramachar [1980](#page-151-7) (Fig. [63](#page-127-1)-u), *Milesites* Ramanujam & Ramachar [1980](#page-151-7) (Fig. [63-](#page-127-1)v), *Pucciniasporonites* Ramanujam & Ramachar and *Shuklania* J.N. Dwivedi.

### 2.4. Class: *Ustilaginomycetes* R. Bauer et al.

This class is represented by the following genera: *Chlamydosporites* Paradkar 1975 (Fig. [63](#page-127-1)-w), *Mundkurella* Thirum., *Teliosporites* R. Kar et al. [2010](#page-142-8) (Fig. [63](#page-127-1)-x), *Thecaphora* Fingerh. and *Ustilago* (Pers.) Roussel.

#### 3. Phylum: *Blastocladiomycota* T.Y. James

*Retesporangicus* Strullu-Derr. is assignable to phylum *Blastocladiomycota*.

#### 4. Phylum: *Chytridiomycota* Doweld

The following genera have affinity with phylum *Chytridiomycota* but their resemblance with taxa of lower ranks could not be established: *Brijax* M. Krings & C.J. Harper, *Globicultrix* M. Krings et al., *Illmanomyces* M. Krings & T.N. Taylor, *Nimbosphaera* C.J. Harper & M. Krings, *Palaeozoosporites* Strullu-Derr., *Perexifasca* M. Krings et al., *Rhizophydites* M. Krings et al. Fossil chytridiomycetus fungi are represented by only one class, viz., *Chytridiomycetes* M. Möbius.

#### 4.1. Class: *Chytridiomycetes* M. Mӧbius

This class is represented by the following genera: *Cultoraquaticus* Strullu-Derr., *Entophlyctis* A. Fisch., *Grilletia* Renault & C.E. Bertrand, *Guizhounema* X. Mu, *Krispiromyces* T.N. Taylor et al., *Lyonomyces* T.N. Taylor et al., <span id="page-126-0"></span>**Fig. 61** *Zoopage phanera* Drechsler (redrawn from Drechsler [1935](#page-136-6)). **A** Four specimens: **a**, **b**, **c** and **d** of *Amocba terricola* III captured by the branched hyphae; **e** and **h** of *Zoopage phancra*, showing the stalked botryoid haustoria of the fungus; hyphae **f** and **g**, perhaps also adhering to animal but without having produced haustoria, bear respectively one and two erect conidiiferous branches with long chains of conidia of which from lack of space only the lowermost individuals are shown; in the animal, d is shown its nucleus, and the same structure from a healthy specimen is shown in  $n; \times 500$ . **B** Two hyphae, bearing three conidial chains: **a**–**c** (shown only inpart from lack of space) and producing on diclinous sexual branches two zygosporangia, d and c, shown at early stages of development;  $\times$  500. **C**–**H** Sexual apparatus of *Zoopage phancra* showing diclinous origin of zygophore, inconstancy of septation during earlier stages in development of fusion product, and the frequently contorted condition of one of the zygophores;  $\times 1000$ . **I**–**M** Approximately mature zygospores, each within its collapsing zygosporangial membrane;×1000. **N**, **O** Sexual apparatus, with a germ tube from a conidium functioning directly as a zygophore;  $\times 1000$ 



*Milleromyces* T.N. Taylor et al., *Oochytrium* Renault, *Trewinomyces* M. Krings et al.

# 5. Phylum: *Glomeromycota* C. Walker & A. Schüßler

The following fossil genera exhibit affinity with *Glomeromycota*, but their resemblance with taxa of lower ranks could not be established: *Helmutella* M. Krings & T.N. Taylor, *Kryphiomyces* M. Krings et al., *Palaeomyces* D. Ellis and *Zwergimyces* M. Krings & T.N. Taylor. Fossil glomeromycetus fungi are represented by following two classes, viz., *Archaeosporomycetes* Sieverd et al. and *Glomeromycetes* Caval.-Sm.

5.1. Class: *Archaeosporomycetes* Sieverd et al.

This class is represented by *Archaeosporites* C. Walker et al.

#### 5.2. Class: *Glomeromycetes* Caval.-Sm.

This class is represented by the following genera: *Gigasporites* Carlie J. Phipps & T.N. Taylor, *Glomites* T.N. Taylor et al., *Glomorphites* Garcia Mass., *Palaeogigaspora* R. Kar et al. [2010](#page-142-8) (Fig. [63](#page-127-1)**-y**), *Palaeoglomus* R. Redecker et al. and Scutellosporites Dotzler et al.

#### 6. Phylum: *Mucoromycota* Doweld



<span id="page-127-0"></span>**Fig. 62** *Verrucocephalum latericorvinisporum* (KPM-NC0016331). **a** Sporophores with matured black spores hanging on short side branchlets. One of the spores accompanying a water drop indicated by a dotted line (arrow). **b** The bottom view of a young hyaline spore with initial minute verrucae. **c** The bottom view of a maturing spore with darkly pigmented verrucae. **d** The top view of a maturing black-

Stolophorites Wilh. Bock exhibits affinity with *Mucoromycota*, but its resemblance with taxa of lower ranks could not be established. Fossil mucoromycetus fungi are represented by a single class, viz., *Endogonomycetes* Doweld.

#### 6.1. Class: *Endogonomycetes* Doweld

This class is represented by the following genera: *Chlamydospora* R. Kar et al., *Endochaetophora* J.F. White & T.N. Taylor (Doweld [2014e\)](#page-135-4), *Lithomucorites* R. Kar et al. [2010](#page-142-8) (Fig. [63](#page-127-1)-z)., *Palaeomycites* Mesch. and *Udaria* A. Gupta 1996 (Fig. [63-](#page-127-1)aa).

ish spore with verrucose surface. **e** Whole fgure of a trapped nematode with elongating haustorial assimilative hyphae from appressoria in a cavity of the nematode. Arrows indicate penetration parts with appressoria and post-penetration bulbs. Scale  $bar = a - e$  50  $\mu$ m (redrawn from Degawa [2013\)](#page-135-3)

<span id="page-127-1"></span>**Fig. 63** Diferent morphologies of fossil spores. **a** *Appendicisporon-*◂*ites typicus*. **b** *Callimothallus pertusus*. **c** *Cucurbitariaceites bellus*. **d** *Kalviwadithyrites saxenae*. **e** *Koshalia enigmata*. **f** *Palaeocercospora siwalikensis*. **g** *Palaeoleptosphaeria intertrappeana*. **h** *Parmathyrites indicus*. **i** *Phragmothyrites eocaenicus*. **j** *Polyhyphaethyrites giganticus*. **k** *Ratnagiriathyrites hexagonalis*. **l** *Brachysporisporites pyriformis*. **m** *Diploneurospora tewarii*. **n** *Hypoxylonites brazosensis*. **o** *Meliolinites spinksii*. p *Palaeoamphisphaerella pirozynskii*. **q** *Palaeocirrenalia elegans*. **r** *Palaeocolletotrichum graminioides*. **s** *Palaeosordaria lagena*. **t** *Lithopolyporales zeerabadensis*. **u** *Hapalophragmites cumminsii*. **v** *Milesites irregularis*. **w** *Chlamydosporites gramineus* **a** Reticulate chlamydospore; **b**, **c** Mycelium and hyphae with haustoria and chlamydospores. **x** *Teliosporites globatus*. **y** *Palaeogigaspora excellensa*. **z** *Lithomucorites miocenicus*. **aa** *Udaria singhii*. Scale bars: **a**–**c**, **h**, **k**, **w** 50 μm, **d**, **g**, **r**, **t** 25 μm, **e**, **f**, **l**, **m**, **q**, **s**, **u** 20 μm, **i** 60 μm, **j** 500 μm, **n**–**p**, **v**, **x**–**z** 10 μm, aa 30 μm



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### **Declarations**

**Conflict of interest** The authors declare no confict of interest. The author list includes members of the Editorial Board of Fungal Diversity. They were not involved in the journal's review of, or decisions related to, this manuscript.

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