



Phylogeny and character evolution in the *Dacrymycetes*, and systematics of *Unilacrymaceae* and *Dacryonaemataceae* fam. nov.

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Key words

Carotenoids
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Dacryonaema
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Abstract We present a multilocus phylogeny of the class *Dacrymycetes*, based on data from the 18S, ITS, 28S, *RPB1*, *RPB2*, *TEF-1 α* , 12S, and *ATP6* DNA regions, with c. 90 species including the types of most currently accepted genera. A variety of methodological approaches was used to infer phylogenetic relationships among the *Dacrymycetes*, from a supermatrix strategy using maximum likelihood and Bayesian inference on a concatenated dataset, to coalescence-based calculations, such as quartet-based summary methods of independent single-locus trees, and Bayesian integration of single-locus trees into a species tree under the multispecies coalescent. We evaluate for the first time the taxonomic usefulness of some cytological phenotypic characters, i.e., vacuolar contents (vacuolar bodies and lipid bodies), number of nuclei of recently discharged basidiospores, and pigments, with especial emphasis on carotenoids. These characters, along with several others traditionally used for the taxonomy of this group (basidium shape, presence and morphology of clamp connections, morphology of the terminal cells of cortical/marginal hyphae, presence and degree of ramification of the hyphidia), are mapped on the resulting phylogenies and their evolution through the class *Dacrymycetes* discussed. Our analyses reveal five lineages that putatively represent five different families, four of which are accepted and named. Three out of these four lineages correspond to previously circumscribed and published families (*Cerinomycetaceae*, *Dacrymycetaceae*, and *Unilacrymaceae*), and one is proposed as the new family *Dacryonaemataceae*. Provisionally, only a single order, *Dacrymycetales*, is accepted within the class. Furthermore, the systematics of the two smallest families, *Dacryonaemataceae* and *Unilacrymaceae*, are investigated to the species level, using coalescence-based species delimitation on multilocus DNA data, and a detailed morphological study including morphometric analyses of the basidiospores. Three species are accepted in *Dacryonaema*, the type, *Da. rufum*, the newly combined *Da. macnabbii* (basionym *Dacrymyces macnabbii*), and a new species named *Da. macrosporum*. Two species are accepted in *Unilacryma*, the new *U. bispora*, and the type, *U. unispora*, the latter treated in a broad sense pending improved sampling across the Holarctic.

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INTRODUCTION

The *Dacrymycetes* (*Basidiomycota*) is one of the three widely accepted classes in the *Agaricomycotina*, sister to the *Agaricomycetes*. It is a monophyletic group comprised of widespread, saprotrophic fungi, mainly characterised by the strikingly long bisterigmate (forked) holobasidia (Hibbett et al. 2007), *Unilacryma unispora* being the only known exception (Shirouzu et al. 2013). The class is further characterised at the ultrastructural level by the presence of continuous parenthosomes covering the dolipore-type septa (Oberwinkler 1994, Wells 1994). In addition, basidiomata are gelatinous to waxy or fleshy-cartilaginous, displaying a wide variation in size and shape, and very frequently yellowish to orange due to the presence of carotenoids (Gill & Steglich 1987), the main pigments of this group (Fig. 1).

The *Dacrymycetes* are typically brown-rotters (Worrall et al. 1997), although some have been reported to degrade considerable amounts of lignin (Seifert 1983). Gene contents from two *Calocera* genomes also suggests a brown-rotting lifestyle (Nagy et al. 2015), and *Dacrymycetes* are the clade with the oldest stem age among *Basidiomycota* wood decomposers. As a result, their ancestors were likely among the first basidio-

mycetous fungi able to decay wood (Oberwinkler 1994, Kirk et al. 2008, Floudas et al. 2012). Floudas et al. (2012), based on genome-scale data (26 DNA regions) in which *Dacrymycetes* was represented by a single species, estimated the stem age of the class to around 350 million years ago (Ma). A similar result, based on phylogenetic analysis of up to six DNA regions, was reached by Zhao et al. (2017), who reported a mean stem age of 358 Ma. Garnica et al. (2016) estimated the stem age to around 300 Ma using four DNA regions, but the *Dacrymycetes* clade was represented by only two species in the *Dacrymycetaceae*. Recently, Varga et al. (2019) performed a variety of dating analyses on a phylogenomic dataset including three *Dacrymycetaceae* species and obtained mean stem ages for the *Dacrymycetes* around 278–504 Ma, although mostly in the 360–385 Ma range. In view of this, the ancestors of this class and the *Agaricomycetes* seem to have split during the Upper Devonian or the Carboniferous.

The *Dacrymycetes* was traditionally comprised of a single order, *Dacrymycetales*, with two families, *Dacrymycetaceae* and *Cerinomycetaceae*, but a second order called *Unilacrymales* was erected to accommodate the single known species with simple basidia, *Unilacryma unispora* (Shirouzu et al. 2013). The two orders were accepted in a recent study dealing with high-level taxonomy of the *Basidiomycota* (Zhao et al. 2017), although no species of *Cerinomycetaceae* were included.

Generic boundaries within the *Dacrymycetes* have largely remained unaltered since Martin (1952), except for *Dicellomyces* having been transferred to the *Ustilaginomycotina* (Kirk et al.

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Fig. 1 Morphological variation of the fruitbodies in the *Dacrymycetaceae*. a. Resupinate and almost unpigmented in *Cerinomyces* sp. (O 247959); b. sterile dark conical stalks in *Dacryonaema rufum* (UPS F-941005); c. clavarioid and yellow in *Calocera comea* (UPS F-940775); d. turbinate, stalked and with a yellow-orange hymenium in *Ditiola radicata* (UPS F-939961); e. pulvinate, sessile and brownish in *Unilacryma bispora* (UPS F-941266); f. pulvinate, sessile and bright orange in *Dacrymyces ovisporus* (UPS F-940139). All in fresh conditions except a. — Scale bars = 1 mm.

2008). The shape of the fruitbodies, basidioma growth direction, presence or absence of a stipe, wall thickness of internal hyphae, and cell differentiation of cortical/marginal hyphae (partially related to the presence/absence of a sterile cortex) are among the most important taxonomic characters traditionally used to distinguish the genera (Fig. 1–2). In a revision based mainly on these characters, Oberwinkler (2014) accepted the following 13 genera: *Arrhytidia*, *Calocera*, *Cerinomyces*, *Dacrymyces*, *Dacryomitra*, *Dacryonaema*, *Dacryopinax*, *Dacryoscyphus*, *Ditiola*, *Femsjonina*, *Guepiniopsis*, *Heterotextus*, and *Unilacryma*. However, some of the mentioned characters, e.g., the shape of the fruitbodies, were demonstrated to be homo-

plastic by Shirouzu et al. (2013), and several common genera are polyphyletic (e.g., *Calocera*, *Cerinomyces*, *Dacrymyces*). In the same study, it was shown that other phenotypic traits, such as the presence or absence of clamp connections, not previously used to distinguish genera, are much better correlated with the phylogeny based on DNA data. For instance, species with and without clamp connections in *Calocera*, or in *Cerinomyces*, are not closely related.

Carotenoid pigments typical of the *Dacrymycetaceae* are synthesised by other fungal groups, but they are relatively unusual among non-photosynthetic eukaryotes (Echavarrri-Erasun & Johnson 2002). They are prominent only in a few fungal taxa,

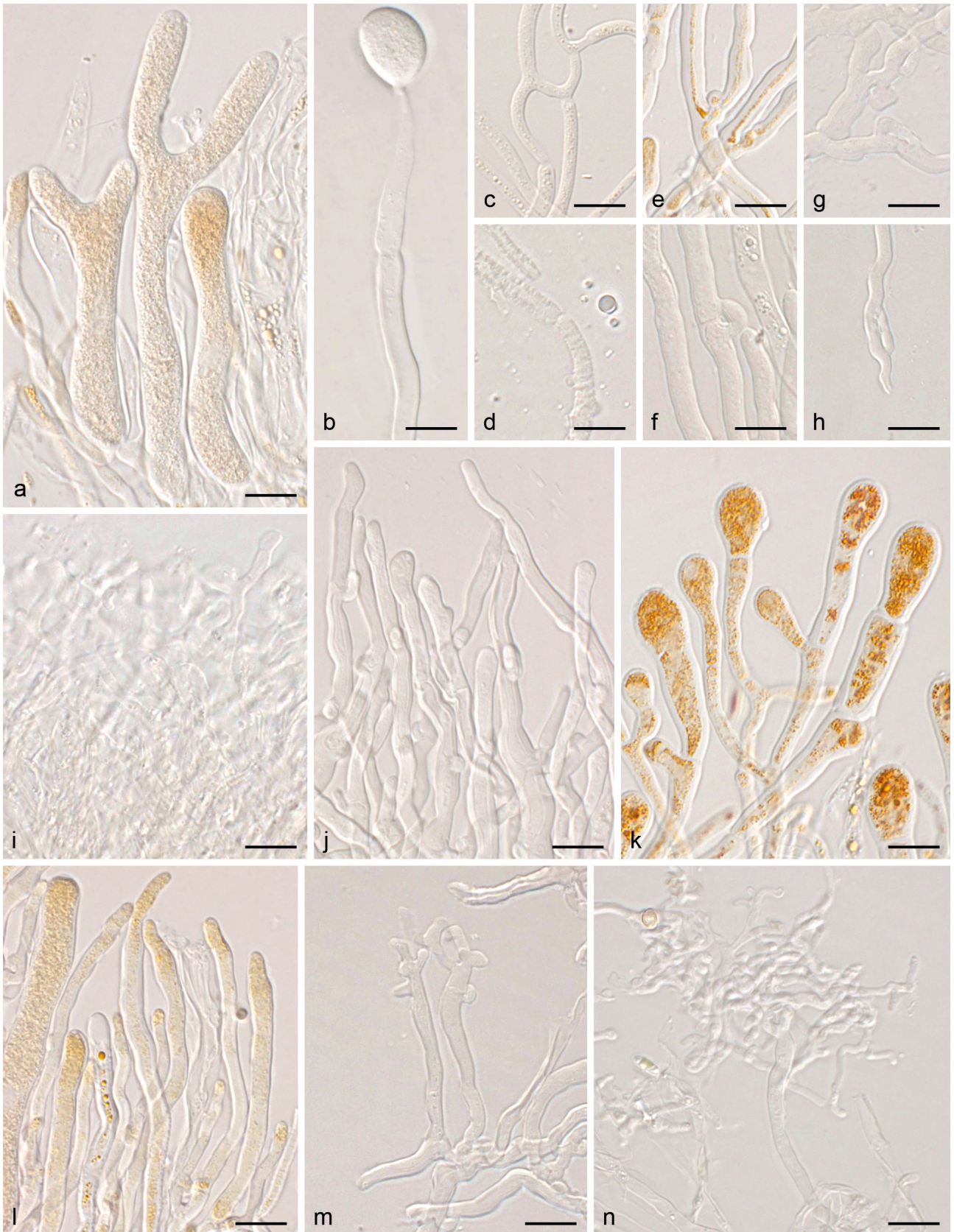


Fig. 2 Micromorphology of traditionally used taxonomic characters. a. Bisterigmate basidia in *Dacrymyces estonicus* (UPS F-941253); b. unisterigmate basidium in *Unilacryma unispora* (UPS F-941283); c–d. clampless septa in *D. stillatus* (c, UPS F-939816) and *Dacryonaema macrosporum* (d, holotype); e–f. short and rounded clamp connections in *D. chrysocomus* s.lat. 2 (e, UPS F-940946) and *U. bispora* (f, UPS F-941259); g–h. long and comparatively angulose clamp connections in *Da. macnabbii* (g, UPS F-940954) and *Da. rufum* (h, UPS F-941005); i–k. terminal cells of cortical/marginal hyphae, branched and gelatinised in *Da. macnabbii* (i, UPS F-940966), simple and not swollen in *D. tortus* s.lat. 4 (j, UPS F-941252), and clearly dilated in *D. estonicus* (k, UPS F-941253); l–n. hyphidia, simple in *D. ovisporus* (l, UPS F-940139), with short apical branches in *D. tortus* s.lat. 3 (m, UPS F-941018), and heavily branched in *Da. macnabbii* (n, UPS F-940950). — Scale bars = 10 μ m.

and as a result they have been used as taxonomic characters both in *Ascomycota* (e.g., Eckblad 1968, Schumacher 1990, Baral 2004) and *Basidiomycota* (e.g., Arpin & Fiasson 1971, Olariaga et al. 2015). Their acquisition and secondary loss in different phylogenetic lineages are well-known (Hansen et al. 2013, Olariaga et al. 2015), but remains to be investigated in the *Dacrymycetetes*.

Species delimitations are challenging in the *Dacrymycetetes*. According to Hibbett et al. (2007), this is the smallest class of the subphylum *Agaricomycotina*, with 110–120 species known so far (Kirk et al. 2008, Shirouzu et al. 2009, 2017). However, judging from recent studies and our experience with fresh and herbarium samples, the real number of species is certainly larger, probably over 200. Basidiospore dimensions and septation, and the morphology of hyphidia are important characters for morphological species delimitation (McNabb 1964, 1965a–e, 1966, 1973), but their variation may be considerable even within a single fruitbody. Shirouzu et al. (2017) provided the most complete phylogeny in terms of included vouchers with both DNA data and morphological data from fruitbodies. Some of the tentatively identified species turned out to be polyphyletic (e.g., *Calocera* cf. *guepinioides*, *Cerinomyces crustulinus*, *Ce. pallidus*, *Dacrymyces microsporus*, *D. minor*, *D. novae-zelandiae*, *D. stillatus*) or appeared on exceptionally long branches within their clade, indicating there is considerable sequence variation (e.g., *Calocera cornea*, *Ca. lutea*, *Dacrymyces capitatus*, *Heterotextus miltinus*). These groups, among others, may contain so far unrecognised species.

The class was subject to a worldwide monograph by Robert McNabb, who described new taxa from a variety of countries and ecozones in a series of studies (McNabb 1964, 1965a–e, 1966, 1973) based on numerous other previous treatments, including those of, e.g., Brasfield (1938) and Kennedy (1959a, b) from the USA, or Kobayasi (1939a, b) from East Asia. During the last decade, significant contributions to the understanding of this group, including several studies based on morphological, molecular and ecological data, mainly on the Japanese and Australasian species, have been published by, e.g., Shirouzu et al. (2009, 2013, 2016, 2017). In Europe, this group sparked the interest of Neuhoﬀ (1936), and was later addressed in considerable detail by Donk (1966) and Raitviir (1967, including a substantial part of northern Asia). Reid (1974) presented a monograph focused on the British taxa, including 20 species, two of which were proposed as new. Since then, however, there have been no thorough revision of the European taxa with modern techniques. In the Nordic countries, Nannfeldt's (1947) monotypic genus *Dacryonaema* stands out as infrequently recorded, mainly from Northern Europe (Torkelsen 1997, <http://www.gbif.org/species/2512811>), although it has also been reported from North America (Brough & Bandoni 1975, Redhead 1997). No DNA data were available for *Dacryonaema* at the beginning of this study, and the phylogenetic relationships of this morphologically deviant genus with the remaining species of *Dacrymycetetes* are largely unknown (Oberwinkler 2014). On the other hand, the monotypic genus *Unilacryma*, described from North America and recorded from Japan, has recently been found also in Europe (Dam & Dam 2014). Shirouzu et al. (2013) commented on the morphological characteristics of the only known species, *U. unispora*, and compared it with another morphologically close species, *Dacrymyces ovisporus*, based on basidia and basidiospore characters. *Dacrymyces ovisporus* was described from Germany and has a wide distribution, including, e.g., northern Europe (Torkelsen 1997) as well as North America (Bandoni 1963). No DNA sequence data of this species are available yet, and its taxonomic position remains unclear.

The objectives of this paper are:

- (i) to establish phylogenetic relationships among genera and species in the *Dacrymycetetes* (including the genus *Dacryonaema*) based on multilocus DNA data;
- (ii) to identify phenotypic characters that allow the recognition of the phylogenetic lineages at family level, paying particular attention to those previously overlooked; and
- (iii) to revise the systematics and nomenclature of *Dacryonaemataceae* and *Unilacrymaceae* based on an integrative taxonomy approach, with special emphasis on the European taxa.

The third aim was taxonomically and geographically restricted because a wider scope would have been hampered by the extensive diversity and the large number of published names. The systematics of the larger *Cerinomycetaceae* and *Dacrymycetaceae* will be investigated in collaboration with other researchers in forthcoming studies.

MATERIAL AND METHODS

Taxon and marker sampling

We sampled representatives for an overall phylogeny based on their morphology and phylogenetic position in previous studies. Due to the similarity of names, *Calocera* is abbreviated as 'Ca.', *Cerinomyces* as 'Ce.', *Dacrymyces* as 'D.', *Dacryonaema* as 'Da.', *Ditiola* as 'Di.', whereas *Dacryomitra*, *Dacryopinax* and *Dacryoscyphus* are not abbreviated. We generated new data from five species in the *Cerinomycetaceae*, viz., *Cerinomyces* sp. (resupinate morphology and ± dry fruitbodies) and *D. tortus* s.lat. (four species with 'dacrymycetoid' and 'coalescing' morphologies, with ± gelatinous fruitbodies); the following species in the *Dacrymycetaceae*, viz., *Ca. cornea*, *Ca. viscosa*, *D. chrysocomus* s.lat., *D. aff. chrysospermus*, *D. estonicus*, *D. cf. minutus*, *D. pinacearum*, *D. stillatus*, *Dacryomitra pusilla*, *Dacryopinax elegans*, *Ditiola radicata*, *Femsjonina peziziformis* s.lat., *Guepiniopsis buccina*, and *Heterotextus miltinus* s.lat.; *Unilacryma unispora* s.lat. in the *Unilacrymaceae*, and *Dacryonaema rufum* with an unknown placement. Additionally, we included *Dacrymyces macnabbii* and *D. ovisporus*, which have not been included in any phylogeny yet, and two unidentified species that, according to morphological and preliminary molecular data, may belong in *Dacryonaema* or *Unilacryma*. We paid particular attention to selecting only visibly clean and non-parasitised fruitbodies. In addition, we added all available *Dacrymycetetes* samples in GenBank being represented by at least three of the eight DNA regions selected (see below), for this purpose considering the ITS (ITS1 + 5.8S + ITS2) as a single region (Table 1). This includes mostly samples from the AFTOL project (Hibbett et al. 2007), the studies on the *Dacrymycetetes* by Shirouzu et al. (2007, 2009, 2013, 2016), and four *Dacrymycetetes* species in MycoCosm (Grigoriev et al. 2014). In order to better assess the consistency of some phenotypic characters at the family level, we expanded the taxonomic sampling to include most available species with both a reasonable amount of molecular and morphological data. For samples with at least two DNA regions available in GenBank, we included data of up to two individuals of each taxon identified to species level when it was either not previously included or represented by a single individual. When selecting among multiple available sequences, preference was given first to type specimens and second to the longest sequences; most of these additional sequences were generated by Shirouzu et al. (2017). Finally, and as an exception, a partial 28S sequence of *Dacryoscyphus chrysochilus* (AY604567, from the holotype specimen) was included because this was the only available DNA sequence of the type of *Dacryoscyphus*.

Table 1 GenBank accession numbers for *Dacrymycetes* sequences used in the phylogenetic analyses (except those from the four genomes indicated in Table 2). Taxon names refer to our post-hoc knowledge and are frequently provisional pending specific studies on certain groups. Environmental nucleic acid sequences without vouchers are indicated as 'ENAS', and are tentatively assigned to a species name based on sequence similarity only. New sequences generated for this study are marked in **bold**. *Holotype specimen. Unpub. = Unpublished DNA sequence from the AFTOL database.

Taxon	Herbarium voucher or isolate	Country	GenBank accession numbers									
			18S	ITS	28S	TEF-1 α	RPB1	RPB2	12S	ATP6		
Ceratomyces												
<i>Ceratomyces albosporus</i> s.lat. (including <i>C. aculeatus</i>)	TNS-F-15706, TUFC 12991, MAFF 240121	Japan	AB712482	AB712440	AB299050	—	—	—	—	—	—	—
<i>Ceratomyces cf. canadensis</i>	TNS-F-21034, TUFC 12876, MAFF 241162	Japan	AB712483	AB712441	AB472696	—	—	—	—	—	—	—
<i>Ceratomyces cf. crustulinus</i>	TUFC30545	Canada	AB712485	AB712443	AB712423	—	—	—	—	—	—	—
<i>Ceratomyces cf. pallidus</i>	FP-150848	Belize	AB712488	AB712446	AB712426	—	—	—	—	—	—	—
<i>Ceratomyces</i> sp. (resupinate)	O 247959	Norway	MN593435	MN595619	MN595619	MN580318	MN580218	MN580218	MN580258	MN580258	MN593502	MN567220
<i>Dacrymyces</i> sp. (as <i>D. punctiformis</i>)	TNS-F-15723, TUFC 13053, MAFF 240138	Japan	AB712504	AB712462	AB299052	—	—	—	—	—	—	—
<i>Dacrymyces tortus</i> s.lat. 1	UPS F-940777	Sweden	MN593436	MN595620	MN595620	MN580319	MN580219	MN580219	MN580259	MN580259	MN593503	MN567221
<i>Dacrymyces tortus</i> s.lat. 2	UPS F-940948	Sweden	MN593437	MN595621	MN595621	MN580320	MN580220	MN580220	MN580260	MN580260	MN593503	MN567221
<i>Dacrymyces tortus</i> s.lat. 3	UPS F-941016	Sweden	MN593438	MN595622	MN595622	MN580321	MN580221	MN580221	MN580261	MN580261	MN593504	MN567221
<i>Dacrymyces tortus</i> s.lat. 3	UPS F-941017	Sweden	MN593439	MN595623	MN595623	MN580322	MN580222	MN580222	MN580262	MN580262	MN593505	MN567221
<i>Dacrymyces tortus</i> s.lat. 4	UPS F-941019	Spain	MN593440	MN595624	MN595624	MN580323	MN580223	MN580223	MN580263	MN580263	MN593505	MN567221
<i>Dacrymyces tortus</i> s.lat. 4	UPS F-941020	Sweden	MN593441	MN595625	MN595625	MN580324	MN580224	MN580224	MN580264	MN580264	MN593505	MN567221
Unidentified <i>Ceratomyces</i>	NBRC 110591	Japan	LC004021	LC004000	LC003883	—	—	—	—	—	—	—
Dacrymycetaceae												
<i>Calocera arborea</i>	*INPA 241458	Brazil	—	AB744230	AB723514	—	—	—	—	—	—	—
<i>Calocera cornea</i>	CBS124.84	Canada	AB712479	AB472738	AB472738	—	—	—	—	—	—	—
	UPS F-940774	Sweden	MN593442	MN595626	MN595626	MN580325	MN580225	MN580225	MN580265	MN580265	MN593506	MN567221
	UPS F-940775	Sweden	MN593443	MN595627	MN595627	MN580326	MN580226	MN580226	MN580266	MN580266	MN593507	MN567221
	GEL 5359, AFTOL-ID 438	Sweden	AY771610	AY789083	AY701526	AY881019	AY857980	AY857980	AY536286	AY536286	MN593507	MN567221
<i>Calocera fusca</i>	PDD 107930	New Zealand	—	LC131405	LC131364	—	—	—	—	—	—	—
	New Zealand	New Zealand	—	LC131406	LC131365	—	—	—	—	—	—	—
	PDD 107972, ICMP 21238	New Zealand	—	LC131411	LC131370	—	—	—	—	—	—	—
<i>Calocera guepinioides</i> s.lat. 1	PDD 107969, ICMP 21236	New Zealand	—	LC131412	LC131371	—	—	—	—	—	—	—
	PDD 107981, ICMP 21240	New Zealand	—	LC131407	LC131366	—	—	—	—	—	—	—
<i>Calocera guepinioides</i> s.lat. 2	PDD 105005, ICMP 20480	New Zealand	—	LC131409	LC131368	—	—	—	—	—	—	—
	PDD 107874, ICMP 21226	New Zealand	—	LC131408	LC131367	—	—	—	—	—	—	—
<i>Calocera guepinioides</i> s.lat. 3	PDD 105033, ICMP 20502	New Zealand	—	LC131410	LC131369	—	—	—	—	—	—	—
	PDD 107929, ICMP 21231	New Zealand	—	LC131411	LC131370	—	—	—	—	—	—	—
<i>Calocera lutea</i> s.lat. 1	CBS291.82	Australia	AB712480	AB712438	AB712379	—	—	—	—	—	—	—
<i>Calocera lutea</i> s.lat. 2	PDD 107841, ICMP 21221	New Zealand	—	LC131413	LC131372	—	—	—	—	—	—	—
	PDD 107842, ICMP 21221	New Zealand	—	LC131414	LC131373	—	—	—	—	—	—	—
	PDD 107830	New Zealand	—	LC131415	LC131374	—	—	—	—	—	—	—
<i>Calocera pedicellata</i>	PDD 107925, ICMP 21230	New Zealand	—	LC131416	LC131375	—	—	—	—	—	—	—
<i>Calocera viscosa</i> s.lat.	TNS-F-15704, TUFC 12873, MAFF 240119	Japan	AB712481	AB712439	AB299048	—	—	—	—	—	—	—
	UPS F-940773	Sweden	MN593444	MN595628	MN595628	MN580327	MN580227	MN580227	MN580267	MN580267	MN593508	MN567222
	TUB 020317, MW591, AFTOL-ID 1679	Sweden	DQ520102	DQ520102	DQ520102	Unpub.	FJ623634	FJ623634	FJ623649	FJ623649	MN593508	MN567222
	HHB-8969	USA	AB712484	AB712442	AB712422	—	—	—	—	—	—	—
	HHB-6908	USA	AB712486	AB712444	AB712424	—	—	—	—	—	—	—
<i>Ceratomyces ceraceus</i>	RLG-13487	USA	AB712487	AB712445	AB712425	—	—	—	—	—	—	—
<i>Ceratomyces cf. grandinoides</i>	TNS-F-21045, TUFC 12845, MAFF 241172	Japan	AB712489	AB712447	AB472707	—	—	—	—	—	—	—
<i>Ceratomyces lagerheimii</i>	*TNS-F-21051, MAFF241177	Japan	AB712490	AB712448	AB472713	—	—	—	—	—	—	—
<i>Dacrymyces cf. adpressus</i>	CBS293.82	Japan	AB712492	AB712450	AB472741	—	—	—	—	—	—	—
<i>Dacrymyces ancylius</i>	CBS280.84	Canada	AB712493	AB712451	AB712427	—	—	—	—	—	—	—
<i>Dacrymyces capitatus</i> s.lat.	UPS F-940136	Spain	MN593445	MN595629	MN595629	MN580328	MN580228	MN580228	MN580268	MN580268	MN593509	MN567223
<i>Dacrymyces chrysocornus</i> s.lat. 1	UPS F-940134	Sweden	MN593446	MN595630	MN595630	MN580329	MN580229	MN580229	MN580269	MN580269	MN593510	MN567223
<i>Dacrymyces chrysocornus</i> s.lat. 2	TNS-F-15712, TUFC 13115, MAFF 240127	Japan	AB712494	AB712452	AB299073	—	—	—	—	—	—	—
<i>Dacrymyces chrysospermus</i> s.lat.	FPL 11353	Japan	—	AF287855	AF287855	—	—	—	—	—	—	—
	FPL 8953, AFTOL-ID 528	Japan	AY705954	DQ205684	AY691892	DQ028587	DQ0785785	DQ0785785	DQ381845	DQ381845	MN593511	MN567223
<i>Dacrymyces aff. chrysospermus</i>	UPS F-593536	New Zealand	MN593447	MN595631	MN595631	MN580330	MN580230	MN580230	MN580270	MN580270	MN593511	MN567223
<i>Dacrymyces citrinus</i>	*PDD 107915, ICMP 21227	New Zealand	—	LC131417	LC131376	—	—	—	—	—	—	—
	PDD 107979, ICMP 21239	New Zealand	—	LC131418	LC131377	—	—	—	—	—	—	—
	*PDD 105052, ICMP 20517	New Zealand	—	LC131419	LC131378	—	—	—	—	—	—	—
<i>Dacrymyces cylindricus</i>	PDD 107989	New Zealand	—	LC131420	LC131379	—	—	—	—	—	—	—

Table 1 (cont.)

Taxon	Country	Herbarium voucher or isolate	GenBank accession numbers											
			18S	ITS	28S	TEF-1 α	RPB1	RPB2	12S	ATP6				
<i>Femisia peziziformis</i> s.lat. (cont.)	Japan	TNS-F-15737, TUF-C 13833, MAFF 240152	AB712518	AB712476	AB299080	—	—	—	—	—	—	—	—	—
<i>Femisia uniseptata</i>	Japan	TNS-F-54019, MAFF 246082	—	LC222844	LC222843	—	—	—	—	—	—	—	—	—
<i>Guepinopsis buccina</i>	Japan	TNS-F-15738, TUF-C 13344, MAFF 240153	AB712519	AB712477	AB299085	—	—	—	—	—	—	—	—	—
	Spain	UPS F-940947	MN593464	MN595643	MN580346	MN580246	—	—	—	—	—	—	—	—
	USA	PBM 2264 (WTU), AFTOL-ID 888	DO667157	DO206986	AY745711	DO028588	—	—	—	—	—	—	—	—
<i>Heterotextus militinus</i> s.lat. 1	New Zealand	TENN 42208	MN593465	MN595644	MN580347	MN580247	—	—	—	—	—	—	—	MN593522
	New Zealand	PDD 104962, ICMP 20435	—	LC131437	LC131401	—	—	—	—	—	—	—	—	—
<i>Heterotextus militinus</i> s.lat. 2	New Zealand	ICMP 16702	AB712520	AB712478	AB712436	—	—	—	—	—	—	—	—	—
	New Zealand	PDD 107924, ICMP 21229	—	LC131438	LC131402	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 1	Japan	TNS-F-61298, NBRC 110568	LC004012	LC004009	LC003901	—	—	—	—	—	—	—	—	—
	Japan	TNS-F-61318	LC004015	LC004001	LC003914	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 2	Japan	TNS-F-61311	LC004014	LC004002	LC003915	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 3	Japan	TNS-F-61312	LC004011	LC004010	LC003921	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 4	Japan	TNS-F-61322	LC004013	LC004004	LC003925	—	—	—	—	—	—	—	—	—
	Japan	10141-1D7	LC004022	LC003999	LC003882	—	—	—	—	—	—	—	—	—
	Japan	NBRC 110589	LC004025	LC003996	LC003879	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 5	Japan	NBRC 110590	LC004024	LC003997	LC003880	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 6	Japan	NBRC 110592	LC004020	LC003977	LC003884	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 7	Japan	NBRC 110594	LC004018	LC004006	LC003887	—	—	—	—	—	—	—	—	—
	Japan	NBRC 110595	LC004016	LC004008	LC003889	—	—	—	—	—	—	—	—	—
	Japan	NBRC 110596	LC004019	LC004005	LC003886	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 8	Japan	NBRC 110598	LC004027	LC003994	LC003890	—	—	—	—	—	—	—	—	—
Unidentified <i>Dacrymycetaceae</i> 9	Japan	NBRC 110599	LC004026	LC003995	LC003891	—	—	—	—	—	—	—	—	—
<i>Dacryonaemataceae</i>														
<i>Dacryonaema macnabii</i>	Sweden	UPS F-940949	MN593472	MN595650	MN595650	MN580353	—	—	—	—	—	—	—	MN567232
	Sweden	UPS F-940951	MN593473	MN595651	MN595651	MN580354	—	—	—	—	—	—	—	—
	Sweden	UPS F-940952	MN593474	MN595652	MN595652	MN580355	—	—	—	—	—	—	—	MN567233
	Sweden	UPS F-940952	MN593475	MN595653	MN595653	MN580356	MN580211	—	—	—	—	—	—	—
	UK	*K(M) 81678	—	MN595681	MN593499	—	—	—	—	—	—	—	—	—
	Sweden	UPS F-940954	MN593476	MN595654	MN595654	MN580357	MN580212	—	—	—	—	—	—	MN567234
	Sweden	UPS F-940993	MN593477	MN595655	MN595655	—	MN580213	—	—	—	—	—	—	MN567235
	Finland	UPS F-940997	MN593478	MN595656	MN595656	MN580358	—	—	—	—	—	—	—	MN567236
	Sweden	UPS F-729963	—	MN595682	—	—	—	—	—	—	—	—	—	—
	Sweden	UPS F-940980	—	MN595657	MN595657	—	—	—	—	—	—	—	—	MN567237
	Sweden	UPS F-940990	—	MN595658	MN595658	—	—	—	—	—	—	—	—	MN567238
	Sweden	UPS F-940991	—	MN595659	MN595659	—	—	—	—	—	—	—	—	MN567239
	Norway	O 160045	—	MN595660	MN595660	MN580359	MN580214	—	—	—	—	—	—	—
<i>Dacryonaema macrosporium</i>	Finland	UPS F-940998	MN593479	MN595661	MN595661	MN580360	MN580215	—	—	—	—	—	—	MN567240
	Finland	UPS F-941001	MN593480	MN595662	MN595662	MN580361	MN580216	—	—	—	—	—	—	MN567241
	Finland	UPS F-940999	MN593481	MN595663	MN595663	—	—	—	—	—	—	—	—	MN567242
	Finland	UPS F-941002	—	MN595664	MN595664	—	—	—	—	—	—	—	—	MN567243
	Finland	*UPS F-941000	—	MN595665	MN595665	—	—	—	—	—	—	—	—	MN567244
	Sweden	UPS F-941003	MN593466	MN595664	MN595664	MN580348	MN580217	—	—	—	—	—	—	MN567228
<i>Dacryonaema rufum</i>	Sweden	UPS F-730180	MN593467	MN595665	MN595665	—	—	—	—	—	—	—	—	—
	Sweden	UPS F-941005	MN593468	MN595667	MN595667	—	—	—	—	—	—	—	—	—
	Canada	TRTC 48002 in UPS	MN593469	MN595669	MN595669	—	—	—	—	—	—	—	—	—
	Sweden	UPS F-941005	—	MN595647	MN595647	MN580349	MN580209	—	—	—	—	—	—	MN567229
	Norway	O 295442	—	MN595646	MN595646	—	—	—	—	—	—	—	—	—
	Norway	O 295528	—	MN595648	MN595648	MN580350	—	—	—	—	—	—	—	—
	Finland	UPS F-941012	MN593470	MN595649	MN595649	MN580351	—	—	—	—	—	—	—	MN567230
	Sweden	UPS F-941011	MN593471	MN595680	MN593498	MN580352	—	—	—	—	—	—	—	MN567231
<i>Uniliacrymaeaceae</i>														
<i>Uniliacryma bispora</i>	Sweden	UPS F-941254	MN593488	MN595670	MN595670	MN580367	MN580253	—	—	—	—	—	—	—
	Sweden	UPS F-941255	MN593489	MN595671	MN595671	MN580368	MN580254	—	—	—	—	—	—	—
	Sweden	UPS F-941268	MN593490	MN595672	MN595672	MN580369	MN580255	—	—	—	—	—	—	—

Table 1 (cont.)

Taxon	Herbarium voucher or isolate	Country	GenBank accession numbers									
			18S	ITS	28S	TEF-1 α	RPB1	RPB2	12S	ATP6		
<i>Unilacryma bispora</i> s.lat. (cont.)	*UPS F-941257	Sweden	MN593491	MN595673	MN595673	MN580370	MN580256	MN580315				
	UPS F-941266	Sweden	MN593492	MN595674	MN595674	MN580371	MN580257	MN580316				
	UPS F-941274	Finland	MN593493	MN595675	MN595675	MN580372		MN580317				
	454_577 (ENAS)	Sweden		KM494136								
<i>Unilacryma unisporea</i> s.lat.	TNS-F-15731, TUFJFC 13832, MAFF 240146	Japan	AB712510	AB712468	AB712432			AB712552				
	TNS-F-38904, TUFJFC:13839	Japan	AB712511	AB712469	AB712432			AB712553				
	1310149-4 (ENAS)	Japan			LC003981							
	1307158-5 (ENAS)	Japan			LC003963							
	13061411-6 (ENAS)	Japan			LC003959							
	13061410-8 (ENAS)	Japan			LC003958							
	1306145-2 (ENAS)	Japan			LC003956							
	1305158-9 (ENAS)	Japan			LC003948							
	1311137-5 (ENAS)	Japan			LC003984							
	1310144-8 (ENAS)	Japan			LC003978							
	1312125-3 (ENAS)	Japan			LC003986							
	1307153-10 (ENAS)	Japan			LC003961							
	UPS F-941277	Sweden	MN593483	MN595665	MN595665	MN580362	MN580248	MN580307	MN567245			
UPS F-941278	Sweden	MN593484	MN595666	MN595666	MN580363	MN580249	MN580308	MN567246				
UPS F-941279	Sweden	MN593485	MN595667	MN595667	MN580364	MN580250	MN580309	MN567247				
UPS F-941291	Sweden	MN593486	MN595668	MN595668	MN580365	MN580251	MN580310	MN567248				
UPS F-941298	Finland	MN593487	MN595669	MN595669	MN580366	MN580252	MN580311					
Unknown family												
Unidentified species	NBRC 110597	Japan	LC004017	LC004007	LC003888							

This sampling comprised c. 90 ingroup species (not all of them identified) and included, at least in a broad sense, the types of all genera accepted by McNabb (1964, 1965a–e, 1966, 1973), and all genera accepted by Oberwinkler (2014) except *Arrhytidia*. The type of *Arrhytidia* (*A. flava*) was considered a species of *Dacrymyces* (*D. marginatus*), placed close to the *D. chrysocomus* group by McNabb (1973). No recent collections identified as such are known.

We selected eight DNA regions for phylogenetic analyses: three linked, multicopy regions in the nuclear ribosomal DNA, viz.,

- 1) the nuclear ribosomal small subunit (18S);
- 2) the internal transcribed spacer (ITS) region (including the ITS1, 5.8S, and ITS2); and
- 3) the D1–D3 domains of the nuclear ribosomal large subunit (28S).

We also selected three unlinked, low-copy, protein-coding, nuclear regions, viz.,

- 4) a c. 900–1100 bp long fragment from the largest subunit of the DNA-directed RNA polymerase II gene between domains A and C (*RPB1*);
- 5) a c. 1500–1700 bp long fragment from the second largest subunit of the DNA-directed RNA polymerase II gene between domains 6 and 11 (*RPB2*); and
- 6) a c. 900–1000 bp long fragment at the end of the translation elongation factor 1 alpha (*TEF-1 α*) gene.

Finally, we included two mitochondrial regions, viz.,

- 7) the mitochondrial ribosomal small subunit (12S); and
- 8) the ATP synthase subunit 6 (*ATP6*).

An outgroup was selected by retrieving from MycoCosm (Grigoriev et al. 2014) representatives from 20 of the c. 26 orders currently accepted of *Tremellomycetes* (three species) and *Agaricomycetes* (17 species). Samples were selected (Table 2) to minimise the amount of missing data in the outgroup. For each sample, nrDNA and mitochondrial regions were retrieved through BLAST searches using already generated sequences (e.g., from GenBank) of the same species or close relatives, keeping the best scoring hit. *RPB1*, *RPB2*, and *TEF-1 α* sequences were retrieved directly from the annotated contigs through the MycoCosm search option.

For morphological and cytological studies, the following fresh samples from Sweden of species also molecularly analysed were used: *Dacrymyces chrysocomus* s.lat. (Uppland, Uppsala, Lunsen, N59.79390° E17.68667°, on *Pinus sylvestris* branches, 19 Nov. 2017, J.C. Zamora, UPS F-940946); *D. chrysocomus* s.lat. (Uppland, Solna, Järva, N59.381147° E17.998386°, on *Pinus sylvestris* branches, 08 June 2018, J.C. Zamora, UPS F-940135); *D. estonicus* (Uppland, Uppsala, Lunsen, N59.79390° E17.68667°, on *Pinus sylvestris* branches, 19 Nov. 2017, J.C. Zamora, UPS F-941253); *D. stillatus* (Uppland, Funbo, c. 400 m north of Fjällnora farm, N59.840762° E17.913327°, on *Pinus sylvestris* branch, 14 Nov. 2017, J.C. Zamora & S. Ekman, UPS F-941285); *D. stillatus* (Uppland, Solna, Hagalund, N59.361045° E18.029935°, on an old *Picea abies* trunk, 01 July 2018, J.C. Zamora & I. Olariaga, UPS F-939950); *D. tortus* s.lat. 3 (Södermanland, Stockholm, Årsta, on *Pinus sylvestris* branches, 10 Sept. 2017, J.C. Zamora & P. Posio, UPS F-941018); *D. tortus* s.lat. 4 (Södermanland, Stockholm, Masmo, N59.25429° E17.89356°, on *Pinus sylvestris* branches, 15 Oct. 2017, J.C. Zamora & P. Posio, UPS F-941252); *D. tortus* s.lat. 4 (Uppland, Solna, Bergshamra, on *Pinus sylvestris* branches, 23 June 2018, J.C. Zamora, UPS F-941021).

We also performed an independent and denser sampling to investigate species delimitations in *Dacryonaema* and *Unilacryma*. In total, we included DNA data of eight individuals of *Da. rufum* from Canada, Finland, Norway, and Sweden, 12 of *Da. macnabbii* from the UK (holotype), Finland, and Sweden, six of an unidentified and potentially new *Dacryonaema* spe-

Table 2 Species for which DNA sequence data were retrieved from the genome database at MycoCosm (or partially GenBank), and references of the place of publication of the genomes. Data was downloaded on or prior to 1 December 2018.

Class	Order	Species	References
<i>Agaricomycetes</i>	<i>Agaricales</i>	<i>Agaricus bisporus</i>	Morin et al. (2012)
<i>Agaricomycetes</i>	<i>Amylocorticiales</i>	<i>Plicaturopsis crispa</i>	Kohler et al. (2015)
<i>Agaricomycetes</i>	<i>Atheliales</i>	<i>Fibulorhizoctonia</i> sp.	Nagy et al. (2015)
<i>Agaricomycetes</i>	<i>Auriculariales</i>	<i>Auricularia subglabra</i>	Floudas et al. (2012)
<i>Agaricomycetes</i>	<i>Boletales</i>	<i>Suillus luteus</i>	Kohler et al. (2015)
<i>Agaricomycetes</i>	<i>Cantharellales</i>	Unknown (as <i>Clavulina</i> sp.)	Unpublished*
<i>Agaricomycetes</i>	<i>Corticiales</i>	<i>Vuilleminia comedens</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Gastrales</i>	<i>Sclerogaster hysterangioides</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Gloeophyllales</i>	<i>Heliocybe sulcata</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Gomphales</i>	<i>Ramaria rubella</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Hymenochaetales</i>	<i>Phellinidium ferrugineofuscum</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Jaapiales</i>	<i>Jaapia argillacea</i>	Riley et al. (2014)
<i>Agaricomycetes</i>	<i>Phallales</i>	<i>Mutinus elegans</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Polyporales</i>	<i>Fomes fomentarius</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Russulales</i>	<i>Russula vinacea</i>	Unpublished*
<i>Agaricomycetes</i>	<i>Sebacinales</i>	<i>Serendipita vermifera</i>	Kohler et al. (2015)
<i>Agaricomycetes</i>	<i>Thelephorales</i>	<i>Thelephora ganbajun</i>	Nuclear genome unpublished* Mitochondrial genome in Wang et al. (2017)
<i>Dacrymycetes</i>	<i>Dacrymycetales</i>	<i>Calocera</i> cf. <i>furcata</i> (as <i>Ca. cornea</i>)	Nagy et al. (2015)
<i>Dacrymycetes</i>	<i>Dacrymycetales</i>	<i>Calocera viscosa</i>	Nagy et al. (2015)
<i>Dacrymycetes</i>	<i>Dacrymycetales</i>	<i>Cerinomyces ceraceus</i>	Unpublished*
<i>Dacrymycetes</i>	<i>Dacrymycetales</i>	<i>Dacryopinax primogenitus</i>	Floudas et al. (2012)
<i>Tremellomyces</i>	<i>Filobasidiales</i>	<i>Filobasidium wieringae</i>	Unpublished*
<i>Tremellomyces</i>	<i>Tremellales</i>	<i>Tremella mesenterica</i>	Floudas et al. (2012)
<i>Tremellomyces</i>	<i>Trichosporonales</i>	<i>Cutaneotrichosporon oleaginosus</i>	Kourist et al. (2015)

* Unpublished genome data are used here with permission from their respective authors (see acknowledgements).

cies from Finland and Norway, seven of *Unilacryma unispora* s.lat. from Finland, Japan, and Sweden, and six of a bisporic *Unilacryma* from Finland and Sweden. In addition, all available GenBank sequences that had a high similarity in BLAST searches ($\geq 99\%$) to any of the studied DNA regions were added (Table 1). Several additional specimens of *Da. macnabii*, *Da. rufum*, *U. unispora* s.lat., and the bisporic *Unilacryma* were collected and morphologically studied to gather a better knowledge of their intraspecific variation (see taxonomic part).

Morphological study

Whenever possible, samples were studied morphologically and photographed when fresh, either immediately after being collected or otherwise kept in a refrigerated (4 °C) humid chamber for a maximum of 48 h. Alternatively, when samples were found in a dry state in the field, they were either studied directly, if the structures were well-developed, or moistened with tap water by submerging the entire substrate a few minutes, and then placed in a refrigerated humid chamber for one to seven days before study. General terminology largely follows Oberwinkler (1994), and for clamp connections we follow Stalpers (1978). In particular, 'hyphidia' is preferred over 'dikaryophyses', as the last term is not consistently applied for dikaryotic hyphidia in other groups of the *Agaricomycotina*, particularly in aphyllorphoid fungi. Also, we prefer the term 'young basidium' instead of 'probasidium' to avoid confusion with the thick-walled structures in *Pucciniomycotina* and *Ustilaginomycotina*, from which the basidium forms (Clemençon 2004).

Macromorphology was studied under a Leica MZ 75 dissecting microscope, and macroscopic photographs were taken with a coupled Infinity 1 macro camera. Micromorphological study was performed by mounting thin hand-cut sections of the fruitbodies in water or a 5–10 % w/v aqueous solution of KOH, and gently squeezing them under the cover slip. All measurements were taken at 630 \times magnification in KOH solution. The hilar appendix

was not included in basidiospore measurements. Microscopic photographs were taken in a Zeiss Axio Imager A1 microscope, fitted with differential interference contrast (DIC) and equipped with an AxioCam ICc3 digital camera. For some of the most voluminous microscopic structures, a series of photographs were stacked using Combine ZP.

An aqueous solution of 1 % w/v potassium triiodide (KI₃, Lugol's solution) was used to check the reaction of the carotenoid contents in some species. Nuclear staining was done by using toluidine blue. The best results were achieved when fresh samples were pre-treated with 5–10 % KOH, quickly rinsed with tap water, and stained with a recently prepared alkaline solution (pH \approx 10.5–11) of 0.1 % toluidine blue.

We performed analyses of variance (ANOVA) on the basidiospore length, basidiospore width, and the length/width ratio (Q) of *Dacryonaema* spp. and *Unilacryma* spp. samples, in order to detect significant differences ($p < 0.001$) among the species in each genus. Diagnostic plots for residuals (residuals vs fitted values, normal Q-Q, scale-location, and residuals vs leverage) were checked to ensure that no model assumptions were severely violated. Cohen's f was used as an estimator of effect size. Since *Dacryonaema* included three species, when any significant difference was found in the ANOVA, the Tukey's honest significant difference (Tukey's HSD) post hoc test was applied to identify pairs of means different at $p < 0.001$. All analyses were done using the basic package of R (R Core Team 2018) and sjstats v. 0.17.4 (Lüdecke 2019). For each species, we measured 100–170 basidiospores from randomly selected fruitbodies (1–30 basidiospores per fruitbody). Measurements were taken either directly under the light microscope (630 \times , immersion oil), or on photographs taken as previously indicated, with the aid of Piximètre v. 5.9 (Henriot & Cheyfe 2016). Length and width values were rounded to the nearest 0.1 μ m, and Q values to the nearest 0.01.

After study, specimens were dried at room temperature and deposited in UPS. Additional herbarium specimens from F, K, NCU, O, S, TENN, UBC, and UPS were studied (herbarium acronyms following Thiers 2019). Geographical coordinates are provided in the World Geodetic System 1984 (WGS84).

Lab work and primer design

Genomic DNA was extracted from a single fruitbody or part of a fruitbody (c. 1 mm³), rarely (species with very small fruitbodies like *Dacryonaema rufum*) from 2–5 adjacent fruitbodies. The samples were disrupted in 1.5 ml microcentrifuge tubes with a plastic micropipette either after being frozen in dried state with liquid nitrogen, or directly in the lysis buffer. We used the DNeasy Plant Mini Kit (Qiagen, Germany), following the manufacturers' instructions, but increased the lysis buffer incubation time to up to two hours. For some samples, including the type specimens of *Dacrymyces macnabbii* (K(M) 81678) and *Unilacryma unisporea* (NCU F-0026842), we followed the Chelex extraction protocol of Ferencova et al. (2017), in order to minimise the amount of material used.

Forty-six different primers were used for amplifying the DNA regions mentioned above (Table 3), comprising both general primers used in other studies as well as newly designed primers for 12S, *RPB1*, *RPB2*, and fragments of ITS1 and ITS2 (specific primers for these last two loci designed only for *Dacryonaema* and *Unilacryma*). ITS4B (Gardes & Bruns 1993) could not be

used because the primer site has four mismatches in the genus *Unilacryma*, one of them affecting the last base at the 3'-end, and three mismatches in the genus *Dacryonaema*, in both cases resulting in unsuccessful PCR amplifications. Primer design was carried out by identifying conserved regions in *Dacrymycetes* alignments containing already generated sequences of those loci from MycoCosm, GenBank and own sequences obtained using general primers. We tried to design the primers so that their CG content was as close as possible to 50 %, with a length in the range of 17–25 bp, melting temperatures differing less than 5 °C within a pair, and avoiding hairpin and duplex formation. Calculations were done using OligoAnalyzer v. 3.1 (Integrated DNA Technologies®).

PCR reactions were performed in a final volume of 20–25 µl using MasterMix (Qiagen, Germany). Numerous primer combinations and PCR conditions were tried for each DNA region, so we only report the most successful PCR conditions used for amplifying the majority of samples. The PCR programmes always consisted on an initial denaturation of 2 min at 94 °C and a final extension step of 7 min at 72 °C. The central cycles and primer pairs for the different loci greatly varied and are summarised in Table 4. When needed, nested PCRs were used for *RPB1* (first PCR using DacryRPB1-1F/DacryRPB1-2r, second PCR with DacryRPB1-A/DacryRPB1-C), *RPB2* (first PCR with DacryRPB2-6F/DacryRPB2-11aR, second PCR with DacryRPB2-6.2F/DacryRPB2-11bR), and *TEF-1α* (first PCR with EF1-1018F/EF1-2218R, second PCR with Efdf/EF1-1953R).

Table 3 Primers used in this study. Within each DNA region, primers are ordered by primer site from 5' to 3', and then by increasing specificity. F = forward, R = reverse.

Locus	Primer name	Sequence (5' → 3')	Sense	Comment	References
18S (nrSSU)	NS1	GTAGTCATATGCTTGTCTC	F		White et al. (1990)
	NS21UCB	GAATAATAGAATAGGACG	F		Gargas & Taylor (1992)
	NS4	CTTCGGTCAATTCCTTTAAG	R		White et al. (1990)
	SR6	TGTTACGACTTTTACTT	R		Vilgalys, unpublished
	NS8	TCCGCAGGTTACCTACGGA	R		White et al. (1990)
nrITS	ITS1F	CTTGCTCATTAGAGGAAGTAA	F		Gardes & Bruns (1993)
	ITS2	GCTGCGTTCTTCATCGATGC	R		White et al. (1990)
	ITS3	GCATCGATGAAGAACGCAGC	F		White et al. (1990)
	ITS4	TCCTCCGCTTATTGATATGC	R		White et al. (1990)
	ITS1DacryoUnil-F	CTGTTGCTGAGAAGCTGATCA	F	Designed for <i>Dacryonaema/Unilacryma</i>	This study
	ITS1Dacryo-r	TTTYGTTTATGACCGCTTTTCG	R	Designed for <i>Dacryonaema</i>	This study
	ITS1Unil-r	TTTGGTTACGACCGCTTGC	R	Designed for <i>Unilacryma</i>	This study
	ITS2Dacryo-F	ATGCCTGTTTGAATGTGCGCA	F	Designed for <i>Dacryonaema</i>	This study
ITS2Unil-F	GAGTGTGTGCTGTGCCT	F	Designed for <i>Unilacryma</i>	This study	
ITS2DacryoUnil-r	CGAAGACGGTTGGAGAGC	R	Designed for <i>Dacryonaema/Unilacryma</i>	This study	
28S (nrLSU)	LR0R	ACCCGCTGAACCTAAGC	F		Cubeta et al. (1991)
	LR3	CCGTGTTCAAGACGGG	R		Vilgalys & Hester (1990)
	LR3R	GTCTTGAAACACGGACC	F		Vilgalys, unpublished
	LR5	TCCTGAGGGAAACTTCG	R		Vilgalys & Hester (1990)
<i>TEF-1α</i>	EF1-1018F	GAYTTCATCAAGAACATGAT	F		Stielow et al. (2015)
	Efdf	AAGGAYGGNCARACYCNGARCAAYGC	F		Rehner, unpublished
	EF1-1953R	CCRGCRACRGRTRGTCTCAT	R		Rehner, unpublished
	EF1-2218R	ATGACACCRACRGRACRGTYTG	R		Rehner & Buckley (2005)
<i>RPB1</i>	DacryRPB1-1F	GAAGATCGAGCATTCCGGAGGT	F	Designed for <i>Dacrymycetes</i>	This study
	gRPB1-A	GAKTGTCCCKGGWCATTTTGG	F		Stiller & Hall (1997)
	DacryRPB1-A	GAYTGYCCKGGRCAYTTTGG	F	Improved for <i>Dacrymycetes</i>	This study
	fRPB1-C	CNGCDATNTRTRTCCATRTA	R		Matheny et al. (2002)
	DacryRPB1-C	CRGCAATCTCGTTGCCATGTA	R	Improved for <i>Dacrymycetes</i>	This study
DacryRPB1-2r	GTTCRGGRTATGTHAGCGTCAT	R	Designed for <i>Dacrymycetes</i>	This study	
<i>RPB2</i>	RPB2-6F	TGGGGKWTGGTYTYGCCCTGC	F		Liu et al. (1999)
	bRPB2-6F	TGGGGYATGGTNTGYCCYGC	F		Matheny (2005)
	DacryRPB2-6F	TGGGGNATGGTVTYGCCYGC	F	Improved for <i>Dacrymycetes</i>	This study
	DacryRPB2-6.2F	TTYGTSAAAYGGKNTTGGATGG	F	Designed for <i>Dacrymycetes</i>	This study
	fRPB2-7cF	ATGGGYAARCAAGCYATGGG	F		Liu et al. (1999)
	bRPB2-10.9R	GTRAASGGYGTGGCRTCYCC	R		Hall, unpublished
	DacryRPB2-11bR	TGRATCTTRTCRTCVACCATRTGCT	R	Designed for <i>Dacrymycetes</i>	This study
	fRPB2-11aR	GCRTGGATCTTRTCRTCSACC	R		Liu et al. (1999)
DacryRPB2-11aR	GCACGAGCRTGRATCTTRTCRTC	R	Designed for <i>Dacrymycetes</i>	This study	
12S (mrSSU)	MS1	CAGCAGTCAAGAATATTAGTCAATG	F		White et al. (1990)
	DacryMS1	CAGCAGCTAAGAATWTTTACAATG	F	Improved for <i>Dacrymycetes</i>	This study
	Dacry12S-4r	AAYAACATRCTTCACTTCGT	R	Designed for <i>Dacrymycetes</i>	This study
	MS2	GCGGATTATCGAATTAATAAC	R		White et al. (1990)
	Dacry12S-2r	AGARATGGTAAGRTTTTACG	R	Designed for <i>Dacrymycetes</i>	This study
<i>ATP6</i>	ATP6-3	TCTCCTTTAGAACAATTTGA	F		Kretzer & Bruns (1999)
	ATP6-4	AAGTACGAAWACWTGWGMMTG	R		Kretzer & Bruns (1999)
	ATP6-2	TAATTCTANWGCATCTTTAATRTA	R		Kretzer & Bruns (1999)

PCR products were purified using ExoSAP-IT™ (Applied Biosystems) or Illustra™ ExoProStar™ (GE Healthcare), following the manufacturer's instructions but adding 1/4 of the recommended volume of enzyme mix and extending the reaction time to 30 min. Purified PCR products were sequenced by Macrogen with the same primers used for amplification.

Sequence edition and alignment

Raw chromatograms were assembled and edited using Sequencher v. 4.1.4 (Gene Codes, USA). Even if most DNA extractions were generated from a single fruitbody, chromatograms with scattered double peaks were found in several samples. Positions with a double signal were often similar or identical among different samples (thus, from independent extractions and PCRs) of a single species, more abundant in introns and, when affecting the exons of protein coding genes, they were almost always restricted to the third codon position, resulting in alternative codons for the same amino acid (synonymous mutations). This observation suggests that more than one allele was present for a particular DNA region in some samples. For the purpose of phylogenetic reconstruction, these positions were coded using ambiguity codes. In case there was a polymorphism affecting the length of the alleles (e.g., in ITS, and often only a single base indel), the shortest copy was retained.

Two datasets were assembled and analysed independently, one for higher level taxa (overall phylogeny) and one for the species delimitation analyses of the genera *Dacryonaema* and *Unilacryma*. The ITS1 region was considered to start immediately after the highly conserved motif 'AGGACATTA' at the end of the 18S. The 5' limit of the 5.8S region was defined by comparing the aligned sequences to annotated sequences from ITSoneBD (Santamaria et al. 2017). The 3'-end of the

5.8S and the 5' start of the 28S were defined using the annotate option of the ITS2 database (Keller et al. 2009). The ITS1 and ITS2 outgroup sequences were excluded for being largely unalignable among the distantly related taxa included in the dataset for the overall phylogeny. Likewise, repetitions at the end of the 28S (in *Cerinomyces* cf. *crustulinus*, *C. lagerheimii*, *Dacrymyces flabelliformis*, *D. subantarcticensis*, and *Dacryopinax indacocheae*) were removed, as well as two intron regions above 600 bp each in the 12S (present in *Vuilleminia comedens* and *Cerinomyces ceraceus* genomic data, and in the newly generated sequences of *Calocera cornea*, *Cerinomyces tortus* s.lat. 1, and *Dacrymyces* cf. *minutus* s.lat. 2). Additionally, we excluded a highly variable and gappy region in the 12S appearing between two easily alignable regions (the one in the 5' ending with the motif 'AGGGTTYGTAG', and the one in 3' starting with the motif 'TAGAATCW'). For the protein-coding DNA regions, intron positions were identified using open reading frames from the JGI and annotated sequences from GenBank as references (with few exceptions), and also excluded from the matrices. All sequences were trimmed to fit to the longest sequences generated by us and to remove poor read ends.

For the overall phylogeny, all alignments were estimated using PASTA v. 1.7.8 (Mirarab et al. 2015). We performed three independent runs with 10 iterations, maximum subset size being half of the sequences included in the alignment, the mask option deactivated, and keeping the best likelihood score alignment. MAFFT (L-INS-i option) was used as the aligner, OPAL as the merger, and RAxML as the tree estimator. For the ribosomal DNA regions we used the GTR + Γ model, and for the exon regions of the protein coding genes, the WAG + I + Γ model for the datasets temporarily translated into amino acids. The alignment with the best score after the three runs was manually

Table 4 Most successful primer combinations and central cycles of the PCR programmes.

DNA region	Primers	Central cycles
18S	Whole region: NS1/NS8 or NS1/SR6	
	Two parts: 1st: NS1/NS4 2nd: NS21/SR6	(i) 5 cycles of 45 s at 94 °C, 45 s at 53 °C (decreasing 1 °C/cycle), 1.5 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 48 °C, 1.5 min at 72 °C
ITS+28S	Both regions: ITS1F/LR5	
	Two parts: ITS: ITS1F/ITS4 28S: LR0R/LR5	(i) 5 cycles of 45 s at 94 °C, 45 s at 55 °C (decreasing 1 °C/cycle), 1.5 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 50 °C, 1.5 min at 72 °C
	Other smaller parts: ITS1F/ITS2, ITS3/ITS4, LR0R/LR3, and LR3R/LR5	Same as above, but extension reduced to 45 s at 72 °C
	Specific for <i>Dacryonaema/Unilacryma</i> : ITS1: ITS1DacryoUnil-F/ITS1Dacryo-r or ITS1Unil-r ITS2: ITS2Dacryo-F or ITS2Unil-F/ITS2DacryoUnil-r	40 cycles of 45 s at 94 °C, 45 s at 60 °C, 30 s at 72 °C
<i>RPB1</i>	gRPB1-A/fRPB1-C, or DacryRPB1-1F/DacryRPB1-2r, or DacryRPB1-A/DacryRPB1-C	(i) 5 cycles of 45 s at 94 °C, 45 s at 58 °C (decreasing 1 °C/cycle), 1.5 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 53 °C, 1.5 min at 72 °C
<i>RPB2</i>	RPB2-6F or bRPB2-6F or fRPB2-7cF/fRPB2-11aR or bRPB2-10.9R (all six combinations), or DacryRPB2-6F/DacryRPB2-11aR, or DacryRPB2-6.2F/DacryRPB2-11bR	(i) 5 cycles of 45 s at 94 °C, 45 s at 58 °C (decreasing 1 °C/cycle), 1.5 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 53 °C, 1.5 min at 72 °C
<i>TEF-1α</i>	EF1-1018F or Efdf/EF1-1953R or EF1-2218R (all four combinations)	(i) 5 cycles of 45 s at 94 °C, 45 s at 60 °C (decreasing 1 °C/cycle), 1.5 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 55 °C, 1.5 min at 72 °C
12S	MS1/MS2, or DacryMS1/Dacry12S-2r, or DacryMS1/Dacry12S-4r	(i) 5 cycles of 45 s at 94 °C, 45 s at 53 °C (decreasing 1 °C/cycle), 1 min at 72 °C (ii) 35 cycles of 45 s at 94 °C, 45 s at 48 °C, 1 min at 72 °C
<i>ATP6</i>	ATP6-3/ATP6-2 or ATP6-3/ATP6-4	Programme according to Kretzer & Bruns (1999)

corrected (minor modifications) and submitted to a last run in PASTA, using the parameters indicated above. Since PASTA already optimises the likelihood under a phylogenetic framework, and the resulting alignments seem to be phylogenetically accurate (Mirarab et al. 2015), neither further manual refinement was done nor parts of the alignments were excluded. All the matrices independently aligned in PASTA contained less than 15 % of missing data at the ends of the sequences, except for the *RPB2* dataset where c. 1/3 of the sequences (all from GenBank) were only represented by a fragment of 843 bp covering an easily alignable exon region (a single indel due to three additional codons in the *Tremellomyces* sequences).

For the *Dacryonaema* + *Unilacryma* dataset, the 5.8S, *TEF-1 α* , and *ATP6* alignments contained no indels. The *RPB2* alignment contained a single, unambiguous indel of three bases in length, corresponding to a codon position. The 18S and 28S alignments contained only three and two unambiguous single-base indels, respectively, and were manually aligned. The *Dacryonaema* and *Unilacryma RPB1* datasets contained a single, unambiguous codon indel and were aligned and analysed separately because the incongruence detected in the overall phylogeny (see ‘Congruence among datasets’ below) suggests that they may be non-orthologous. The ITS1 and ITS2 regions were independently aligned using BALi-Phy v. 3.3 (Suchard & Redelings 2006). Since BALi-Phy is computationally intensive, we reduced the datasets by selecting two or three sequences within the same species, keeping the maximum variation but avoiding those sequences with more missing data or ambiguous base calls. A GTR + I + Γ model was used for the substitutions, whereas an rs07 model was used for insertions and deletions. For each dataset, we executed 6 simultaneous runs with 5×10^4 iterations per run, checked convergence, and computed the alignments discarding the first 25 % of the runs as burn-in. The average standard deviation of splits in the posterior sample turned out to be smaller than 0.005, and all parameters analysed in Tracer v. 1.7 (Rambaut et al. 2018) had effective sample sizes exceeding 1000 for individual runs (≥ 7000 when combined). The resulting alignments were used as backbones for manually re-introducing the remaining sequences. Indels were coded as a separate binary dataset (gap present = 1, gap absent = 0) in FastGap v. 1.2 (Borchsenius 2009), using the simple indel coding method (Simmons & Ochoterena 2000). All final alignments were deposited in TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S24147>).

Congruence among datasets

Dataset congruence was assessed for both the overall phylogeny and the *Dacryonaema* + *Unilacryma* datasets through maximum likelihood (ML) analyses using IQ-TREE v. 1.6.8 (Nguyen et al. 2015). We performed standard, non-parametric bootstrap analyses with 200 replicates for the nrDNA dataset, each of the three nuclear protein-coding genes, and the mitochondrial dataset, assuming no recombination within each dataset. Each of the protein coding genes was partitioned by codon position, the nrDNA dataset was originally divided into five partitions (18S, ITS1, 5.8S, ITS2, and 28S), and the 12S and *ATP6* genes were considered as potentially different partitions. Additionally, we also tested for incongruence among partitions of the nrDNA dataset. Incongruence was considered significant when a strongly supported clade from one phylogeny was contradicted by another strongly supported clade in another phylogeny (Mason-Gamer & Kellogg 1996). A bootstrap support (BS) threshold of 75 % was used, at or above which a clade was considered strongly supported. A significant case of incongruence (*Dacryonaema* in the *RPB1* dataset) in the overall phylogeny analyses was inspected for alternative explanations.

Possible deviations from nucleotide composition heterogeneity among taxa were checked with the IQ-TREE composition test. The amount of recombination, if any, was tested with GARD (Kosakovsky Pond et al. 2006) and SlidingBayes v. 0.94 (Paraskevis et al. 2005). Finally, we checked for heterotachy by performing analyses under the edge-unlinked partition model (Lopez et al. 2002) and the GHOST model (Crotty et al. 2017) in IQ-TREE. In the end, these processes as well as incomplete lineage sorting turned out to be unlikely candidates for the observed topological conflict, and consequently the *Dacryonaema RPB1* sequences were excluded from downstream analyses.

Phylogenetic analyses for the overall phylogeny

Two approaches were used for calculating phylogenies: (i) tree estimation based on the concatenated dataset of all loci; and (ii) coalescent-based species tree estimation joining independent single-locus tree phylogenies. In this set of analyses, gaps were treated as missing data. For the concatenated analyses, two strategies were used for phylogenetic reconstructions: maximum likelihood (ML) and Bayesian Inference (BI). ML analyses were performed using IQ-TREE v. 1.6.8 (Nguyen et al. 2015), and the model selection and partitioning scheme for the mitochondrial and nuclear regions were calculated by using the Bayesian Information Criterion in the integrated version of ModelFinder (Kalyaanamoorthy et al. 2017). Fourteen potential partitions were defined for the nuclear regions: 18S, ITS1, 5.8S, ITS2, 28S, and one for each codon position of *RBP1*, *RPB2*, and *TEF-1 α* , and four potential partitions were defined for the mitochondrial regions: 12S and each codon position of *ATP6*. Branch lengths were linked and proportional across partitions. The analysis was repeated twice, starting from random trees. Branch support was assessed by standard non-parametric bootstrap, performing 500 replicates, and through the Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT, Guindon et al. 2010), performing 1000 replicates. The purpose of using the SH-aLRT was to identify very short branches (with no signal from the data) that may have received spuriously high BS support (Guindon et al. 2010). Significant support is considered when BS ≥ 75 % and SH-aLRT ≥ 85 %.

Bayesian inference on the concatenated dataset was carried out using MrBayes v. 3.2.6 (Ronquist et al. 2012). We applied the same partitioning scheme obtained in the previous analysis with IQ-TREE, unlinking model parameters across partitions, and using model jumping to integrate over models and their parameter values in each partition (Huelsenbeck et al. 2004), allowing a gamma distributed rate heterogeneity across sites (approximated by four categories) and a proportion of invariant sites. We used a (1, 1, 1, 1, 1, 1) Dirichlet prior for the substitution rates, a (1, 1, 1, 1) Dirichlet prior on the state frequencies, and a uniform (0, 1) prior for the proportion of invariable sites. Branch lengths were linked and proportional across partitions. We used a compound Dirichlet prior on branch lengths in the overall phylogeny (1, 0.0588, 1, 1), based on tree length estimates from the best tree of the ML analysis. We set the temperature increment parameter to 0.1 after having observed swap rates between the cold and hot chains in preliminary runs. Four parallel runs starting with random trees were executed, each with four chains, for a maximum of 1×10^8 generations. The analysis was automatically stopped before reaching the maximum number of generations if the average standard deviation of split frequencies dropped below 0.01. Sampling was done every 1000th tree. The first 50 % of the analysis was eliminated as burn-in, and Tracer v. 1.7 (Rambaut et al. 2018) was used to ensure that effective sample size (ESS) for each parameter exceeded 200. RWTY (Warren et al. 2017) was used to check the convergence of tree topologies. The 50 % majority-rule tree with posterior probabilities (PP^{MrBayes}, considered signi-

ficant when ≥ 0.95) and average branch lengths was calculated from the post-burn-in trees, and visualised in FigTree v. 1.4 (Rambaut 2016).

ML and Bayesian approaches were also used for inferring a coalescent-based species tree from the overall phylogeny data. An ML species tree was estimated with ASTRAL III (Zhang et al. 2018, Rabiee et al. 2019) v. 5.6.3, using the independent ML phylogenetic trees obtained for each locus in IQ-TREE. Since ASTRAL uses a summary method for computing a species tree based on single-locus trees, limited accuracy is expected when the number of loci is low. Considering this limitation, we conducted two analyses, first by considering the effectively linked parts of the nrDNA as a single locus (analysis assuming five loci), and secondly by splitting the nrDNA into its units: 18S, ITS1, 5.8S, ITS2, and 28S (analysis of nine loci). The second analysis has the advantage of accounting for any unsupported discordances among those nrDNA regions, but has the shortcoming of giving too much weight to the nrDNA compared to other loci if recombination is absent among the mentioned nrDNA regions. Branches receiving $\leq 10\%$ BS support in each single-locus tree were collapsed using Newick Utilities v. 1.6 (Junier & Zdobnov 2010), as recommended by Zhang et al. (2018). Branch support was calculated as local posterior probabilities based on quartet support (QS, Sayyari & Mirarab 2016), and was considered significant when $QS \geq 0.95$. Species assignments of individuals mostly followed our identifications and those from the literature (e.g., Shirouzu et al. 2009, 2017), and tended to be conservative, i.e., in some cases very closely related and potentially cryptic species might have been merged: *Calocera viscosa* s.lat., *Dacrymyces chrysospermus* s.lat., and *Unilacryma unisporea* s.lat. In these three cases, all samples in each group formed a fully supported clade in the previous concatenated ML and Bayesian analyses. Exceptions were made when we detected either non-monophyly or a striking divergence among individuals in the ML analysis, e.g., the three clades of *Calocera* cf. *guepinioides* were considered as three independent species, and *Dacrymyces* cf. *minutus* s.lat. 1 and *D. aff. stillatus*, from Shirouzu et al. (2009), were treated as different species to our *D. cf. minutus* s.lat. 2 and *D. stillatus*, respectively.

StarBEAST 2, implemented in BEAST 2.5.1 (Bouckaert et al. 2014, Ogilvie et al. 2017), was used to infer a dated species chronogram for the overall phylogeny, based on the multispecies coalescent. In the absence of fossils for the included taxa, we used a secondary calibration for the *Dacrymycetes*, setting the stem age of the group by using a normal prior distribution with values for $\mu = 350.0$ Ma and $\sigma = 50.0$, according to the results from Floudas et al. (2012) (also largely agreeing with Zhao et al. 2017), under an uncorrelated lognormal relaxed clock model (Drummond et al. 2006). We used the uninformative 1/X prior on clock rates to allow the calibration to guide those clock rates, and a calibrated Yule tree model. Clock and tree models were unlinked among the following loci: nrDNA, *RPB1*, *RPB2*, *TEF-1 α* , and the mitochondrial regions. The original partitions (18S, ITS1, 5.8S, ITS2, 28S, 12S, and codon positions of each protein coding gene) were used since it is biologically implausible to combine partitions from different loci under the coalescent (e.g., codon positions of unlinked protein coding genes). We used bModelTest (Bouckaert & Drummond 2017) for coestimating site models during the runs, averaging among all extended named models. The result of model averaging was summarised using BModelAnalyser (part of the BEAST 2 package, Bouckaert et al. 2014). Given the general life cycle in *Agaricomycotina*, we considered a biparental inheritance for the mitochondrion, and we used the analytical population size integration as the population model, since the number of individuals per species was low, and we were not specifically interested in per-

species population sizes. Individuals were assigned to species as indicated above for the ML analysis in ASTRAL III. Initially, we executed six parallel runs starting from random trees for 5×10^8 generations, sampling every 10^{4th} tree, then checked the traces and ESSs in Tracer v. 1.7, and discarded three of them due to failure to reach convergence for some parameters (different parameters in each run and none affecting the statistics of the species tree, see Results and Discussion). The other three runs were resumed until completing 1×10^9 generations. The first 50% of these three runs was eliminated as burn-in. Tracer v. 1.7 (Rambaut et al. 2018) was used to check effective sample size (ESS) of model parameters (ideally > 200). RWTY (Warren et al. 2017) was used to check the convergence of the species tree topologies. The maximum clade credibility species tree with posterior probabilities (considered significant when ≥ 0.95) and node ages was calculated using TreeAnnotator v. 2.5.1 (part of the BEAST 2 package, Bouckaert et al. 2014) and visualised in FigTree v. 1.4.

Character evolution analyses

We inferred the history of selected morphological characters, including transformation counts and ancestral states at the nodes corresponding to the *Dacrymycetes* and the four major families *Cerinomycetaceae*, *Dacrymycetaceae*, *Dacryonaemataceae*, and *Unilacrymaceae*. We avoided further analyses of characters for which convincing inferences were already provided by Shirouzu et al. (2013), and not expected to be altered by our sampling, e.g., basidiospore septation pattern, fruitbody morphology, and presence or absence of a stalk. The following traits were analysed: (i) basidium morphology (0 = 2-spored, 1 = 1-spored); (ii) clamp connections presence and morphology (0 = clamps absent, 1 = short and rounded clamps present at most septa, 2 = long and angulose clamps mainly restricted to the bases of basidia and hyphidia, and subhymenial hyphae); (iii) hyphidia (0 = conspicuous and branched, 1 = unbranched or indistinct); (iv) terminal cells of cortical/marginal hyphae (0 = filamentose and simple, 1 = clearly dilated and simple, 2 = filamentose and branched); (v) carotenoids (0 = absent or indistinct, 1 = conspicuous at least in some cells); (vi) cytoplasmic lipid content (0 = indistinct and with no visible lipid drops in the cytoplasm, 1 = lipid drops visible when the lipid bodies coalesce); (vii) number of nuclei in recently discharged basidiospores before the formation of the first septum (0 = uninucleate, 1 = binucleate). Taxa with unknown states were coded as having prior probabilities equally divided among all known states. For simplicity, the character state matrix is shown next to the species tree obtained in the StarBEAST 2 analysis (Fig. 9). Our hypotheses, based on prior knowledge of the group, were as follows: (i) the two-sterigmate basidia, presence of clamps, branched hyphidia, filamentose and simple terminal cells of cortical/marginal hyphae, absence of conspicuous carotenoids, high cytoplasmic lipid content, and uninucleate young basidiospores are plesiomorphic in the *Dacrymycetes*; (ii) the long, angulose clamp connections at subhymenial hyphae and the bases of basidia and hyphidia, and the filamentose and branched terminal cells of cortical/marginal hyphae are plesiomorphic in the *Dacryonaemataceae*; (iii) binucleate young basidiospores and low cytoplasmic lipid content are acquired states plesiomorphic in the *Cerinomycetaceae*; (iv) the presence of carotenoids and unbranched hyphidia are acquired states plesiomorphic in the *Dacrymycetaceae*; (v) the 1-spored basidium, the dilated terminal cells of cortical/marginal hyphae, and the clampless septa are apomorphic conditions within different families in the *Dacrymycetes*.

Character transformation counts and ancestral state reconstructions were performed by stochastic mapping with SIMMAP (Bollback 2006), using the *makeSimmap()* function of

the R package phytools (Revell 2012). We performed the analysis on a subsample of 1 000 trees from both the posterior distribution of the MrBayes phylograms and the StarBEAST 2 chronograms to explore whether there is any difference between assuming that the phenotypic change is proportional to genetic change or to time, following Kistenich et al. (2018). The MrBayes phylograms were pruned to retain only one sample per species (the one on the shortest branch) and to exclude the outgroup. Outgroup taxa were also excluded from the StarBEAST 2 chronograms. Since the analytical approaches used for obtaining the phylogenetic trees were also different (supermatrix vs supertrees), the resulting topologies may differ as well. Under these circumstances, it may be difficult to know whether any difference in results are caused by the topology or by the scaling of branch lengths. Therefore, in case a contrasting result was found for a given character, a third group of analyses was performed using chronograms generated from the MrBayes phylograms. These chronograms were calculated using penalised likelihoods under a correlated model (Paradis 2013) with the *chronos()* function of the R package ape v. 5.0 (Paradis & Schliep 2019). In all cases, ten simulations were performed for each tree. Otherwise, Kistenich et al. (2018) was followed, including the use of an instantaneous rate matrix, Q, estimated from the empirical data, and a prior distribution of states on the root node estimated from the stationary distribution of Q. However, we summarised the inferred ancestral states at the nodes of interest from all mapped trees, instead of later downsampling them, since the number of initial simulations was smaller. Reconstructions of ancestral states were focused on five nodes in the backbone of the *Dacrymycetes*, i.e., the crown node for the class *Dacrymycetes* and the four crown nodes for each named family. We found a quite obvious artefact in the reconstruction of the number of basidiospore nuclei at the root node (*Dacrymycetes*), which was inferred with 100 % probability as binucleate with the settings indicated above. This problem was caused by an unusual calculation of the Q matrix and was solved by using a symmetric model for that particular character. The problem can also be fixed by sampling Q from the posterior probability distribution of Q using MCMC, but this is much more computationally costly and impractical for large datasets. Following Kistenich et al. (2018), we also checked whether ancestral state reconstructions at or near the root node were influenced by the assumptions made about the distribution of states at the root. The *rayDISC()* function of the corHMM v. 1.22 R package (Beaulieu et al. 2013) was used for this purpose. Reconstructions were made on the pruned majority-rule consensus tree from the MrBayes analyses and on the maximum clade credibility tree from the StarBEAST 2 analysis. The same coding and state characters used for the stochastic mapping were applied. The three different assumptions about the weight given to the states at the root node allowed in *rayDISC* were used for calculating marginal reconstructions: (i) equal weighting among all states; (ii) based on the estimated transition rates (Yang 2006); and (iii) estimated from the data and the model (Maddison et al. 2007, FitzJohn et al. 2009).

Phylogenetic and species delimitation analyses in *Dacryonaema* and *Unilacryma*

The phylogeny of *Dacryonaema* and *Unilacryma* species based on the concatenated dataset was inferred following the same methods indicated for the overall phylogeny, i.e., ML analyses in IQ-TREE v. 1.6.7 and BI in MrBayes v. 3.2.6. The only differences were that we used the compound Dirichlet prior on branch lengths (1, 1.3333, 1, 1) for the Bayesian analysis, based on the different tree length estimates from the best tree of the ML analysis, and the temperature increment parameter was set to 0.2, resulting in good mixing.

For the purpose of delimiting species in *Dacryonaema* and *Unilacryma*, we used the multispecies coalescent approach implemented in STACEY v. 1.2.4 (Jones 2017). Clock and tree model parameters were estimated independently for the nrDNA and each protein coding locus. We used an uncorrelated lognormal relaxed clock model (Drummond et al. 2006) and a Yule tree model. The nrDNA was divided into three partitions, (i) 18S + 5.8S + 28S, (ii) ITS1 + ITS2 and (iii) coded indels, and each protein coding locus was divided in (i) 1st + 2nd codon positions, (ii) the 3rd codon position, and (iii) coded indels for each locus, if present. With this simplified scheme we tried to avoid overpartitioning, since the amount of variation in the *Dacryonaema* + *Unilacryma* dataset was relatively low, particularly the first and second codon positions of the protein coding genes, and parameter estimates when treating those partitions independently were sometimes poor in preliminary analyses. Parameters of the site models were independently estimated for each DNA partition using bModelTest (Bouckaert & Drummond 2017), taking into consideration all transition/transversion split models, while the Lewis MK model with two states was used for the binary partitions. The most visited models were summarised using BModelAnalyser (part of the BEAST 2 package, Bouckaert et al. 2014). We used the CIPRES Science Gateway (Miller et al. 2010) to run ten MCMC parallel analyses for 1×10^9 generations, sampling every 5×10^{4th} tree, and with the collapse height parameter $\epsilon = 10^{-4}$. Considering the amount of missing data (most individuals from GenBank were represented by a single locus) and the taxonomic information from both morphology and the phylogenetic analyses on the concatenated dataset, we used an informative prior for the collapse weight parameter (ω). Since the most likely number of species according to the tree topology and information from morphology was around five or six, and impossible below two (that would imply considering all samples of *Dacryonaema* and *Unilacryma* not only as a single genus, but as a single species), we set $\omega = 0.908$ (highest probability around $\kappa = 5-6$), with 0.5 ($\kappa = 25$) and 0.98 ($\kappa = 2$) as the lower and upper values of the interval, respectively. The first half of each run was discarded as burn-in, and the remaining posterior sample was used to calculate the most likely number of clusters (i.e., putative species), using SpeciesDelimitationAnalyzer (Jones et al. 2014). The matrix of pairwise posterior probabilities of belonging to the same cluster was subsequently visualised and plotted in R (R Core Team 2018) following Jones et al. (2014). The chronogram showing the phylogenetic position of all included samples was drawn with FigTree v. 1.4.

RESULTS

Morphological results

Vacuolar cell contents and its variation with mounting media

The cytoplasm of living cells in all studied species of *Dacrymycetes* was found to be filled with conspicuous, highly refractive vacuolar bodies (VBs) that are variable in size and shape within specimens but not between species (Fig. 3a, c, g, i, m, o, s, u). The VBs disappear when treated with alkaline lethal media, such as KOH, or when cells die by other causes, e.g., desiccation (Fig. 3b, d–f, h, j–l, n, p–r, t, v–x).

Lipid bodies (LBs) have been observed in nearly all studied samples but, unlike VBs, they show considerable variation. Conspicuous LBs are always absent in living cells, being reduced to hardly discernible vacuoles of granular appearance ($\leq 0.2 \mu\text{m}$ diam, unaffected by KOH treatment) due to the presence of the much more conspicuous VBs. When cells die, the LBs become more visible and tend to gradually fuse, forming larger oil drops when the amount of LBs allows so. The difference in

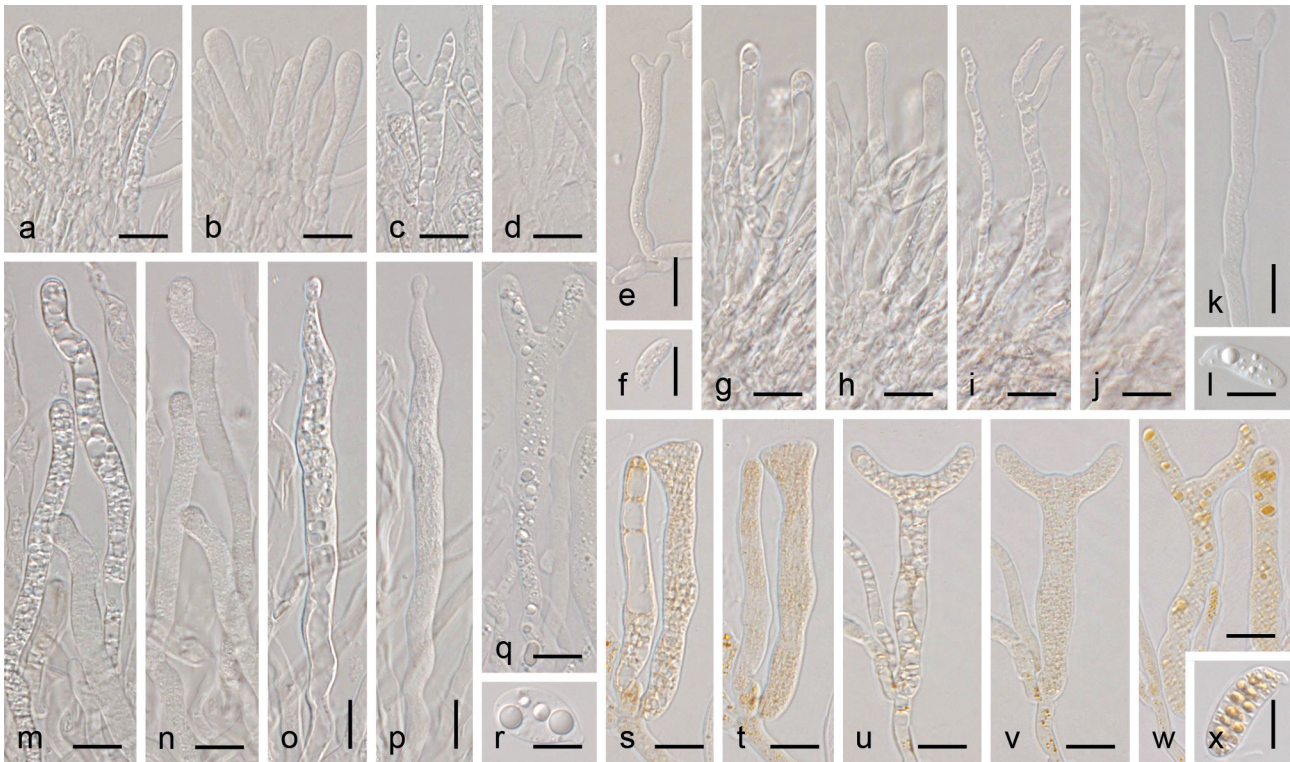


Fig. 3 Vacuolar cell contents in the *Dacrymycetaceae*. a–d, g–j, m–p, s–v. Vacuolar contents in alive (mounted in water) and recently killed (after adding 10 % KOH) young and mature basidia; e–f, k–l, q–r, w–x. vacuolar contents in dead and relatively old basidia and basidiospores after coalescence of the lipid bodies, if present. a–d. *Dacrymyces tortus* s.lat. 4 (UPS F-941021); e–f. *D. tortus* s.lat. 3 (UPS F-941018); g–j. *Dacryonaema rufum* (UPS F-941006); k. *Da. macnabbii* (UPS F-940954); l. *Da. macrosporum* (UPS F-941001); m–p. *Unilacryma unispورا* (UPS F-941286); q–r. *U. bispora* (holotype); s–v. *Dacrymyces chrysocomus* s.lat. 2 (UPS F-940135); w–x. *D. chrysocomus* s.lat. 2 (UPS F-940946). — Scale bars = 10 µm.

the amount of visible cytoplasmic lipid content among groups is more pronounced in basidiospores, less so in basidia, and even less in somatic cells.

In *Cerinomycetaceae*, the cytoplasm of samples treated with KOH appears either as undifferentiated or, at most, as only slightly granular, with an inconspicuous lipid content (Fig. 3b, d–e). Big oil drops (> 2 µm diam) are rarely seen in dead cells of samples of this family (Fig. 3f), neither in rapidly desiccated samples nor in samples exposed to reagents that favour the coalescence of LBs, such as ethanol or HCl. However, exceptions occur, and *D. tortus* s.lat. 1 displayed a rather high lipid content, comparable to some species in the *Dacrymycetaceae*.

In *Dacryonaemataceae*, there is a large variation among the three species (Fig. 3h, j–l). *Dacryonaema rufum* has a comparatively low cytoplasmic lipid content, and LBs are not visible or hardly discernible in dead cells, even after forcing their coalescence (Fig. 3h, j). *Dacryonaema macnabbii* and especially *Da. macrosporum* may show a considerable amount of LBs, and some big lipid drops may form when the LBs coalesce (Fig. 3k–l).

In all studied species of *Dacrymycetaceae* and *Unilacryma-ceae* the cytoplasm of various cells (particularly in basidia and basidiospores) appears clearly granular when samples are mounted in KOH (Fig. 3n, p, t, v). In addition, a few big oil drops (> 2 µm diam) are commonly observed in at least some of the dead cells (Fig. 3q–r, w–x).

Coalesced LBs are often visible in well-preserved herbarium samples, even decades after being collected. However, the cytoplasm tends to become rather uniformly refractive with age in some old herbarium specimens, in which case internal details are lost.

Other variations with mounting media

Cell size and shape displayed little variation among living or dead cells mounted in water and dead cells mounted in KOH (Fig. 3). On the other hand, due to the commonly large size of many structures and the elasticity of the cell walls, dimensions are easily altered by simply applying pressure on the cover slip, or by premature mounting of insufficiently rehydrated specimens. The terminal cells of cortical/marginal hyphae in *Dacryonaema* and a few other genera (e.g., *Heterotextus*) are an exception, since walls become distinctly gelatinised and expanded in KOH.

Hyphae with somewhat roughened walls (e.g., some internal hyphae in various species, Fig. 2d) are often more distinctly roughened when the sections are mounted in KOH, perhaps due to the precipitation of some substances of the intercellular gelatinous matrix.

Nuclei

Nuclei are masked by the VBs in the living state, and are best studied in recently dead cells. In species with highly pigmented cytoplasm, nuclei appear as 'empty' areas due to the absence of pigments, and are relatively easy to observe in fresh samples mounted in KOH (Fig. 4c, e–g, k–m). They are also observable under the same conditions in species with a very low amount of lipid contents (Fig. 4j, n, q). Nuclei can sometimes be seen in slides prepared with Lugol's solution as slightly paler areas of the cytoplasm when the cells are still alive or have recently died (Fig. 4t). Sometimes, they appear as slightly darker areas (due to staining) when the cells are dead. The best contrast is often achieved by using alkaline toluidine blue solutions, where the nuclei (or at least the nucleoli) stain deeply blue, while the cytoplasm stains less intensely and often more purplish, depending on the species and small variations in pH (Fig. 4a–b, d, h–i, o–p, r–s, u–v).

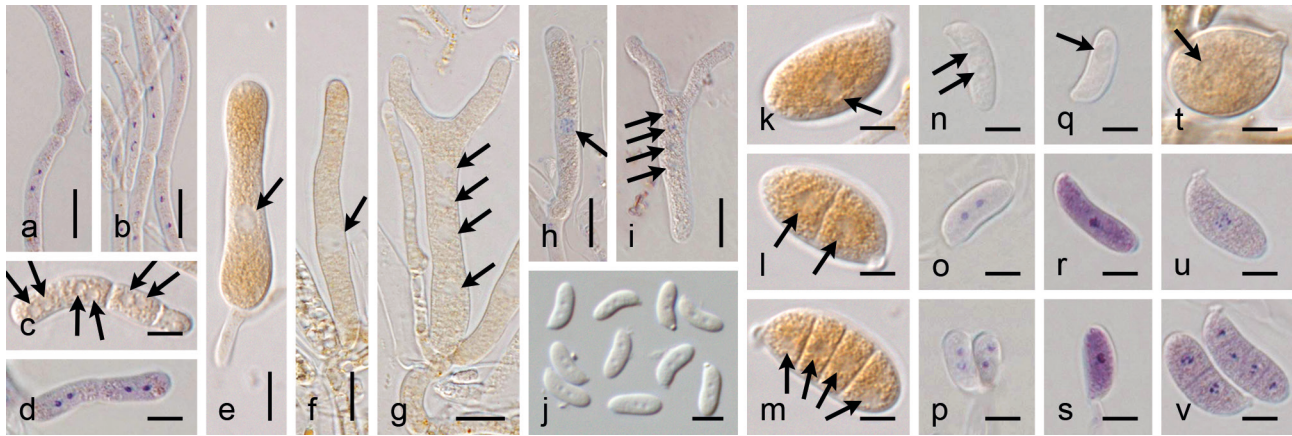


Fig. 4 Nuclei in some cells in the *Dacrymycetaceae*. a–b. Dikaryotic vegetative hyphae; c–d. dikaryotic arthrospores; e–f, h. young basidia showing the single diploid nucleus after cariogamy; g, i. basidia showing the four post-meiotic haploid nuclei; j, n–p. binucleate basidiospores; k–m, j–n, q. in KOH solution; t. basidiospore with a slightly visible (slightly paler) nucleus. a–b, d, h–i, o–p, r–s, u–v. Stained with toluidine blue; c, e–g, j–n, q. in KOH solution; t. stained with Lugol's solution. a–d, h–i, u–v. From *Dacrymyces stillatus* (UPS F-939950); e, k–m. from *D. estonicus* (UPS F-940137); f–g. from *D. chrysocomus* s.lat. (UPS F-940135); j. from *Dacrymyces tortus* s.lat. 3 (UPS F-941016); n–p. from *D. tortus* s.lat. 4 (UPS F-941021); q. from *Dacryonaema rufum* (UPS F-941006); r–s. from *Da. macnabbii* (UPS F-940954); t. from *Unilacryma unispora* (UPS F-941284). — Scale bars: a–b, e–j = 10 μ m, c–d, k–v = 5 μ m.

Vegetative cells in all fresh samples studied are dikaryotic, and mitospores formed directly from those hyphae, e.g., the arthrospores of *Dacrymyces stillatus* are dikaryotic as well (Fig. 4a–d). Basidia under formation are originally dikaryotic, but the two haploid nuclei quickly fuse into a single, large, diploid nucleus (Fig. 4e–f, h). At a certain state of development, often when the sterigmata are already formed, meiosis occurs and sometimes the four postmeiotic nuclei are revealed (Fig. 4g, i), the position of which varies within the mature basidium.

In *Cerinomycetaceae*, young basidiospores constantly have two nuclei, even when still attached to the basidium (Fig. 4j, n–p, observed also in two additional species not included in phylogenetic analyses).

In *Dacrymycetaceae* and *Dacryonaemataceae*, a single nucleus passes into each basidiospore and is not immediately followed by another nuclear division, resulting in young basidiospores that are always uninucleate (Fig. 4k, q–s, u, observed also in c. 15 additional *Dacrymycetaceae* species not included in the phylogenetic analyses). After discharge, they remain uninucleate until mitosis occurs (including septation), and septate basidiospores contain a single nucleus in each daughter cell (Fig. 4l–m, v). This has been clearly observed in basidiospores with up to eight cells. When the number of cells is higher, nuclei are obscured by the narrowness of each individual cell. Scattered binucleate, aseptate basidiospores have been seen in samples where several other basidiospores were already septate, probably representing a developmental stage immediately prior to septation.

In *Unilacrymaceae*, nuclei are very difficult to observe in the basidiospores due to the combination of large volume, high content of lipid bodies, and absence of a colourful cytoplasm (see 'pigments' below). In addition, the differential staining of nuclei with toluidine blue is not as effective as in samples of the other families. Nevertheless, our observations indicate that basidiospores are very likely uninucleate prior to septation (Fig. 4t).

Nuclear details are often lost some time after desiccation, except typically for xerotolerant species (e.g., in *Dacryonaema*), where nuclei can be visible at least in the basidiospores for longer periods after being collected.

Pigments

There is considerable colour variation in the fresh fruitbodies of the four families of the *Dacrymycetaceae*. Fruitbodies in the

Dacrymycetaceae are often yellow or orange (Fig. 5d), although other colours, e.g., brownish or whitish, occur. In the *Cerinomycetaceae*, *Dacryonaemataceae*, and *Unilacrymaceae* duller colours, e.g., pale cream, ochre, brown or grey, are the rule (Fig. 5a–c), although young fruitbodies of some species may be rather orange. The spore print is often yellow to orange in many species of *Dacrymycetaceae* with also yellow to orange fruitbodies (Fig. 5e), pale cream or whitish in paler species and in all samples of *Cerinomycetaceae*, *Dacryonaemataceae* and *Unilacrymaceae* from which we were able to obtain a spore print.

Yellow-orange carotenoid pigments were observed mainly in the lipid bodies of the cytoplasm (Fig. 5f, i–l, p–q, s). In some species, these pigments are only clearly observable after having been concentrated by the coalescence of the lipid bodies and are easily overlooked in living or recently dead cells (Fig. 5p–q). A distinct carotenoid content was observed in most fresh specimens of *Dacrymycetaceae* and also in some herbarium samples. In *Calocera cornea*, the yellowish colour may be particularly inconspicuous and is best seen in the basidium cytoplasm. In *Unilacrymaceae* and *Da. macrosporum*, carotenoids are typically not visible (Fig. 5r), but an inconspicuous yellowish cream tinge (never bright yellow or bright orange) after the fusion of lipid bodies may be seen in fruitbodies growing in very bright places and receiving more sun-light. In *Cerinomycetaceae*, *Da. rufum*, and *Da. macnabbii*, yellow or orange colours were not discernible within the cytoplasm and, as noted above, no big oil drops that could concentrate these pigments are normally formed when the cells die (Fig. 5n–o). These observations do not imply that *Cerinomycetaceae*, *Dacryonaemataceae*, and *Unilacrymaceae* are necessarily devoid of carotenoids, but indicate that the amount of carotenoids, if present, is distinctly lower than in most species of *Dacrymycetaceae*, or of a different composition. The carotenoids progressively degrade after the samples are collected. For this reason, the yellow or orange colours tend to slowly fade in herbarium specimens and may disappear after a few decades.

The carotenoid contents in the studied species of *Dacrymycetaceae* show a distinct greenish to greenish grey reaction with Lugol's solution (Fig. 5g, m), although in *Calocera cornea* it is often inconspicuous (it may be seen on spore prints). This reaction has not been observed in any species of *Cerinomycetaceae*, *Dacryonaemataceae*, or *Unilacrymaceae* (Fig. 5h),

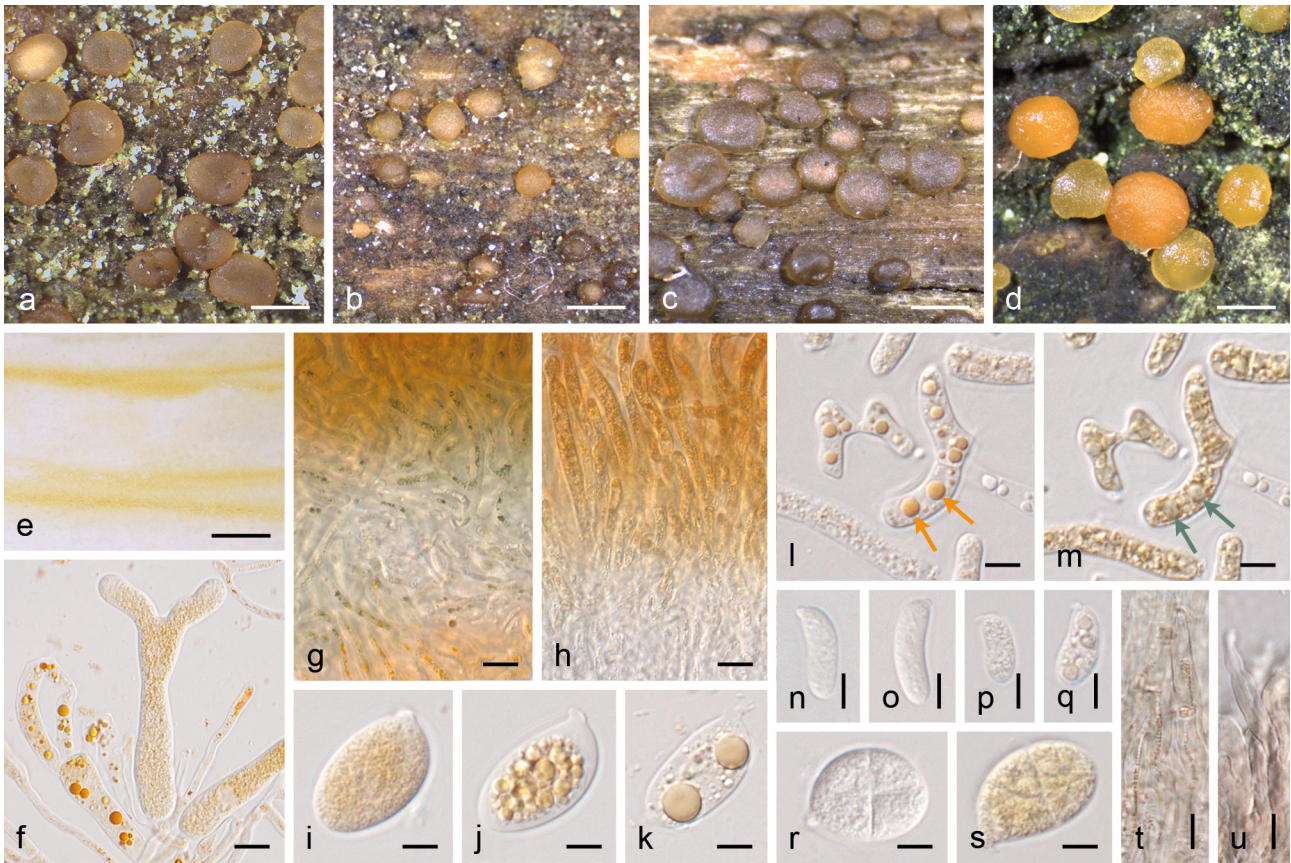


Fig. 5 Pigments in the *Dacrymycetes*. a–d. Representative species with pulvinate and sessile ('dacrymycetoid') fruitbodies in the *Dacryonaemataceae* (a. *Dacryonaema macnabbii*, UPS F-940954), *Unilacrymaceae* (b. *Unilacryma unispورا*, UPS F-941284), *Cerinomycetaceae* (c. *Dacrymyces tortus* s.lat. 3, UPS F-941018), *Dacrymycetaceae* (d. *D. stillatus*, UPS F-939816); e. yellow-orange spore print of *Calocera viscosa* (UPS F-940773); f, i–k. basidia and basidiospores of *D. estonicus* (UPS F-940137); g–h. reaction of the carotenoid contents with Lugol's solution when applied from above to cuttings of *D. estonicus* (g, UPS F-940137) and *U. unispورا* (h, UPS F-941284); i–m. dead arthrospores of *D. stillatus* (UPS F-941285) in water and after the application of Lugol's solution, arrows depict the change in colour of lipid drops with carotenoids; n–o. basidiospores of *Dacryonaema rufum* (n, UPS F-941005) and *Dacrymyces tortus* s.lat. 4 (o, UPS F-941252); p–q. basidiospores of *Calocera viscosa* (UPS F-940773), before (p) and after (q) the coalescence of the lipid bodies; r–s. similarly shaped but differently pigmented basidiospores of *Unilacryma unispورا* (r, UPS F-941280) and *Dacrymyces ovisporus* (s, UPS F-940139); t. cortical/marginal hyphae in *Dacryonaema macrosporum* (UPS F-941000), with brownish intracellular pigments; u. marginal hyphae in *Da. rufum* (UPS F-941005) with diffuse, brownish parietal pigment. — Scale bars: a–e = 1 mm, f–h = 10 μ m, i–u = 5 μ m.

not even when a pale yellowish cream tinge was present in the cytoplasm.

Additional pigments, both intracellular and parietal, are sometimes seen in various species of all four families. An intracellular brownish pigment, seen as small granules in water but partially soluble in KOH, has been noted in samples of all three species of *Dacryonaema*, and it is particularly conspicuous in some marginal cells of *Da. macrosporum* (Fig. 5t). Parietal diffuse pigments were observed in *Dacrymyces tortus* s.lat. and all species of *Dacryonaema* and *Unilacryma*, in *Dacryopinax elegans*, and in a few *Dacrymyces* s.lat. species not included in the present phylogenetic analyses. They are more conspicuous in the sterile marginal cells and may vary from clearly brownish to yellowish brown, brownish orange, or reddish brown. In *Dacryonaema*, they are sometimes better distinguished from the intracellular pigments when mounting samples in KOH (Fig. 5u). These pigments seem to provide at least part of the dark colour in species like *Da. rufum*, and are more persistent than the carotenoids in herbarium specimens. All pigments tend to be more abundant in fruitbodies growing in sun-lit places.

Dacryonaema and *Unilacryma* morphometric analyses

A total of 625 basidiospores were measured (375 for *Dacryonaema* spp. and 250 for *Unilacryma* spp.). ANOVA analyses of the three measured characters (basidiospore width, basidiospore length, and basidiospore Q, Fig. 6) showed significant

differences in basidiospore width and Q in the *Unilacryma* dataset ($p < 0.001$, Cohen's $f > 0.97$), but not in basidiospore length ($p = 0.011$, Cohen's $f = 0.163$), while there were significant differences among the three species of *Dacryonaema* in all three characters ($p < 0.001$, Cohen's $f > 1.66$ for basidiospore length and width, and Cohen's $f = 0.496$ for basidiospore Q). Further analysis by Tukey's HSD test on the *Dacryonaema* dataset found significant differences ($p < 0.001$) in all pairwise comparisons except between *Da. macnabbii* and *Da. rufum* basidiospore length ($p = 0.002$) and *Da. macrosporum* and *Da. rufum* basidiospore Q ($p = 0.0014$). *Dacryonaema macrosporum* has longer and wider basidiospores than *Da. macnabbii* and *Da. rufum*. *Dacryonaema macnabbii* basidiospores are also wider than those of *Da. rufum*. *Dacryonaema macnabbii* has basidiospores tending to be ellipsoid to cylindrical-allantoid, whereas those of *Da. macrosporum* and *Da. rufum* are cylindrical to narrowly cylindrical-allantoid. *Unilacryma bispورا* basidiospores are significantly narrower than *U. unispورا* basidiospores, which is reflected in the significantly different Q, most basidiospores being narrowly ovoid to ellipsoid in *U. bispورا* and subglobose to ovoid in *U. unispورا*.

Molecular results

Molecular data

Counting each of the three parts of the nrDNA as independent sequences, 402 new sequences were generated in this study

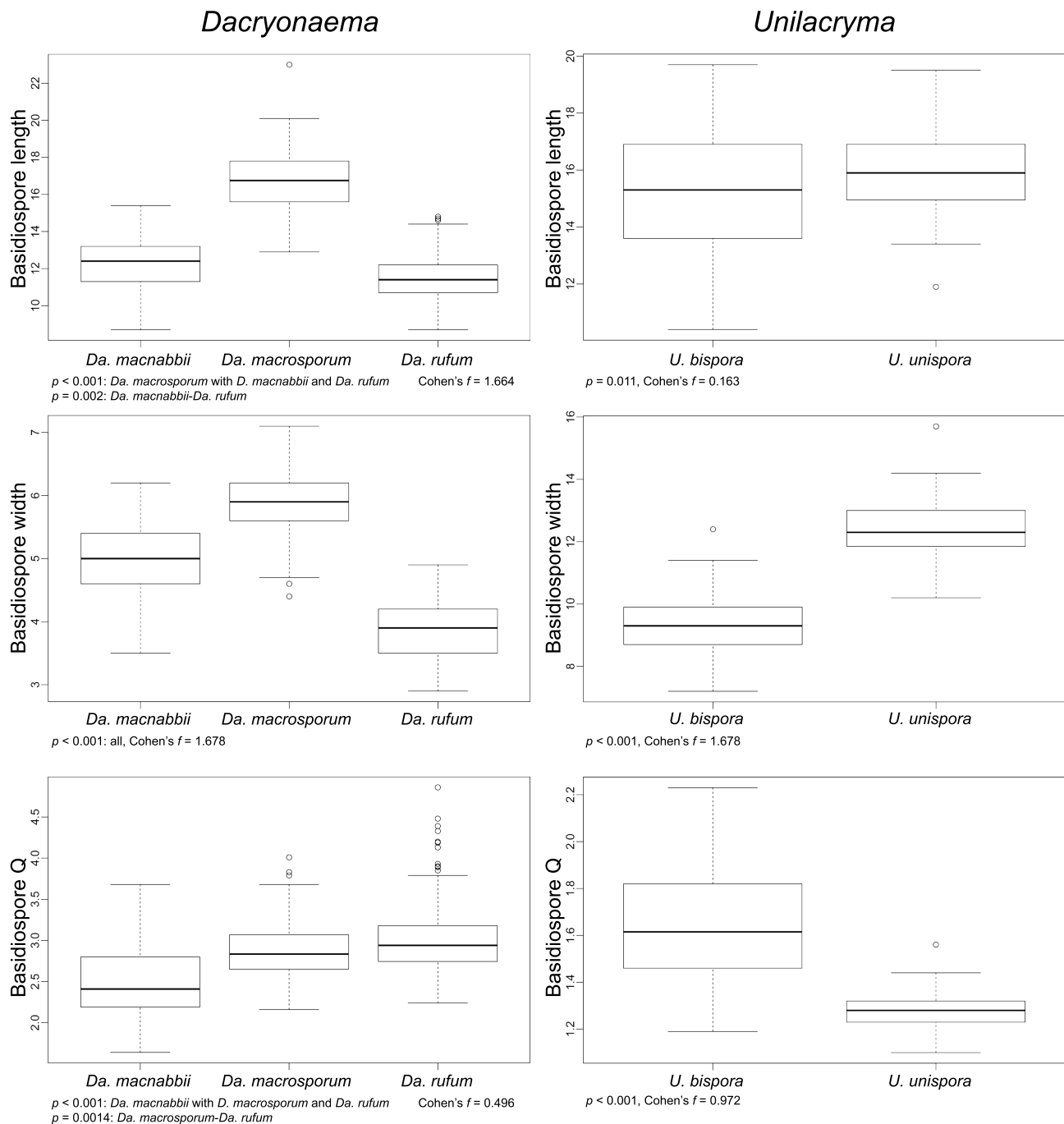


Fig. 6 Morphometric comparisons of *Dacryonaema* spp. and *Unilacryma* spp.

(Table 1). The final dataset for the overall phylogeny included 188 individuals (20 in the outgroup and 168 in the ingroup) that represent around 110 putative species (three in *Tremellomycetes*, 17 in *Agaricomycetes*, and c. 90 in *Dacrymycetes*). The combined matrix consisted of 8890 characters (767 12S, 694 *ATP6*, 1715 18S, 282 ITS1, 171 5.8S, 296 ITS2, 947 28S, 1155 *RPB1*, 1848 *RPB2*, 1015 *TEF-1 α*), of which 4029 were variable and parsimony informative (323 12S, 462 *ATP6*, 392 18S, 158 ITS1, 57 5.8S, 224 ITS2, 389 28S, 644 *RPB1*, 971 *RPB2*, 409 *TEF-1 α*) and 718 variable and parsimony uninformative (90 12S, 61 *ATP6*, 100 18S, 45 ITS1, 10 5.8S, 20 ITS2, 76 28S, 88 *RPB1*, 170 *RPB2*, 58 *TEF-1 α*).

The final dataset for the *Dacryonaemataceae* + *Unilacrymaceae* analyses included 50 terminals. The combined matrix consisted of 7796 characters (638 *ATP6*, 1653 18S, 146 ITS1, 154 5.8S, 292 ITS2, 886 28S, 700 *RPB1* (*Dacryonaema*), 697 *RPB1* (*Unilacryma*), 1620 *RPB2*, 913 *TEF-1 α* , 97 indels), of which 1283 were variable and parsimony informative (163 *ATP6*, 36 18S,

30 ITS1, 8 5.8S, 37 ITS2, 65 28S, 61 *RPB1* (*Dacryonaema*), 154 *RPB1* (*Unilacryma*), 490 *RPB2*, 154 *TEF-1 α* , 85 indels) and 167 variable and parsimony uninformative (0 *ATP6*, 1 18S, 1 ITS1, 0 5.8S, 4 ITS2, 9 28S, 66 *RPB1* (*Dacryonaema*), 6 *RPB1* (*Unilacryma*), 55 *RPB2*, 13 *TEF-1 α* , 12 indels).

No introns were detected in the nrDNA or in the *ATP6* sequences of any of the included species. Five introns were identified in the amplified *RPB1* fragment. All species in *Dacrymycetes* lack the third *RPB1* intron (spliceosomal intron 2 in Matheny et al. 2002), which is consistently present in all species in the outgroup. All species included in the study have a single intron in *RPB2* at domain 7 (*Arabidopsis* intron 16).

In *TEF-1 α* , 12 introns were identified in the alignment, seven of them exclusively present in the *Tremellomycetes*. The second closest intron to the 5'-end is present in some lineages of the *Dacrymycetes* (*Dacryonaemataceae*, *Unilacrymaceae*, *Cerinomycetaceae*, and the *D. chrysocomus* clade), as well as in all included species of *Agaricomycetes* and in *Filobasidium*

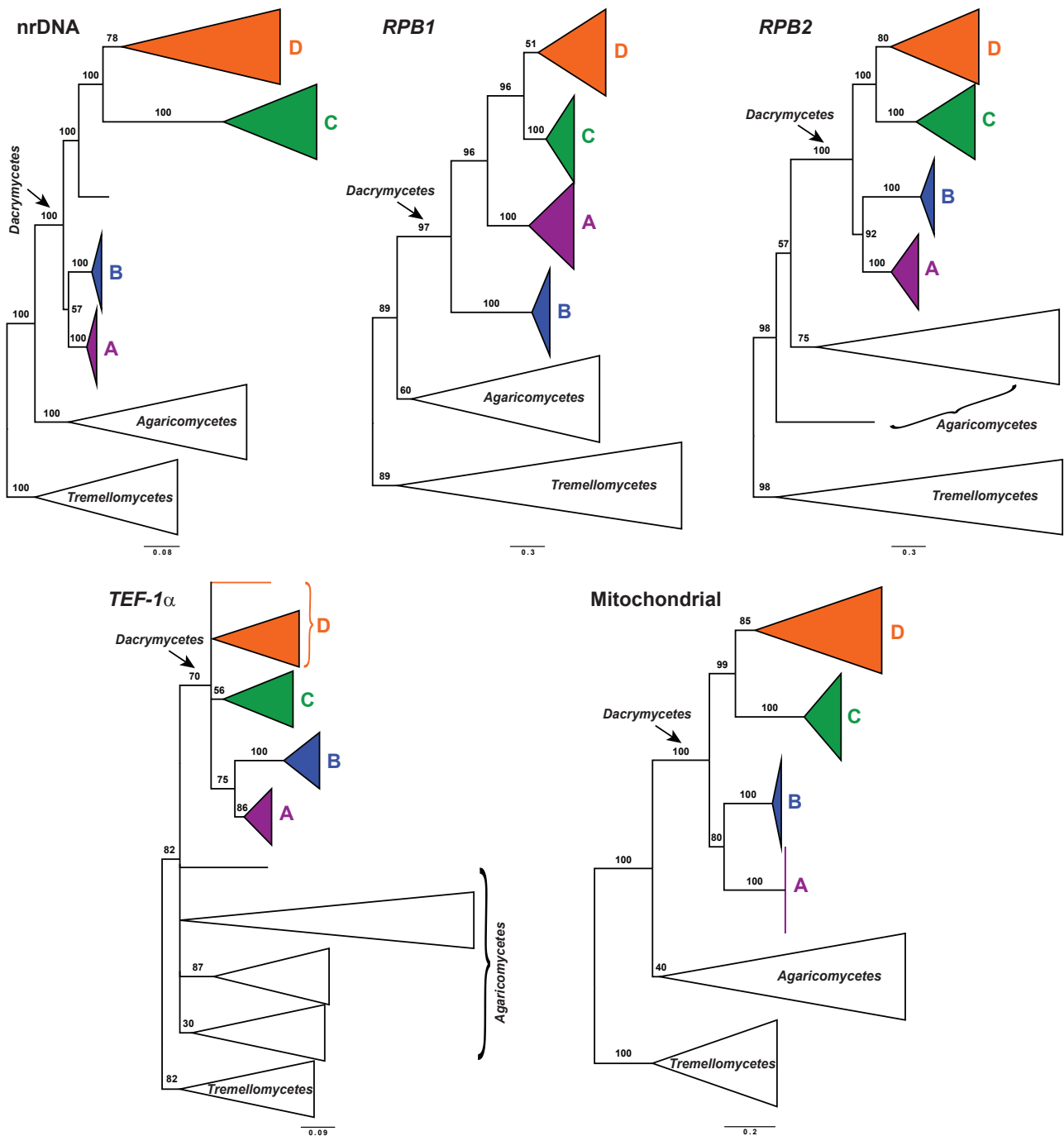


Fig. 7 Maximum likelihood phylogenetic trees from each DNA locus including all samples for the overall phylogeny. A = *Unilacrymaceae*, B = *Dacryonaema*, C = *Cerinomycetaceae*, D = *Dacrymycetaceae*. Numbers on branches indicate ML bootstrap support. Branches with BS ≤ 20 % are collapsed for clarity. Note the incongruent phylogenetic position of the *Dacryonaema* clade in the *RPB1* phylogram, sister to all other *Dacrymycetes* with high support.

wieringae. It is absent in most *Dacrymycetaceae* species and in *Tremella mesenterica* and *Cutaneotrichosporon oleaginosus*. The third intron in *TEF-1α* is present in *Dacryonaemataceae*, *Unilacrymaceae*, and *Cerinomycetaceae*, and absent in all *Dacrymycetaceae*. In the outgroup, it is present in most species except *Plicaturopsis crispa* and *Filobasidium wieringae*. The sixth intron is only present in *Dacryonaemataceae* within the *Dacrymycetes*, and its presence or absence varies in the outgroup, appearing in about 2/3 of the included species.

Three introns were identified in 12S. The first intron is exclusively present in some *Dacrymycetes* species (*Calocera cornea*, *Cerinomyces ceraceus*, *Cerinomyces tortus* s.lat. 1, and *Dacrymyces* cf. *minutus* s.lat. 2) and was located in the motif 'AACAGGAT[I]TWGRKACCCAGG', where '[I]' denotes the intron. The *Vuilleminia comedens* sequence had another exclusive intron (over 1.5 kb) in the motif 'GCAAARBT[I]GAAAWCAAAAAA'.

In both *Calocera cornea* sequences, a third intron was present at the end of the sequence, probably immediately before the primer site for Dacry12S-2r.

Overall phylogeny

Preliminary analyses of all data showed a significant incongruence in the *RPB1* gene tree involving the placement of *Dacryonaema*. *RPB1* data placed *Dacryonaema* as sister to all other *Dacrymycetes* with high support (BS = 96 % for the clade sister to *Dacryonaema* and 100 % for the *Dacryonaema* clade itself), while in all other gene trees, *Dacryonaema* and *Unilacryma* often formed a well-supported clade (BS = 57 % for the nrDNA, 92 % for *RPB2*, 75 % for *TEF-1α*, and 80 % for the mitochondrial dataset), as sister to a generally well-supported clade (99–100 % for the nrDNA, *RPB2*, and the mitochondrial dataset, unsupported for *TEF-1α*) containing the remaining

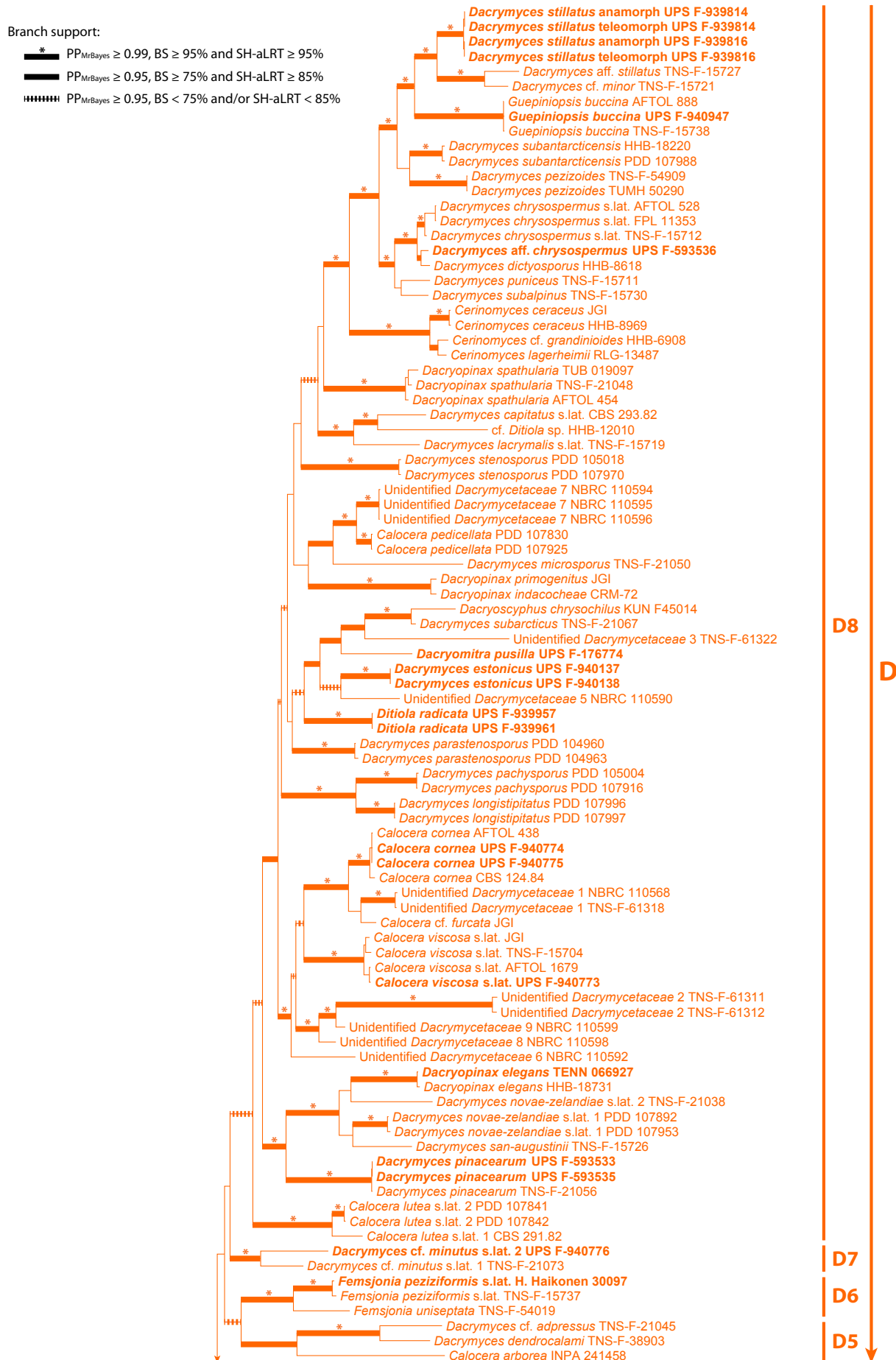


Fig. 8 Fifty percent majority-rule phylogenetic tree of the *Dacrymycetaceae* from the Bayesian analysis, based on the concatenated dataset. Numbers above branches indicate Bayesian posterior probability values (PP_{MrBayes}), and numbers below branches indicate maximum likelihood bootstrap (BS) and Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT) values. Branch support is indicated in the figure for branches above the species level. Samples with newly generated sequences are marked in **bold**. The scale bar represents the average number of substitutions per site. Clade A = *Unilacrymaceae*, clade B = *Dacryonaemataceae*, clade C = *Cerinomycetaceae*, clade D = *Dacrymycetaceae*. Subclades are explained in the text.

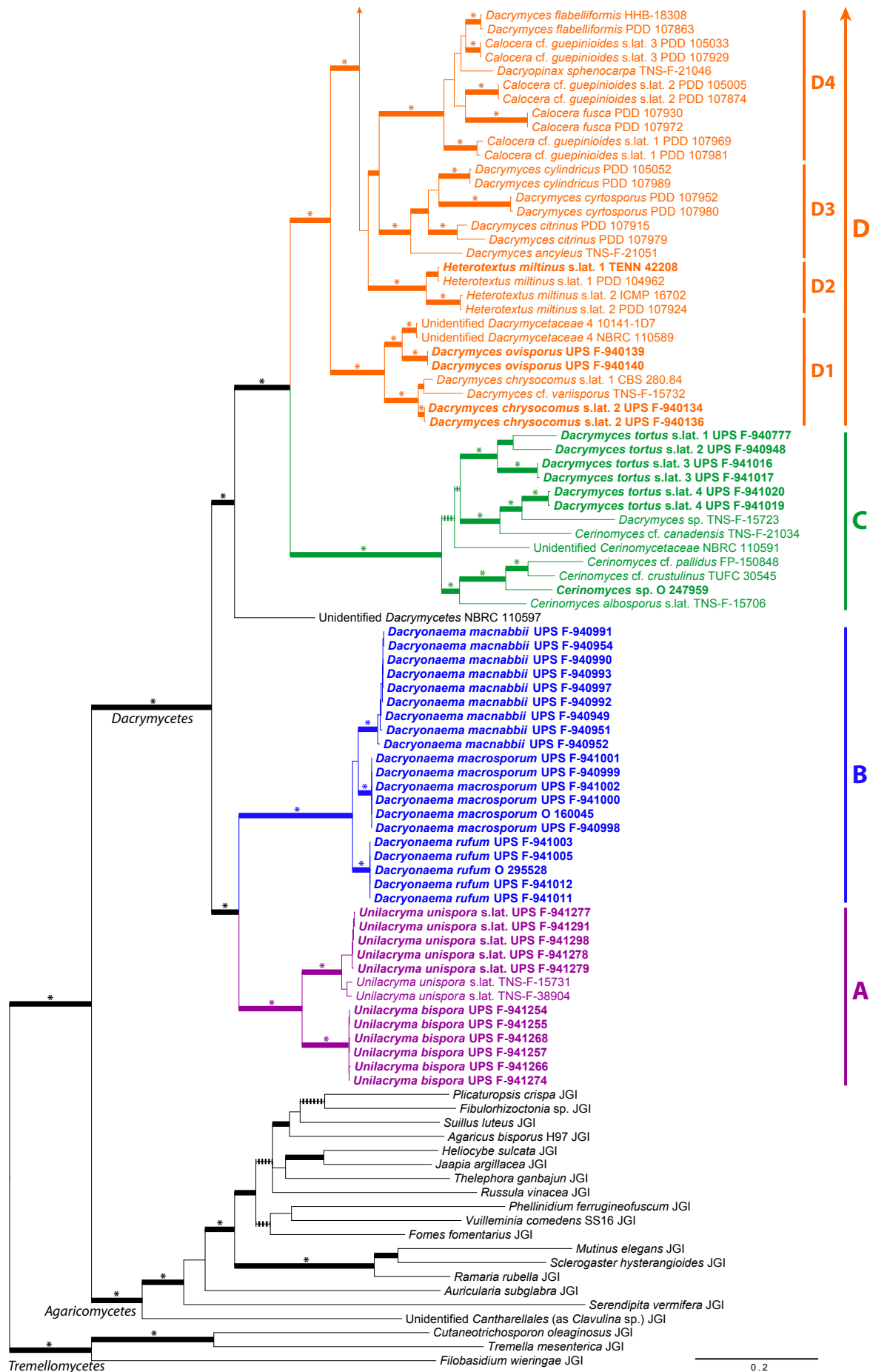


Fig. 8 (cont.)

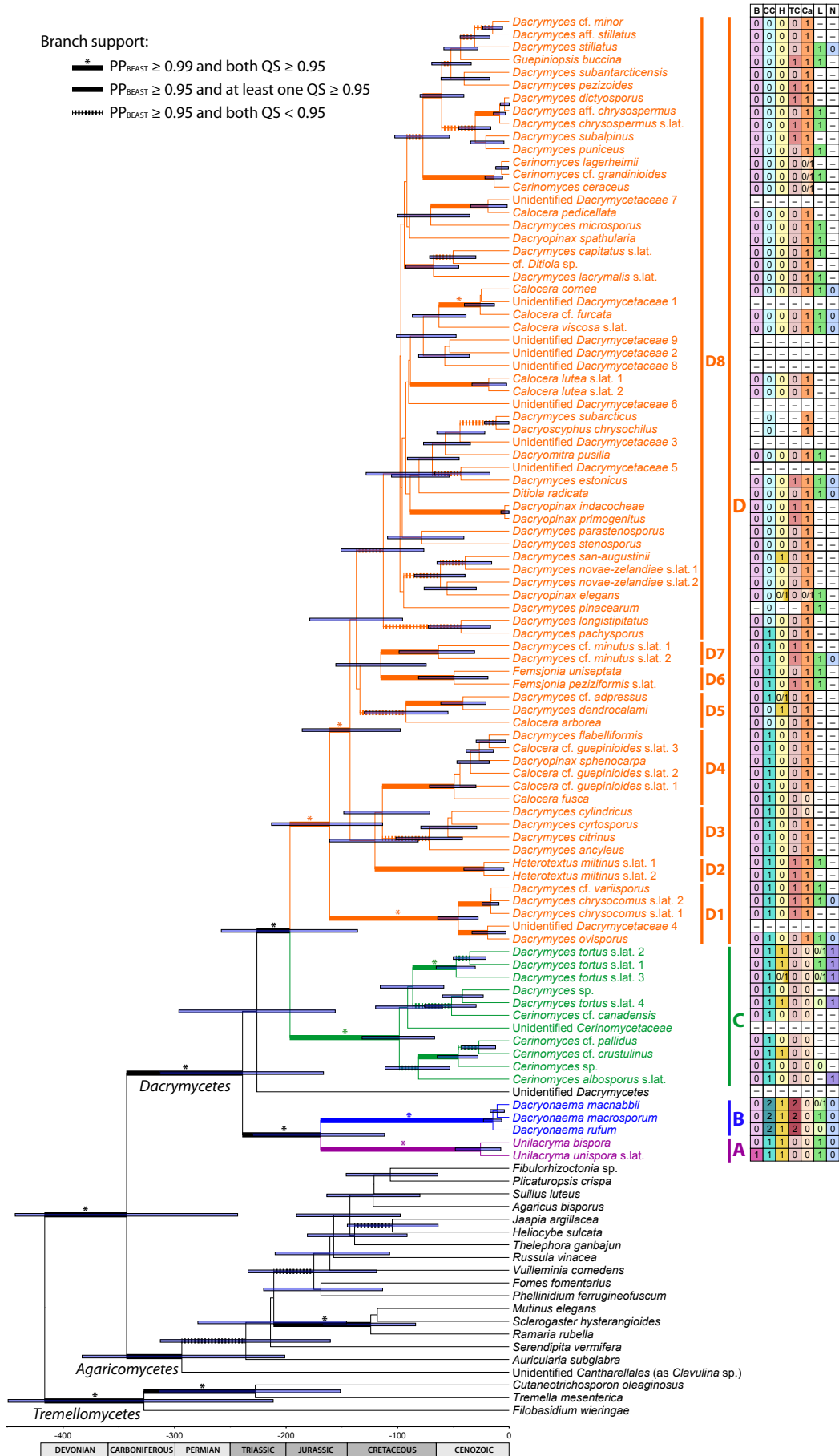


Table 5 Node support, stem ages from the phylogenetic analyses, and median character state probabilities (in percentage) from the SIMMAP analyses on selected clades of interest. PMB = MrBayes phylograms, CMB = MrBayes chronograms, BEAST = StarBEAST 2 chronograms. Probability values $\geq 95\%$ are marked in **bold**.

Clade	Support: BS, SH-aLRT, PP ^{MrBayes} , QS ^{ASTRAL5} , QS ^{ASTRAL9} , PP ^{BEAST}	Mean of stem age (95% HDP interval)	State	Basidium type ¹		Clamp connections ²		Hyphidia ³		Terminal cells of cortical/marginal hyphae ⁴		Carotenoids ⁵		Cytoplasmic lipid content ⁶		Nuclei in young basidiospores ⁷	
				PMB	BEAST	PMB	BEAST	PMB	CMB	BEAST	PMB	BEAST	PMB	BEAST	PMB	BEAST	PMB
<i>Cerinomycetaceae</i>	100, 100,	197	0	99	99	0	0	35	83	100	100	100	71	65	1	1	
	1.00, 0.98,	(136, 258)	1	1	100	100	65	17	0	0	0	0	29	35	99	99	
	1.00, 1.00		2	-	0	0	-	-	0	0	-	-	-	-	-	-	
<i>Dacrymycetaceae</i>	100, 100,	197	0	99	99	0	0	96	86	100	96	2	1	3	100	100	
	1.00, 0.99,	(136, 258)	1	1	100	100	4	14	0	7	4	98	97	99	97	0	
	1.00, 1.00		2	-	0	0	-	-	0	0	-	-	-	-	-	-	
<i>Dacrymycetes</i>	100, 100,	343	0	100	100	0	0	67	87	98	74	92	1	0	99	98	
	1.00, 0.99,	(243, 443)	1	0	98	98	33	13	2	25	8	16	99	100	1	2	
	1.00, 1.00		2	-	2	2	-	-	1	1	-	-	-	-	-	-	
<i>Dacryonaemataceae</i>	100, 100,	169	0	100	99	0	0	0	18	0	0	0	33	42	100	100	
	1.00, 0.98,	(112, 230)	1	0	1	0	0	100	82	100	0	0	67	58	0	0	
	1.00, 1.00		2	-	100	100	-	-	100	100	-	-	-	-	-	-	
<i>Unilacrymaceae</i>	100, 100,	169	0	81	87	0	0	1	2	0	97	100	1	1	100	100	
	1.00, 0.95,	(112, 230)	1	19	13	100	100	99	98	100	3	0	0	0	0	0	
	1.00, 1.00		2	-	0	0	-	-	0	0	-	-	-	-	-	-	

¹ 0 = 2-spored; 1 = 1-spored.
² 0 = absent; 1 = at most septa and short; 2 = subhyphal and long.
³ 0 = simple or absent; 1 = branched.
⁴ 0 = filamentose, simple; 1 = swollen, simple; 2 = filamentose, branched.
⁵ 0 = indistinct; 1 = distinct.
⁶ 0 = no visible lipid drops; 1 = visible lipid drops.
⁷ 0 = one nucleus; 1 = two nuclei.

Dacrymycetes (Fig. 7). We did not find any evidence of heterogeneous nucleotide composition, recombination, or heterotachy that could explain the topological differences. One possibility is that an *RPB1* paralogue was sequenced in *Dacryonaema*. The *RPB1* sequence of one sample of *Dacryonaema macnabii* (UPS F-940952), amplified and sequenced twice, contained at its end substitutions that resulted in codon positions coding for unique amino acids, and a sequence from another sample (UPS F-940954) had a single base indel towards the end (both strands sequenced and no double signal noted), affecting the open reading frame. These problems were not noted in any of the other *Dacrymycetes RPB1* sequences. Excluding suspicious parts of the alignment did not change the topological incongruence of the phylogeny. We consider it unlikely that the incongruence was caused by incomplete lineage sorting, since it affected very deep and comparatively long branches in the backbone, while relationships among closely related species were congruent. No other cases of incongruence were detected.

The best partitioning scheme and models for each partition in the final concatenated ML analysis were: (i) 12S + *ATP6* 1st, GTR + I + Γ 4, (ii) *ATP6* 2nd, GTR + Γ 4, (iii) *ATP6* 3rd, TIM + Γ 4, (iv) 18S, TIMe + I + Γ 4, (v) ITS1, TIM2 + I + Γ 4, (vi) ITS2, TPM2 + F + I + Γ 4, (vii) 5.8S + 28S, TN93 + I + Γ 4, (viii) *RPB1* 1st + *RPB2* 1st, GTR + I + Γ 4, (ix) *RPB1* 2nd + *RPB2* 2nd, GTR + I + Γ 4, (x) *RPB1* 3rd + *RPB2* 3rd, GTR + I + Γ 4, (xi) *TEF-1 α* 1st, GTR + I + Γ 4, (xii) *TEF-1 α* 2nd, TVM + I + Γ 4, and (xiii) *TEF-1 α* 3rd, GTR + Γ 4. The two ML tree replicates had a similar topology, with lnL = -157377.647 for the best one. The concatenated Bayesian analysis finished after 1.8×10^7 generations (ASDF < 0.01); all parameters had a post-burn-in ESS > 1000, all PSRFs = 1.000 \pm 0.007 (most of them 1.000 \pm 0.002), and RWTY plots did not show convergence problems with the tree topologies. The topology of the 50 % majority-rule consensus tree was similar to that of the ML trees, and thus only the Bayesian tree with PP, BS and SH-aLRT values is shown in Fig. 8.

The final normalised quartet score in ASTRAL III was 0.983 in the five-locus analysis and 0.955 in the nine-locus analysis, indicating very low discordance among the single-locus trees. The trees computed from each of those two analyses were nearly identical. Only three runs reached convergence in the StarBEAST 2 analysis. Nearly every parameter had an ESS > 200 (often > 1000), including species tree parameters (posterior, likelihood, tree height, tree length), except the cySpeciationRate, which did not fully converge (100 < ESS < 200, but see Discussion). RWTY plots did not show convergence problems among the species tree topologies. The ASTRAL III and StarBEAST 2 species trees were very similar, with minor differences only in unsupported groups. Thus, only the species tree from StarBEAST 2 with QS values from ASTRAL and PP values from StarBEAST analyses is shown in Fig. 9.

In all analyses, the *Tremellomycetes*, *Agaricomycetes* and *Dacrymycetes* were resolved as monophyletic groups and generally received high support (BS = 100 %, SH-aLRT = 100 %, PP^{MrBayes} = 1.00, QS^{ASTRAL5} = 0.99, QS^{ASTRAL9} = 0.99–1.00, PP^{BEAST} = 1.00, except the *Agaricomycetes* in the ASTRAL III analysis assuming five loci, with QS^{ASTRAL5} = 0.69). The *Dacrymycetes* and the *Agaricomycetes* are sister groups and form a clade with high support (BS = 100 %, SH-aLRT = 100 %, PP^{MrBayes} = 1.00, QS^{ASTRAL5} = 0.99, QS^{ASTRAL9} = 1.00, PP^{BEAST} = 1.00). Within *Dacrymycetes*, we distinguish five main groups: (i) *Unilacrymaceae* (clade A), comprising *Unilacryma unisporea* s.lat. and an unnamed *Unilacryma* with bisporic basidia; (ii) genus *Dacryonaema* (clade B), with three species; (iii) an isolated sample from the culture NBRC 110597; (iv) *Cerinomycetaceae* (clade C, several species); and (v) *Dacrymycetaceae* (clade D), including several genera and the majority of species. Support values and stem ages for the mentioned clades are summarised in Table 5.

Table 6 Character state probabilities (in percentage) from the *rayDisc* analyses on selected clades of interest. PMB = MrBayes phylograms, CMB = MrBayes phylograms, BEAST = StarBEAST 2 chronograms. Results are given ordered according to the following assumptions about the distribution of states at the root: equal distribution of states/estimated from the data and the model/based on the transition rates. Probability values $\geq 95\%$ are marked in **bold**. Clade 1 = *Dacrymycetes*, clade 2 = *Dacryonaemataceae*, clade 3 = *Unilacrymycetaceae*, clade 4 = *Cerionomycetaceae*, clade 5 = *Dacrymycetaceae*.

Clade	State	Basidium type ¹		Clamp connections ²		Hyphidia ³		Terminal cells of cortical/marginal hyphae ⁴		Carotenoids ⁵		Cytoplasmic lipid content ⁶		Nuclei in young basidiospores ⁷	
		PMB	BEAST	PMB	BEAST	PMB	CMB	BEAST	BEAST	PMB	BEAST	PMB	BEAST	PMB	BEAST
1	0	0/100/100	32/50/100	0/0/0	0/0/0	4/0/59	50/50/91	80/99/69	74/97/60	96/100/96	91/100/90	63/100/2	100/100/4	100/100/100	100/100/98
1	1	100/0/0	68/50/0	98/100/100	98/100/100	96/100/41	50/50/9	20/1/31	25/3/40	4/0/4	9/0/10	37/0/98	0/0/96	0/0/0	0/0/2
2	2	-	-	2/0/0	2/0/0	-	-	0/0/0	1/0/0	-	-	-	-	-	-
2	0	99/100/100	100/100/100	0/0/0	0/0/0	0/0/0	1/1/1	0/0/0	0/0/0	100/100/100	100/100/100	66/100/42	100/100/46	100/100/100	100/100/100
1	1	1/0/0	0/0/0	0/0/0	0/0/0	100/100/100	99/99/99	0/0/0	0/0/0	0/0/0	0/0/0	34/0/58	0/0/54	0/0/0	0/0/0
2	2	-	-	100/100/100	100/100/100	100/100/100	100/100/100	100/100/100	100/100/100	-	-	-	-	-	-
3	0	0/96/72	28/89/89	0/0/0	0/0/0	0/0/1	1/1/1	99/99/99	97/98/96	100/100/100	100/100/100	8/1/2/2	1/1/1	100/100/100	100/100/100
1	1	100/4/28	72/11/11	100/100/100	100/100/100	100/100/99	99/99/99	3/2/4	0/0/0	0/0/0	0/0/0	92/88/98	99/99/99	0/0/0	0/0/0
2	2	-	-	0/0/0	0/0/0	0/0/0	0/0/0	0/0/0	0/0/0	-	-	-	-	-	-
4	0	99/100/99	99/99/99	0/0/0	0/0/0	20/18/34	61/61/76	100/100/100	100/100/100	100/100/100	100/100/100	85/100/74	100/100/72	1/1/1	1/1/1
1	1	1/0/1	1/1/1	100/100/100	100/100/100	80/82/66	39/39/24	0/0/0	0/0/0	0/0/0	0/0/0	15/0/26	0/0/28	99/99/99	99/99/99
2	2	-	-	0/0/0	0/0/0	-	-	0/0/0	0/0/0	-	-	-	-	-	-
5	0	95/100/99	94/99/99	0/0/0	0/0/0	90/90/94	57/57/76	93/98/91	93/98/91	2/3/1	4/5/2	9/1/7/3	25/25/7	100/100/100	100/100/100
1	1	5/0/1	6/1/1	100/100/100	100/100/100	10/10/6	43/43/24	7/4/9	7/4/9	98/97/99	96/95/98	91/83/97	75/75/93	0/0/0	0/0/0
2	2	-	-	0/0/0	0/0/0	-	-	0/0/0	0/0/0	-	-	-	-	-	-

¹ 0 = 2-spored; 1 = 1-spored.
² 0 = absent; 1 = at most septa and short; 2 = subhymenial and long.
³ 0 = simple or absent; 1 = branched.
⁴ 0 = filamentose, simple; 1 = swollen, simple; 2 = filamentose, branched.
⁵ 0 = indistinct; 1 = distinct.
⁶ 0 = no visible lipid drops; 1 = visible lipid drops.
⁷ 0 = one nucleus; 1 = two nuclei.

Table 7 Median transformation counts (and 95% HDP interval) of ancestral states for the overall phylogeny of the *Dacrymycetes*. PMB = MrBayes phylograms, CMB = MrBayes phylograms, BEAST = StarBEAST 2 chronograms.

State change	Basidium type ¹	Clamp connections ²		Hyphidia ³		Terminal cells of cortical/marginal hyphae ⁴		Carotenoids ⁵		Cytoplasmic lipid content ⁶		Nuclei in young basidiospores ⁷	
		PMB	BEAST	PMB	BEAST	PMB	BEAST	PMB	BEAST	PMB	BEAST	PMB	BEAST
0→1	3 (1–5)	7 (1–23)	2 (1–3)	2 (0–3)	6 (3–10)	15 (6–25)	7 (6–10)	14 (10–18)	1 (0–2)	4 (1–9)	5 (0–10)	3 (1–6)	3 (1–6)
0→2	-	-	0 (0)	0 (0)	-	-	-	0 (0)	-	-	-	-	-
1→0	2 (0–6)	7 (0–24)	3 (2–4)	3 (1–5)	6 (0–10)	12 (2–23)	0 (0)	4 (0–9)	3 (2–5)	6 (3–10)	7 (3–11)	0 (0–2)	0 (0–2)
1→2	-	-	1 (1–2)	1 (1–2)	-	-	-	1 (1–2)	-	-	-	-	-
2→0	-	-	0 (0)	0 (0)	-	-	-	0 (0)	-	-	-	-	-
2→1	-	-	0 (0)	0 (0–1)	-	-	-	0 (0–1)	-	-	-	-	-

¹ 0 = 2-spored; 1 = 1-spored.
² 0 = absent; 1 = at most septa and short; 2 = subhymenial and long.
³ 0 = simple or absent; 1 = branched.
⁴ 0 = filamentose, simple; 1 = swollen, simple; 2 = filamentose, branched.
⁵ 0 = indistinct; 1 = distinct.
⁶ 0 = no visible lipid drops; 1 = visible lipid drops.
⁷ 0 = one nucleus; 1 = two nuclei.

Dacryonaema and *Unilacrymaceae* are sister groups forming a clade with high support (BS = 100 %, SH-aLRT = 99 %, PP_{MrBayes} = 1.00, QS_{ASTRAL5} = 0.98, QS_{ASTRAL9} = 0.99, PP_{BEAST} = 1.00), and so are *Cerinomycetaceae* and *Dacrymycetaceae* in most analyses (BS = 100 %, SH-aLRT = 100 %, PP_{MrBayes} = 1.00, QS_{ASTRAL5} = 0.67, QS_{ASTRAL9} = 0.99, PP_{BEAST} = 1.00). The clade formed by the sample NBRC 110957 + *Cerinomycetaceae* + *Dacrymycetaceae* is highly supported in the analyses using the concatenated dataset but not in two of the coalescence-based analyses (BS = 99 %, SH-aLRT = 96 %, PP_{MrBayes} = 1.00, QS_{ASTRAL5} = 0.67, QS_{ASTRAL9} = 0.95, PP_{BEAST} = 0.86). The nodes in the *Dacrymycetes* backbone, uniting groups i–v, are consistent among all analyses (Fig. 8–9).

Several internal subclades can be identified in the *Dacrymycetaceae*, and we highlight eight of them with taxonomic or nomenclatural relevance: D1–D8. All are well-supported except D8 that received high support only from SH-aLRT (100 %), PP_{MrBayes} (1.00), and PP_{BEAST} (1.00), and D3 and D5 that received high support in all analyses except ASTRAL III. For details on the included species and morphological characterization of these subclades, see Discussion.

Character evolution

Posterior probabilities of character states (pp) from the SIMMAP simulations at selected nodes were similar between the MrBayes phylograms and the StarBEAST 2 chronograms, except for the hyphidia (Table 5). The ancestor of the *Dacrymycetes* was inferred to have simple hyphidia with 67 % pp in the analysis using MrBayes phylograms, 87 % pp based on MrBayes chronograms, and 97 % pp when using Star BEAST 2 chronograms. The ancestor of *Cerinomycetaceae* was inferred to have branched hyphidia based on MrBayes phylograms (65 % pp), while using chronograms from MrBayes and StarBEAST 2 simple hyphidia was the most probable state, with 83 % and 98 % pp, respectively.

The ancestor of the *Dacrymycetes* was inferred to have 2-spored basidia, hyphae with short clamp connections at most septa, not visible carotenoids, conspicuous cytoplasmic lipid contents, and uninucleate young basidiospores (84–100 % pp). The ancestor of *Dacryonaemataceae* had 2-spored basidia, hyphae with long clamp connections mostly at the subhymenium and the bases of basidia and hyphidia, branched hyphidia, cortical/marginal hyphae with branched, and undilated terminal cells, no visible carotenoids, and uninucleate young basidiospores (83–100 % pp). The ancestor of *Unilacrymaceae* had 2-spored basidia, hyphae with short clamp connections at most septa, branched hyphidia, simple and undilated terminal cells of cortical/marginal hyphae, invisible carotenoids, conspicuous cytoplasmic contents, and uninucleate young basidiospores (81–100 % pp). The ancestor of *Cerinomycetaceae* had 2-spored basidia, hyphae with short clamp connections at the majority of septa, cortical/marginal hyphae with simple and undilated terminal cells, invisible carotenoids, and binucleate young basidiospores (99–100 % pp). Finally, the ancestor of *Dacrymycetaceae* had 2-spored basidia, hyphae with short clamp connections at most septa, hyphidia simple or absent, cortical/marginal hyphae with simple and undilated terminal cells, visible carotenoids, conspicuous cytoplasmic lipid contents, and uninucleate young basidiospores (86–100 % pp).

RayDISC results (Table 6) largely agreed with those from SIMMAP, including the different inferences for the hyphidia when a different tree was used. However, state reconstructions for the basidium type and the amount of lipid content varied considerably at the root node (ancestor of the *Dacrymycetes*) depending on the assumed distribution of states at that node. A 1-spored basidium was inferred as the ancestral state with 100 % pp assuming an equal distribution of states in MrBayes

phylograms, while a 2-spored basidium was inferred with 100 % pp both when the distribution of states is calculated using the estimated transition rates (both trees) and when it is estimated from the data and the model in MrBayes phylograms. The amount of cytoplasmic lipid content is inferred as high when the distribution of states is calculated using the estimated transition rates (96–98 % pp), while it is inferred as low when it is estimated from the data and the model (100 % pp in both trees) and also assuming an equal distribution of states for the BEAST chronograms (100 % pp).

The median transformation counts (Table 7) show that the 1-spored basidium seems to be a rarely acquired apomorphic state, probably as a reduction of the 2-spored basidium, although there is a substantial uncertainty in the counts from the chronograms. Clamp connections were lost only in a couple of internal groups in the *Dacrymycetaceae* (remarkably in subclade D8, Fig. 9), and reversions to the state with clamp connections were uncommon. The hyphidium branching seems to have increased in most clades, although the acquisition of branched hyphidia was rare in *Dacrymycetaceae*. The terminal cells of cortical/marginal hyphae showed some flexibility, as the transformation counts show numerous transitions from simple and not swollen cells into more or less conspicuously dilated cells in the *Dacrymycetaceae*. Some reversions to the ancestral state were also inferred. The branched and not swollen cells most probably appeared only once (in the *Dacryonaemataceae*). The carotenoids were most likely acquired only once, by the ancestor of the *Dacrymycetaceae*, with a few secondary losses. The visible lipid content of the cytoplasm in basidia and basidiospores was reduced a few times (e.g., *Dacryonaema rufum* and some species in the *Cerinomycetaceae*), although increases in visibility also occurred. The binucleate young basidiospore state is an acquired state in the *Cerinomycetaceae*.

Dacryonaema and *Unilacryma* phylogeny and species delimitation analyses

No incongruence among loci were detected. Both ML and Bayesian analyses based on the concatenated dataset resulted in trees with a similar topology. The best partitioning scheme and models for each partition in the ML analysis were: (i) 18S + 5.8S + *TEF-1α* 2nd, K2P + I; (ii) ITS1 + *RPB1* (*Dacryonaema*) 1st + *RPB1* (*Unilacryma*) 1st + *RPB2* 1st, TIM2 + I; (iii) ITS2, K2P; (iv) 28S + *TEF-1α* 1st, TN93 + I; (v) *RPB1* (*Dacryonaema*) 2nd + *RPB1* (*Unilacryma*) 2nd + *RPB2* 2nd, TPM3 + F; (vi) *RPB1* (*Dacryonaema*) 3rd + *RPB1* (*Unilacryma*) 3rd + *RPB2* 3rd, HKY + Γ4; (vii) *TEF-1α* 3rd, TPM2 + F + Γ4; (viii) *ATP6* first codon position (1st), TN93 + I; (ix) *ATP6* second codon position (2nd), TPM3u + F + I; (x) *ATP6* third codon position (3rd), TIM3; and (xi) indels, MK + ASC. Both ML replicates yielded trees with a nearly identical topology (minor changes in a few unsupported branches within species), with $\ln L_1 = -18219.068$ and $\ln L_2 = -18219.016$ (best).

The MrBayes analysis was automatically halted after 8×10^6 generations, when the ASDSF dropped below 0.01. At this point, all ESSs after discarding the burn-in were > 1000, and all tree and model parameters had PSRFs = 1.000 ± 0.001 .

The STACEY analysis yielded a chronogram with a similar topology to the phylograms obtained using the concatenated dataset. Thus, only the STACEY chronogram with support values from all analyses is shown in Fig. 10. According to SpeciesDelimitationAnalyzer, the most probable delimitation involves six species (40.7 % posterior probability). All other delimitations had a posterior probability of < 5 %.

Dacryonaemataceae and *Unilacrymaceae* were resolved as monophyletic, highly supported groups. Within the *Dacryonaemataceae*, three fully or nearly fully supported main clades are resolved (BS = 100 %, SH-aLRT = 99–100 %, PP_{MrBayes} = 1.00,

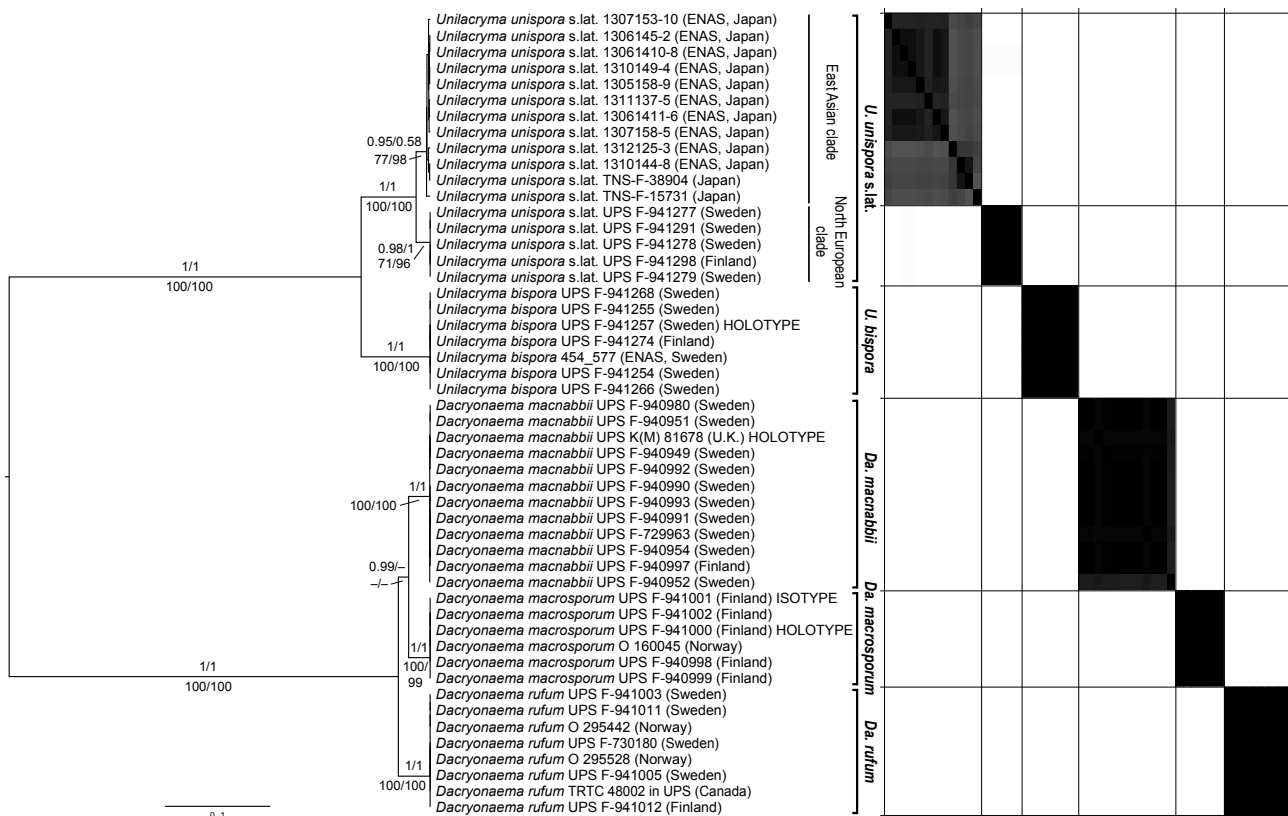


Fig. 10 STACEY species delimitation analysis with chronogram and similarity matrix for the *Dacryonaema* + *Unilacryma* dataset. Numbers above branches indicate Bayesian posterior probability ($PP_{STACEY}/PP_{MrBayes}$) values, and numbers below branches indicate maximum likelihood bootstrap (BS) and Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT) values.

$PP_{STACEY} = 1.00$, and the same clades are inferred as putative species (*Da. macnabii*, *Da. macrosporum*, and *Da. rufum*) in the STACEY species delimitation analysis. The posterior probability among specimens to belong to the same inferred group was high according to the similarity matrix ($pp > 0.88$). The posterior probability that any of the samples in one of the three putative species may belong to any of the other two putative species was zero.

Within *Unilacrymaceae*, two fully supported clades were resolved, representing *U. bisporea* and *U. unisporea* s.lat. The posterior probability of all specimens of *U. bisporea* belonging to the same cluster inferred by STACEY was high ($pp > 0.99$), and the probability that any of the samples belong to any of the *U. unisporea* s.lat. groups was zero. The clade formed by *U. unisporea* s.lat. samples is further divided into two subclades, one including all North European samples (BS = 71 %, SH-aLRT = 96 %, $PP_{MrBayes} = 1.00$, $PP_{STACEY} = 0.98$) and the other including all East Asian samples (BS = 77 %, SH-aLRT = 98 %, $PP_{MrBayes} = 0.58$, $PP_{STACEY} = 0.95$). The STACEY analysis inferred these subclades as two putative species. Within the North European group, all specimens had high posterior probability of belonging to the same group ($pp > 0.99$). Within the East Asian group, the posterior probability of each sample belonging to the same group was highly variable ($0.67 < pp < 0.94$). The posterior probability of North European and Japanese samples belonging to the same cluster was low ($pp < 0.02$).

DISCUSSION

Phenotypic characters

Cytology as a source of taxonomic characters

Vacuolar cell contents have been rarely noted in *Dacrymycetes* by other researchers, and there are scarce records of them in

the literature. For instance, they were probably drawn by Nannfeldt (1947) for some basidiospores of *Dacryonaema rufum* (e.g., his f. 2 showing vacuolated ungerminated basidiospores after 5 days in water). Also, the photograph for explaining the ontogeny of basidia in *Dacrymyces stillatus* included by Oberwinkler (2014: f. 3 of that study) showed conspicuous VBs in some basidia. Unfortunately, we did not see any taxonomically relevant variation of the VBs in the *Dacrymycetes*.

The amount of lipid contents, routinely used as a taxonomic character in several groups of *Ascomycota* (Baral 1992), has been poorly explored in the *Basidiomycota*. The higher lipid content in the cytoplasm of *Dacrymycetaceae* and *Unilacrymaceae* compared to *Cerinomycetaceae* seems useful for identifying the latter family, although *D. tortus* s.lat. 1 is a clear exception in which conspicuous lipid drops are easily observed. This species might be confused with others in the *Dacrymycetaceae*, especially as taxa in the *D. tortus* complex also have dacrymycetoid fruitbodies. They rarely coalesce and may have obvious orange tinges when young. The basidia are sometimes comparatively long, in which case their dimensions and shape overlap with those of some *Dacrymycetaceae*.

There are not many studies focused on the nuclear stages in the *Dacrymycetes*. The first mentions appeared in Dangeard (1895), Istvanffi (1895), Juel (1898), and Maire (1902), who noted that a single nucleus passes into each young basidiospore when attached to the basidium. Bodman (1938) carefully explained the karyological states of basidia and basidiospores of *Dacryopinax spathularia*, and suggested that of the four postmeiotic nuclei, two migrate to each basidiospore (thus, basidiospores are haploid and uninucleate) and two degenerate in the basidium. Lange (1955), for the *Dacrymycetaceae* (as '*Dacryomycetaceae*'), suggested an ontogeny of the basidia in agreement with Bodman (1938), and depicted dikaryotic vegetative cells and 'oidia-like' mitospores, that might correspond to a particular interpretation of the arthrosporic state of, e.g., *D. stillatus*.

Oberwinkler (2014) described nuclear stages in agreement with Bodman (1938) and Lange (1955) when detailing the general life cycle of *Dacrymycetes*, exemplified by *Dacrymyces stillatus*. Duncan & Chisholm (1982) discovered a third nuclear division (pattern B of Duncan & Galbraith 1972) in the basidia of *Dacrymyces stillatus*, but basidiospores were still uninucleate (six nuclei remaining in the old basidium). McLaughlin et al. (2016) studied the ontogeny and cytology of a newly described species, *Dacryopinax primogenitus*, and observed the dikaryotic nature of the vegetative cells and conidia, as well as a similar pattern of post-meiotic nuclear division noted by Duncan & Chisholm (1982).

Most of this information closely agrees with our observations in *Dacrymycetaceae*. Although the observation of some nuclear stages was possible without stains, a clear observation of the nuclei often needs specific staining. Most nuclear staining techniques involve a long process (sometimes days) that requires the fixation of the samples prior to staining, and uses some chemicals unusual in routine taxonomic studies (Bodman 1938). As an alternative, we tried a fast (< 1 min) staining with alkaline toluidine blue, not requiring previous fixation of the samples (see Material and Methods). This gave reasonably good results, allowing us to confirm several of our observations, despite the fact that only a low number of cells were often properly stained. However, we have not been able to observe the pattern B of the third nuclear division reported by Duncan & Chisholm (1982) in any species.

Our observations of the uninucleate state in basidiospores of *Dacrymycetaceae* agree with nearly all previous information about *Dacrymycetaceae* (see above), except for Vandendries (1938), who claimed that basidiospores in *Calocera cornea* are dikaryotic. Nannfeldt (1947) likely illustrated uninucleate basidiospores in *Dacryonaema rufum* when still attached to the basidium (f. 2 on p. 328 in that study), although it was not explicitly indicated whether or not the central, rounded organelle depicted was the nucleus. Although further research is needed to confirm the number of nuclei in *Unilacrymaceae* basidiospores, young (= non-septate) basidiospores in both species seem to be uninucleate (e.g., Fig. 4t). The ancestral state reconstruction infers the uninucleate state of the basidiospores as the most probable one for the ancestors of the *Dacrymycetes* and also of *Dacrymycetaceae*, *Unilacrymaceae*, and the genus *Dacryonaema* (Table 5–6, Fig. 9).

Unlike the other families, binucleate young basidiospores seem to be the rule in the *Cerinomycetaceae*. Maekawa (1987) already indicated that recently discharged basidiospores of *Ce. aculeatus* were mostly binucleate. In addition to this species, we have consistently observed binucleate young basidiospores in five out of six species of *Cerinomycetaceae* that were studied based on living specimens (three of them included in the present phylogenetic analyses). Binucleate basidiospores in the *Basidiomycota* are not uncommon in some taxonomic groups, as noted by Duncan & Galbraith (1972). The binucleate condition is typically a product of postmeiotic nuclear division pattern D, i.e., the original haploid post-meiotic nucleus in the young basidiospore further divides inside the basidiospore and both daughter nuclei remain. *Dacrymyces tortus* s.lat. 1, in which observations of nuclei are difficult because of high amounts of lipid bodies in the basidiospore cytoplasm, showed both uninucleate and binucleate basidiospores when young and aseptate. This would still agree with pattern D, if we consider the third nuclear division more delayed than in other species, and then very recently discharged basidiospores may be uninucleate for a short period of time. Should further studies on a wider number of species confirm our observations, a comparatively long binucleate state of the basidiospores (i.e., not starting only immediately prior to septation) may represent a new synapo-

morphy and distinguishing character for the *Cerinomycetaceae*. With the present data, ancestral state reconstruction analyses indeed infer the binucleate state of the basidiospores as plesiomorphic but unique to the *Cerinomycetaceae*.

Lineage-specific presence of carotenoids

It is generally accepted that the yellow/orange colours in the *Dacrymycetes* are caused by the presence of various carotenoid pigments (Gill & Steglich 1987, Oberwinkler 2014). The precise composition has been investigated in *Calocera*, *Dacrymyces*, and *Dacryopinax* (e.g., Goodwin 1953, Hanna & Bulat 1953, Fiasson 1973, Czczuga 1980), and β -carotene has been found to be among the most abundant substances. Carotenoids are lipidophilous and thus typically found in lipid globules or in membranes (Echavarri-Erasun & Johnson 2002). We clearly observed them only in the lipid bodies of the cytoplasm, as reported for many other fungi (e.g., Cederberg & Neujahr 1970, Riley 1978, Griffin 1994). However, Vail & Lilly (1968) reported the highest amount of carotenoids in the 'cell wall fraction' of *Dacryopinax spathularia*. The colour of fresh fruitbodies in the *Dacrymycetes* varies considerably, especially at the family level, and species outside the *Dacrymycetaceae* are commonly creamish, brownish, or greyish, instead of bright yellow or orange (Fig. 5a–d). Already Nannfeldt (1947) noted that neither the basidia nor the basidiospores of *Dacryonaema rufum* contain yellow lipid drops, in contrast with their common occurrence in many species of *Dacrymycetaceae*. Oberwinkler (2014) reported that some taxa, such as *Ditiola haasii* and *Cerinomyces* species were exceptions to the rule that the basidiocarp pigmentation is caused by yellowish to orange carotenoids in the *Dacrymycetes*. The spore print is often reported as yellow to orange, and this is the case in many species within the *Dacrymycetaceae* (e.g., *Calocera viscosa*, Fig. 5e, *Dacrymyces aureosporus*, *D. chrysospermus*), but there are also reports of species with pale or whitish spore prints (e.g., *Cerinomyces albosporus*, Boidin & Gilles 1986). Other published information may not be as accurate. For instance, some taxa with clearly yellowish spore prints have been incorrectly reported as having a whitish spore print (e.g., the generic descriptions of *Dacrymyces*, *Ditiola*, *Femsjonia* and *Guepiniopsis* in Læssøe & Petersen 2014).

Our results indicate that there has been a sharp increase in the amount of carotenoids in the *Dacrymycetaceae*, whereas these pigments are not clearly observable, perhaps even absent, in *Cerinomycetaceae*, *Unilacrymaceae*, and in *Dacryonaema*. Also, the colour of the fruitbodies and that of colonies grown in pure culture seem to be related. Shirouzu et al. (2009) only reported yellow to orange colours for species in the *Dacrymycetaceae* (pale or whitish colonies also occurred), while the colonies in the clades corresponding to *Cerinomycetaceae* and *Unilacrymaceae* were constantly white.

Carotenoids have been demonstrated to play a role in the cell protection against oxidative stress and photoreactions (Will et al. 1984, Will III & Scovel 1989, Moliné et al. 2010, Avalos & Limón 2015). This function was demonstrated in pure cultures of *Dacryopinax spathularia* (Goldstrohm & Lilly 1965). However, several of the species most frequently found on exposed wood, receiving large amounts of sun-light, possess small or no amounts of carotenoids, e.g., *Dacryonaema*, *Unilacryma*, and at least one *Cerinomyces* species (*D. tortus* s.lat. 3). Therefore, a putative ecological advantage of these pigments may also be related to other functions of carotenoids (e.g., as precursors of some active compounds, Avalos & Limón 2015).

Iodine solutions are known to react with the carotenoid contents in different groups of fungi, typically exemplified by the greening of the paraphyses of several ascomycetes with Lugol's or Melzer solutions (Malençon & Bertault 1971, Schumacher

1990, other coloured reactions also occur in this group). The carotenoid contents in the *Dacrymycetaceae* display a greenish to greenish grey reaction in Lugol's solution, while this reaction has not been observed in any species of *Cerinomycetaceae*, *Dacryonaema*, or *Unilacryma*. The combination of direct observation of the fruitbodies and cell cytoplasm, plus their greenish reaction with iodine, may allow researchers to quickly and objectively assess the presence and relative amount of carotenoids.

There are, however, some species in the *Dacrymycetaceae* with colours other than the common yellow or orange, such as the whitish or very pale coloured *Calocera cavaruae*, *Ca. pallidospatulata* (Reid 1974), *Ca. fusca*, *Ditiola haasii* (Oberwinkler 1989), and *Dacrymyces cylindricus* (Shirouzu et al. 2017). For these taxa, it is reasonable to suggest a secondary loss of the pigmentation, since DNA sequence data have demonstrated that they belong to at least three different internal clades of the *Dacrymycetaceae* dominated by species with yellow-orange pigments (Shirouzu et al. 2013, 2017). This agrees with both the present phylogenies and our ancestral state reconstructions including *Ca. fusca* and *D. cylindricus* (Fig. 9). The loss of carotenoids in certain species or specimens has already been reported in other fungi (e.g., Baral 2004 in *Sarcoscypha*, Hansen et al. 2013 in some clades of *Pyronemataceae*, Olariaga et al. 2015 in *Cantharellus*). In addition, fruitbodies of species that normally produce carotenoids often produce more pigment when exposed to sunlight. In contrast, when they grow in absence of light they are often paler or even whitish, a phenomenon well known in some other *Basidiomycota* (Pilz et al. 2003, Olariaga et al. 2015). The *Dacrymycetes* are not an exception. Such uncommon cases do not invalidate the usefulness of carotenoid content for taxonomic purposes. With the current data, any species with a conspicuous and high carotenoid content would be tentatively best placed in the *Dacrymycetaceae*, since the presence of distinct carotenoids is inferred as being acquired in the ancestor of the family, and few reversions to the plesiomorphic carotenoid-deficient condition occurred (Table 5–7). We recommend that future studies on *Dacrymycetes* include careful observations of fresh, unstained samples and the subsequent reporting of approximate carotenoid content.

Other morphological characters

There is only one previous study including ancestral reconstruction analyses of multiple morphological characters in the *Dacrymycetes* (Shirouzu et al. 2013). They proposed that the macromorphology of the fruitbodies is highly homoplastic and rarely a synapomorphy for any higher clade, and the same was true for the wall thickness of marginal hyphae and, to some extent, also basidiospore septation. By contrast, other characters, such as the presence or absence of clamp connections, turned out to characterise some phylogenetic lineages.

In agreement with Shirouzu et al. (2013), our phylogeny shows that clamped septa seem to be the plesiomorphic state in the *Dacrymycetes*, while simple septa appear in some derived clades only within the *Dacrymycetaceae*. In addition, we found that clamp connection morphology and location may be informative for high-level taxa delimitation. Loop-like, long and angulose clamp connections (i.e., long medallions), almost restricted to subhymenial hyphae and the bases of basidia and hyphidia, are characteristic of the three known species of *Dacryonaema*. This specific morphology and distribution do not consistently appear in any other group, as was inadvertently indicated by Reid (1974) when he commented on the uniqueness of the *Da. macnabbii* (as *Dacrymyces macnabbii*) clamps. Loop-like, but normally shorter and more rounded clamp connections are present in some *Dacrymycetes*, as noted by, e.g., Martin (1949). We have observed them in some species, for instance in the

Dacrymyces tortus complex and in *Unilacryma*, and infrequent transitions to *Dacryonaema*-like, longer clamp connections may occur. It is interesting to note that *Unilacrymaceae*, the sister group to *Dacryonaema*, also lacks clamp connections at some septa, but this feature may occur in some species of *Cerinomycetaceae* and *Dacrymycetaceae* with typically clamped septa, due to the formation of secondary simple septa.

The morphology of cortical/marginal hyphae was used as a key character by Oberwinkler (1994, 2014) to circumscribe some genera, and it is commonly found in the literature as important to characterise, e.g., *Guepiniopsis* and *Heterotextus* (McNabb 1965c–d). Our ancestral state reconstruction analyses show that the presence of branched and anastomosing terminal cells with walls that become gelatinised in KOH is plesiomorphic and synapomorphic for *Dacryonaema* but absent in other clades, where terminal cells of cortical/marginal hyphae tend to be simple and anastomoses rare. Indeed, the ancestral condition predicted for the *Dacrymycetes*, *Cerinomycetaceae*, *Dacrymycetaceae*, and *Unilacrymaceae* is the presence of simple and not swollen terminal cells, although some uncertainty exists depending on the assumptions made at the root node (Table 5–6). These terminal cells are conspicuously dilated and morphologically diverse in some groups (e.g., *Femsjonina*, *Heterotextus*), suggesting that this character should be carefully addressed in future studies, particularly in the *Dacrymycetaceae*.

The type of basidium, i.e., forked or simple, cannot any longer be considered a good character to distinguish *Unilacryma* from other clades, as *Unilacryma bispora* constantly has bisterigmate basidia. Simple, 1-spored basidia are highly unusual in the *Basidiomycota*, and in most cases species characterised by 1-spored basidia belong to genera with a variable number of spores produced per basidium, e.g., *Heteroacanthella*, *Lactarius*, *Oliveonia*, *Russula*, *Septobasidium*, and *Tremella* s.lat. (Zamora et al. 2014, 2016, Vidal et al. 2019). This result, together with the phylogenetic position of *Dacryonaema*, allowed inferring the bisterigmate basidium as the most probable ancestral state for the *Dacrymycetes* in most analyses. However, the *rayDisc* analysis assuming equal distribution of states at the root node unexpectedly reconstructed the 1-spored basidium as the ancestral state for the *Dacrymycetes*. This is the least parsimonious option and also the equally-distributed states assumption is anyway unrealistic.

McNabb (1973) emphasised the degree of branching of the hyphidia for delimiting species within *Dacrymyces* s.lat., and this is still used to characterise some species, e.g., *Dacrymyces dendrocalami* (Oberwinkler & Tschen 1989, Shirouzu et al. 2009). In our analyses, the morphology of the hyphidia has been simplified to only two extremes (absent or simple vs conspicuous and branched) to be able to make use of imprecise descriptions from the literature. For instance, McNabb (1973) commented on the presence (sometimes occasional) of simple hyphidia in the majority of *Dacrymycetes*, in which branched hyphidia are not found, while Oberwinkler (2014) reported that the hymenium is composed of only basidia, a few species excepted. This character should be revisited in the future, since there are some clades, e.g., the one including *D. chrysocomus* and *D. ovisporus*, where hyphidia are often distinct, yet simple. In others, e.g., in *Calocera* s.str., and the *D. chrysospermus* and *D. stillatus* clades, true hyphidia are probably absent and the observed hyphae tend to develop into new basidia and new generative hyphae. While the plesiomorphic state in *Dacryonaema* and *Unilacryma* seems to be the branched hyphidia, and the ancestral state for *Dacrymycetaceae* the simple or absent hyphidia, important differences are found between the ancestors of the *Dacrymycetes* and *Cerinomycetaceae*. These differences are related to both the type of phylogenetic trees and the method

(Table 5–6), a relatively unusual result (Litsios & Salamin 2012, Cusimano & Renner 2014) that should be considered before drawing definitive conclusions. Consequently, given the current data, we consider the ancestral state of the hyphidia in the *Dacrymycetes* and *Cerinomycetaceae* to be uncertain.

There are additional characters that should be investigated in the future. Reid (1974) found that the basidium in *Calocera furcata* usually has an apical protuberance that is normally absent in *C. cornea*. This character was also noted by Van de Put (2014). The morphology of mitotic spores, whether dikaryotic arthrospores, other conidia formed from vegetative cells, or microconidia formed from the basidiospores, seems to be useful for species delimitation. Shirouzu et al. (2009) reported clear differences in mitospore morphology in fruitbodies of *D. pinacearum*, *D. stillatus*, and *D. subarcticus*, and Kirschner & Yang (2006) described morphologically characteristic stauroconidia in *Dacryoscyphus chrysochilus*. Reid (1974) described *Calocera pallidospathulata* by emphasizing the deviant cylindrical microconidia produced by mature basidiospores when compared with other species of *Calocera*, which normally produce rounded microconidia. These traits and others (e.g., characteristics in culture) are promising for delineating taxonomic groups, especially within the *Dacrymycetaceae*.

Phylogenetic relationships and boundaries among *Dacrymycetes* higher-level taxa

Differences among phylogenetic analyses

Coalescence-based phylogenetic inference often outperforms concatenation when discordance among single-locus phylogenies exists, assuming that the conflicts are a consequence of incomplete lineage sorting (Pamilo & Nei 1988, Kubatko & Degnan 2007, Degnan & Rosenberg 2009, Fernández-Mazuecos et al. 2018). Precise estimates of the effective population sizes, through inclusion of several individuals per species, benefit the species tree inference (Heled & Drummond 2010), while the superior accuracy of coalescence vs concatenation is often only relevant for relatively short branches (Ogilvie et al. 2016). With our final dataset containing no incongruence after the exclusion of *Dacryonaema RPB1* sequences, less than ten independent loci, and some species being represented by a single individual, both analytical approaches supported approximately the same groups. The few alternative relationships among some species were unsupported and affected very short branches, particularly in the *Dacrymycetaceae*. On the other hand, despite the improvements in performance in StarBEAST 2, species tree inferences often require very long runs for the MCMC to converge (Ogilvie et al. 2017). Our RWTY analyses indicate that species tree topologies did indeed reach convergence, and we only noted some convergence problems with the speciation rate parameter (cySpeciationRate, with $100 < \text{ESS} < 200$), an observation made previously (Dorsey et al. 2018).

Phylogram and chronogram topologies obtained in the present study are generally congruent with the most recent phylogenies based only on data from the nrDNA (e.g., Shirouzu et al. 2016, 2017), at least concerning the supported groups, and slightly deviate from previous studies including fewer taxa (Shirouzu et al. 2013). Our improved resolution was probably achieved by the combination of more taxa and DNA loci.

Taxonomic boundaries at ordinal and familiar ranks

Nowadays, there are two main currents for describing higher level taxa. One is the most classical, focused on the diagnosability of the resulting named groups (Christenhusz et al. 2015) considering their evolutionary relationships (thus, in agreement with phylogenies based on molecular data), as followed by, e.g., the consortium of experts in The Angiosperm Phylogeny Group

(2016). The alternative method is ‘temporal banding’, which considers estimated stem ages of the clades to define taxonomic ranks (Avice & Johns 1999, Holt & Jönsson 2014). However, adhering too tightly to the temporal banding method may have undesirable taxonomic and nomenclatural consequences. For instance, it may force the joining, under the same higher-ranked taxon name, of lower-ranked taxa that are strikingly different based on phenotypic characters, thus resulting in heterogeneous assemblages that are difficult to recognise. Conversely, it could force the splitting of morphologically homogenous groups for little gain (Lücking 2019). We think that trying to reconcile the systems may be the best solution, i.e., proposing only diagnosable groups while considering their stem ages with some flexibility. As guides for our study, the mean stem age estimates proposed by Kraichak et al. (2018) is in the range 176–194 Ma for orders and 111–135 Ma for families in the *Lecanoromycetidae* and *Ostropomycetidae* (*Ascomycota*). Zhao et al. (2017), using a wide sampling in the *Basidiomycota*, reported stem ages of most orders around 120–290 Ma. Although Zhao et al. (2017) were not focused on the family level, the oldest crown ages estimated for families were c. 150–160 Ma (e.g., in *Moniliellaceae* and *Phleogenaceae*), meaning that their stem ages must be even older and broadly overlapping with the order ages. Considering these premises, we can circumscribe at least five groups within the *Dacrymycetes* that deserve a taxonomic rank above the generic level, agreeing with the current use of ranks in the *Basidiomycota*. Four of them are well characterised by phenotypic characters and highly supported monophyletic groups in all analyses, three of which are already named as families, viz., *Cerinomycetaceae*, *Dacrymycetaceae*, and *Unilacrymaceae*. The specimen NBRC 110597 may represent another family judging from its phylogenetic position, but since we lack morphological data and its phylogenetic placement is based only on nrDNA data from a single sample, we consider it appropriate to leave it unnamed for the time being.

The clade containing the genus *Dacryonaema* includes taxa with long medallion clamps concentrated in the subhymenium and the bases of basidia and hyphidia, with septa normally lacking clamp connections elsewhere. The cortical/marginal hyphae are variable but some terminal cells are always abundantly branched and anastomosed, becoming gelatinised in KOH. Hyphidia are branched and carotenoids are inconspicuous or absent. Basidiospores are \pm cylindrical-allantoid and may remain aseptate or become transversely 1-septate (but rarely up to 3-septate) at maturity. In addition, all *TEF-1 α* sequences have an intron (the sixth in the analysed fragment) that is absent in other samples among the *Dacrymycetes*. This clade is proposed here as a new family, *Dacryonaemataceae*, since its morphological characteristics do not allow the placement of *Dacryonaema* in any of the previously circumscribed families. This decision is also in agreement with the relatively old stem age of the clade, as our results indicate that the ancestors of *Dacryonaemataceae* and *Unilacrymaceae* may have split around 169 Ma. This estimate was calculated setting an age for the *Dacrymycetes* within the ranges recently obtained by He et al. (2019) and Varga et al. (2019).

The clade representing the family *Unilacrymaceae*, which only contains the genus *Unilacryma*, is comprised of species with short clamp connections at most septa and often simple and not swollen terminal cells of marginal hyphae that are not highly gelatinised in KOH. Hyphidia are branched and there is a high lipid content in the cytoplasm of the basidia and basidiospores. Carotenoids are inconspicuous or absent. The basidiospores are ovoid to ellipsoid, presumably uninucleate when young, and become muriform at maturity. The discovery of the new species *U. bisporea* demonstrates that simple basidia, one of the main morphological characters used to recognise the family by

Shirouzu et al. (2013), is not a synapomorphy of this group. On the other hand, the presence of ovoid-ellipsoid basidiospores that become muriform, another important character noted in Shirouzu et al. (2013), is also not enough to distinguish the family, since this morphology is present in *Dacrymyces ovisporus* (Fig. 5s). This species, however, is a typical member of the *Dacrymycetaceae* and is characterised by the conspicuously high carotenoid cell content, and is additionally distinguished from *Unilacryma* by the simple to (rarely) sparingly branched hyphidia (Fig. 2l). In addition, *D. chrysocomus*, a close relative to *D. ovisporus* in our phylogenetic analyses, has predominantly cylindrical-allantoid basidiospores that frequently become muriform at maturity. Scattered shorter, ellipsoid basidiospores may be present, linking the morphology of the cylindrical-allantoid basidiospores with the ovoid ones.

The clade representing the family *Cerinomycetaceae* contains species with short clamp connections at most or all septa and often simple and not swollen terminal cells of cortical/marginal hyphae that are not clearly gelatinised in KOH. The hyphidia are often abundantly to sparingly branched, but sometimes simple or absent. The amount of lipid bodies in the cytoplasm of basidia and basidiospores is normally low (with one exception), and carotenoids are inconspicuous or absent. The basidiospores are \pm cylindrical-allantoid, binucleate during some time before the formation of the first septum, and may remain aseptate or become transversely 1–3-septate at maturity. Basidia tend to be relatively short compared to those of the other clades, especially to *Dacryonaema* and *Unilacryma*. The fruitbody macromorphology varies from resupinate, coalescing to dacrymycetoid, and the consistency is \pm dry, fleshy-cartilaginous or gelatinous.

The clade representing the family *Dacrymycetaceae* includes several genera, e.g., *Calocera*, *Dacrymyces*, *Dacryopinax*, *Femsjonia*, and *Heterotextus*. Some of these genera are non-monophyletic, which has also been demonstrated in previous studies (Shirouzu et al. 2007, 2009, 2013). This clade is very species-rich and morphologically diverse, containing species with both clampless and clamped septa, variously shaped terminal cells of cortical/marginal hyphae. Basidiospore shape and septation are highly variable, although basidiospores tend to be cylindrical-allantoid in most cases. All species studied possess basidia and basidiospore cytoplasm with high lipid contents and young basidiospores with a single nucleus before the formation of the first septum. The presence of conspicuous carotenoids and the absence of highly branched hyphidia are characteristic of most species, but exceptions occur. All *TEF-1 α* sequences obtained for this family lacked the third intron in the analysed fragment.

Taxa at the order level are not well characterised, and some doubts arise when assessing the placement of *Dacryonaemataceae*. A conservative classification that would agree with the tree topology and partly with the estimated node ages, and in addition does not require any nomenclatural changes, would be to adopt the two orders proposed by Shirouzu et al. (2013) and include *Dacryonaemataceae* in the order *Unilacrymales*. However, since the 1-spored basidium is no longer a synapomorphy of the *Unilacrymales*, and since basidiospore shape would also not be diagnostic with the putative inclusion of *Dacryonaemataceae*, the phenotypic characterization, i.e., diagnosability, of *Unilacrymales* becomes problematic. Some characters are shared by the *Dacryonaemataceae* and *Unilacrymaceae*, e.g., the clamped hyphae, branched hyphidia, absence of conspicuous carotenoids, and the similar ecology, but those traits are to some extent shared with members of the *Cerinomycetaceae*. The amount of observable lipids is not diagnostic, as no lipid drops are seen in the cytoplasm of *Da. rufum*. As noted above, basidia are often much larger in *Dacryonaemataceae* and *Unilacrymaceae* when compared with

Cerinomycetaceae, but in the *Dacrymycetaceae* some species also have relatively large basidia. The differences between *Cerinomycetaceae* and *Dacrymycetaceae* are also considerable, making it a challenge to identify diagnostic characters that can include all members of both families while at the same time excluding all taxa in *Dacryonaemataceae* and *Unilacrymaceae*. Consequently, sorting the four named families into two orders seems unpractical.

There are two alternatives instead of accepting the two currently recognised orders in the *Dacrymycetes*. One would be to create a new order for *Dacryonaemataceae*, and the second option to accept the four families (putatively five) within a single order *Dacrymycetales*. Erecting an order for *Dacryonaemataceae* might be justifiable from both a morphological point of view and by using the divergence times from Zhao et al. (2017). However, it would then be logical to also include the *Cerinomycetaceae* in a separate order, resulting in the recognition of four orders, with a single family in each one. We refrain from taking this action in the present study, as it would currently not provide any substantial benefit. The recognition of the four described families in a single order *Dacrymycetales* is the other conservative approach that does not require any nomenclatural change, and seems reasonable based on morphological and molecular data. Considering the ample overlap among stem and crown ages of orders and families in Zhao et al. (2017), and the uncertainty of our age estimates, we believe that this provisional decision does not disagree with the temporal banding method. Further studies based on genome-scale data could provide a more definite classification for the higher ranked taxa, as well as more accurate age estimates.

Taxonomy below the family level and future directions

Relationships among species groups in the *Dacrymycetaceae* are not well resolved, and generic delimitations and species circumscriptions need substantial efforts to reach taxonomic and nomenclatural stability. Subclade D1 includes *D. chrysocomus* s.lat., *D. ovisporus*, and *D. variisporus*, and is characterised by the presence of clamp connections, thick-walled terminal cells of cortical/marginal hyphae, which may form a cortex concolorous with the hymenium, and by the large basidiospores that are either transversely multiseptate and/or muriform at maturity. Subclade D2 corresponds to *Heterotextus* (*H. militinus* s.lat., generic type) and has clamp connections, a cortex of thick-walled dilated terminal cells, concolorous with the hymenium, and multiseptate but not muriform, basidiospores. Subclade D3 is formed by a group of sessile to stalked *Dacrymyces* s.lat. with clamp connections and 0–3-septate, thin-walled basidiospores (*D. ancyleus*, *D. citrinus*, *D. cylindricus*, and *D. cyrtosporus*). Subclade D4 contains a group of stalked species of *Calocera* s.lat., *Dacrymyces* s.lat. and *Dacryopinax* s.lat. with clamp connections and thin-walled, 0–3-septate basidiospores (*Calocera guepinioides* s.lat., *Ca. fusca*, *D. flabelliformis*, and *Dacryopinax sphenocarpa*). Subclade D5 is a morphologically heterogeneous group with sessile and stalked species, with and without clamp connections, with or without branched hyphidia and with thin- to slightly thick-walled, 0–3-septate basidiospores, including *Ca. arborea*, *D. dendrocalami*, and *D. cf. adpressus*. Subclade D6 corresponds to the genus *Femsjonia* (*F. monospora* and *F. peziziformis* s.lat., generic type), which has clamp connections, a contrasting pale cortex of thin- to thick-walled, cylindrical or \pm dilated terminal cells, and 1–multiseptate, thin-walled basidiospores. Subclade D7 includes specimens provisionally identified as *D. minutus*, with clamp connections, a cortex of thick-walled and \pm dilated terminal cells, concolorous with the hymenium, and 3-septate basidiospores. Finally, subclade D8, the ‘core’ *Dacrymycetaceae*, includes most species without clamp connections (with the exception of *D. pachysporus*), and the generic types of

Calocera (*Ca. viscosa*), *Dacrymyces* (*D. stillatus*), *Dacryomitra* (*Dacryomitra pusilla*), *Dacryopinax* (*Dacryopinax elegans*), *Dacryoscyphus* (*Dacryoscyphus chrysochilus*), *Ditiola* (*Di. radicata*), and *Guepinopsis* (*G. buccina*). The macromorphology in this large last group is highly variable, with fruitbodies that may be resupinate, clavarioid, cochleariform, flabelliform, pulvinate, etc., and with thin- to thick-walled basidiospores displaying various septation patterns. While some groups in subclade D8 are morphologically homogeneous, e.g., *Cerinomyces* without clamp connections, or the *D. chrysospermus* group with multi-septate basidiospores, there are also exceptions. For example, *D. novae-zelandiae* s.lat. and *D. san-augustinii*, which also have multi-septate basidiospores, are phylogenetically related to *Dacryopinax elegans*, a species with 3-septate, thick-walled basidiospores and a very different macromorphology. The lack of support for some of these groups is likely worsened by the limited amount of available DNA data. With the primers and PCR programmes provided in the present study, it should be unproblematic to obtain at least three or four unlinked DNA loci from recently collected (≤ 20 years old) samples.

The inclusion of several new samples in our *Dacrymycetes* phylogeny has led us to detect some misidentifications that have taxonomic and nomenclatural relevance. For example, *D. stillatus* and *Di. radicata*, the types of two accepted genera, have been variously interpreted in the literature. *Dacrymyces stillatus* is likely a species complex, and an improved sampling of taxa and markers is needed to clarify its identity. The specimens provisionally identified here as *D. stillatus* s.str. agree well with the protologue and with the current European concept of the species, which is a taxon very commonly found in the Nordic countries. This species has clear molecular differences with the '*D. stillatus*' sample used by Shirouzu et al. (2013), and also a somewhat different morphology according to the published description and iconography. *Ditiola radicata* is a common species in Europe, and the Swedish and Finnish specimens included in our phylogeny agree very well with the general concept of this species. However, this understanding of the species is notably different from '*Di. radicata*' sensu Shirouzu et al. (2013), obtained from culture HHB-12010, which nests in the same clade as specimens previously identified as *D. capitatus* and *D. lacrymalis*. This clade is probably another species complex, within which *D. capitatus* s.lat. displays substantial morphological variation and misidentifications are likely. The sequences retrieved from the JGI as '*Calocera cornea*' were also substantially different from those of our samples of *Ca. cornea* and agree well with preliminary data we have obtained from our own specimens of *Ca. furcata*. The present study is not aimed at resolving the taxonomy of these taxa, but the disagreements mentioned here serve to caution against uncritical taxonomic rearrangements and nomenclatural novelties based on previously generated data.

Species delimitations in *Dacryonaema* and *Unilacryma*

According to our results, the five species of *Dacryonaema* and *Unilacryma* found in the Nordic countries seem to be well differentiated, and distinguishing them by morphological or molecular data (or both) is unproblematic. The three species of *Dacryonaema* accepted here are highly supported by the STACEY species delimitation analysis, receive high support in all other phylogenetic analyses (Fig. 10), and are distinguished by several morphological characters (see Taxonomic part). Therefore, we do not consider them mere infraspecific divergent populations (Sukumaran & Knowles 2017). In addition, distribution ranges overlap at least in northern Europe, yet the three species retain their morphological distinguishing features in sympatry. *Unilacryma bispora* is an easily circumscribed species, with little taxonomic problems except for its potential confusion with

Dacrymyces ovisporus (see Taxonomic part). The STACEY analysis supported this as a distinct species, and it appears monophyletic in all phylogenetic analyses. Furthermore, basidia and basidiospore morphology (Fig. 6, 14–15) clearly separate it from its closest relative, *U. unispora*. Species delimitations in the *Unilacryma unispora* s.lat. clade are much less clear. While, as a group, it is distinguished from *U. bispora*, it is not entirely clear whether more than one species should be recognised here. As noted in the results, the *U. unispora* s.lat. clade is formed by two partially supported subclades, one joining all North European specimen and the other comprising the two Japanese specimens and additional environmental sequences from eastern Asia. It should be noted that the only molecular difference between the two clades shared by all included samples are two transitions (C/T) in the 28S dataset. Additional molecular differences are present between the 18S, ITS, and *RPB2* sequences of the Japanese specimens and those from the Nordic countries. Even if the two clades are distinguished by STACEY as two putative species, this pattern can easily be obtained when analysing few samples of isolated populations in species with a broad distribution (Sukumaran & Knowles 2017, Chambers & Hillis 2019). Also, the few differences detected might just be part of their intraspecific variation. Furthermore, the clade formed by the Japanese samples is poorly supported in the concatenated Bayesian analysis, although it is well-supported in the ML analysis. Since we have no information about other populations of *U. unispora* s.lat. in Eurasia and North America, and the amount of missing data from the East Asian clade is high due to the single region obtained from environmental studies, keeping all of them temporarily as a single species is the only justifiable option with the present data. A phylogeographic study and a careful morphological analysis involving numerous vouchered samples from throughout the northern hemisphere is needed to decide whether *U. unispora* is a single species or a complex of closely related species.

TAXONOMY

We present the following provisional key to families in the class *Dacrymycetes*:

1. Cortical/marginal hyphae with terminal cells branched and frequently anastomosing, not conspicuously dilated, at least some of which become heavily gelatinised in KOH; most hyphae thick-walled and clampless; subhymenial hyphae \pm thin-walled, with long, often angulose or slightly spurred, medallion clamp connections, present also at the bases of basidia and hyphidia, but very sparse or absent elsewhere *Dacryonaemataceae*
 (currently a single genus, *Dacryonaema*)
1. Cortical/marginal hyphae with terminal cells not or only sparingly branched, without or with disperse anastomoses, rarely becoming gelatinised in KOH (and then well-defined and conspicuously dilated); hyphae thin- or thick-walled, either all clampless or with numerous clamp connections in most parts of the fruitbodies (sometimes inconspicuous at internal hyphae); clamp connections, when present, with variable morphology, but not consistently long and angulose if of medallion type 2
2. Basidiospores \pm ovoid-ellipsoid, soon with a single complete transverse septum (very rarely 2–3), becoming muriform due to the formation of additional oblique, longitudinal and partial transverse septa; without conspicuous carotenoid contents (spore print whitish to pale yellowish cream) observable under the light microscope in any cell; hyphidia distinct and abundantly branched *Unilacrymaceae*
 (currently a single genus, *Unilacryma*)

2. Basidiospores commonly cylindrical-allantoid and aseptate or with only transverse septa, very rarely ovoid-ellipsoid and becoming muriform, but then with visible carotenoid contents when fresh (spore print often yellow or orange) that are also present in other cells (basidia, etc.), and with indistinct or simple hyphidia 3
3. Basidiospores with 0–3 transverse septa, never with longitudinal or oblique septa, thin-walled, all or several binucleate when still aseptate; basidia and basidiospores normally without a conspicuously granular/vacuolar content when observed in KOH, but sometimes with some granules that may gradually coalesce in dead state to form small (normally < 2 µm diam) lipid drops, exceptionally with a higher lipid content; cells without conspicuous carotenoid contents; clamp connections always present; basidiomata resupinate to pulvinate, never long stalked *Cerinomycetaceae* (currently a single genus, *Cerinomyces*)
3. Basidiospores with 0–many transverse septa and sometimes with additional longitudinal or oblique septa, thin- to thick-walled, uninucleate when young and still aseptate; basidia and basidiospores with a densely and conspicuously granular/vacuolar content when observed in KOH, which gradually coalesces in dead state to form conspicuous (at least some > 2 µm diam and often much bigger) lipid drops; cells frequently with distinct carotenoid contents, rarely secondary lost; clamp connections present or absent; basidiomata variously shaped *Dacrymycetaceae* (several genera)

Class *Dacrymycetes* Doweld, Prosyllabus Tracheophytorum, tentamen systematis plantarum vascularium (Tracheophyta): LXXVII. 2001

Family *Dacryonaemataceae* J.C. Zamora & S. Ekman, *fam. nov.* — MycoBank MB833097

Type. *Dacryonaema rufum* (Fr.: Fr.) Nannf., Svensk Bot. Tidskr. 41: 336. 1947.

Fruitbodies ± firmly gelatinous when fresh, sometimes cartilaginous, xerotolerant, sessile or stalked, with the fertile parts pulvinate to depressed or subglobose. *Hymenium* amphigenous, or confined to the upper part of the fruitbody when fruitbodies are discoid, then with a distinct sterile cortex. *Cortical/marginal hyphae* variable, frequently clampless, but sparse loop-like clamp connections may occur, some with terminal cells clearly branched and anastomosing, strongly gelatinised in KOH; unbranched cylindrical, thick-walled and slightly dilated terminal cells may also occur, then normally tapering at the apex. *Internal hyphae* thick-walled, clampless or with very sparse clamps. *Subhymenium* composed by more or less thin-walled hyphae, with long and angulose, medallion clamps. *Basidia* 2-spored, bearing a loop-like basal clamp connection; apex frequently U-shaped, sometimes Y-shaped, and more rarely with an apical protuberance. *Hyphidia* abundantly branched and often anastomosing, with a loop-like basal clamp connection. Recently discharged and still aseptate basidiospores uninucleate. *Mature basidiospores* 0–1-septate (exceptionally 3-septate), thin-walled, hyaline, cylindrical-allantoid. *Spore print* whitish. *Microconidia* ellipsoid. *Cell cytoplasm* without conspicuous carotenoids under the light microscope. Brownish intracellular and diffuse parietal pigments often visible at least in some cortical/marginal hyphae. Basidia and basidiospores with or without conspicuous lipid drops when old.

Included genera — *Dacryonaema* Nannf., Svensk Bot. Tidskr. 41: 336. 1947 (type).

Key to species

1. Fruitbodies for a long time appearing as a conical, dark brown to dark reddish brown (rarely amber-coloured), sterile stalk, which finally develops an apical, often rounded or irregularly shaped, but not centrally depressed, pruinose fertile head, ≤ 0.7 mm diam when fresh and mature; dead basidia and basidiospores with a rather even cytoplasm, with invisible or hardly discernible lipid bodies; basidiospores (2.9–)3.2–4.3(–4.9) µm wide *Da. rufum*
1. Fruitbodies very variable in colour, from yellowish or cream coloured to brownish or dark brown, pulvinate to pustulate at first, later slightly convex or becoming flattened or centrally depressed (pezizoid), sessile or sometimes shortly stalked, often ≥ 0.7 mm diam when fresh and mature; dead basidia and basidiospores with a slightly to conspicuously granular cytoplasm, often with some discernible lipid bodies; basidiospores (3.5–)4.3–6.6(–7.1) µm wide 2
2. Fruitbodies yellowish, cream coloured or brownish; cell cytoplasm with distinct or indistinct lipid content; basidiospores on average < 14 × 5.3 µm *Da. macnabbii*
2. Fruitbodies brown to dark brown or blackish; cell cytoplasm normally with a rather distinct lipid content; basidiospores on average > 14.5 × 5.3 µm *Da. macrosporum*

Dacryonaema macnabbii (D.A. Reid) J.C. Zamora & S. Ekman, *comb. nov.* — MycoBank MB833098; Fig. 11

≡ *Dacrymyces macnabbii* D.A. Reid, Trans. Brit. Mycol. Soc. 62: 456. 1974 [basion.]

Typus. UNITED KINGDOM, Wester Ross, Kinlochewe, Coille na Glas-Leitire, on *Pinus sylvestris*, 20 Aug. 1963, R.W.G. Dennis (K(M) 81678! holotype).

Basidiocarps ± firmly-gelatinous, from pale cream coloured to yellowish, ochraceous, or brownish, sometimes with an orange tinge, darker when dried, xerotolerant, at first erumpent or pulvinate, soon discoid, flattened to centrally depressed (pezizoid), sessile or shortly stalked, attached to the substrate by a central point and sometimes rooting, (0.5–)0.7–1.5(–1.8) mm diam when mature. *Cortical/marginal hyphae* conspicuous, variable, at least some with terminal cells abundantly branched and partially anastomosing, with walls that become gelatinised in KOH; some slightly to clearly thick-walled and somewhat dilated cells may also appear. *Internal hyphae* (2–)2.5–7 µm wide, slightly to clearly thick-walled, often roughened, embedded in a gelatinous matrix, clampless or, rarely, with scattered and obscure long clamps. *Hymenium* usually confined to the upper surface of the fruitbodies, rarely becoming amphigenous with age, forming numerous basidia intermixed with hyphidia. *Subhymenium* with thin- to slightly thick-walled hyphae, with long and ± angulose medallions, including at the bases of basidia and hyphidia. *Hyphidia* well-developed, profusely branched, normally abundant, 2–4(–5) µm thick in the upper half, terminal branches 1–2(–2.5) µm thick, often with simple septa throughout their length. *Young basidia* cylindrical to very narrowly clavate, sometimes with a tortuous base. *Mature basidia* 2-spored (very rarely 3-spored basidia have been seen), (42–)45–100(–125) × (4–)4.5–7(–10) µm, sterigmata 11–20 µm long, often forming an obtuse angle or arising from slightly lateral position from the apex, basidia sometimes with a small apical papilla between the sterigmata. *Basidiospores* (8.7–)10.4–13.9(–15.4) × (3.5–)4.3–5.5(–6.2), Q = (1.6–)2–3.2(–3.7), ellipsoid to cylindrical-allantoid. *Basidia* and *basidiospore cytoplasm* hyaline, often slightly granular in KOH due to the presence of some lipid bodies, which may form small lipid drops after coalescence. *Microconidia* from germinating basidiospores rarely seen, hyaline, narrowly ellipsoid to cylindrical, about 3.5–4.5 × 1–1.5 µm.

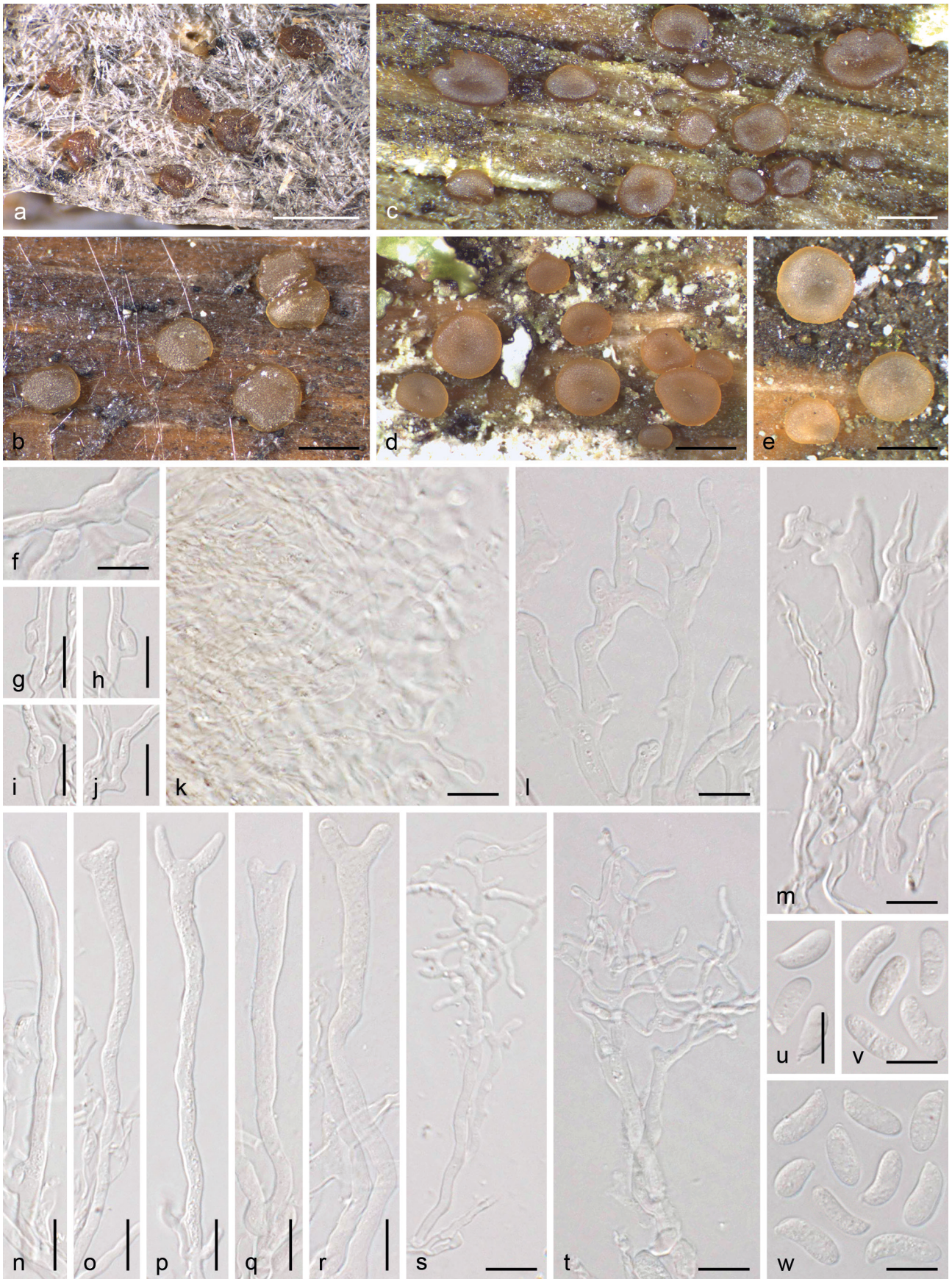


Fig. 11 *Dacryonaema macnabbii*. a–e. Fruitbodies in different degrees of development, all in fresh condition except a; f–w. Micromorphology; f. thick-walled, clampless internal hyphae; g–j. long medallion clamp connections; k. gelatinised and branched terminal cells of cortical hyphae; l. anastomoses; m. thick-walled and slightly dilated marginal hyphae; n. young basidium; o–r. basidia in different degrees of development; s–t. branched hyphidia; u–w. basidiospores. a, u. From the holotype; b, g–j, m–p, s, v. from UPS F-940950; c, k. from UPS F-940966; d–f, l, q–r, t, w. from UPS F-940954. — Scale bars a–e = 1 mm, f–w = 10 μ m.

Ecology & Distribution — Apparently common on exposed coniferous wood, particularly *Pinus sylvestris* branches still hanging from the tree, but also found on fallen wood lying on the ground. Its distribution is uncertain because it has been confused with *Dacrymyces tortus* and other species, but published records and studied specimens demonstrate that it is present in the temperate and boreal biomes of north-western Europe. We have studied specimens of *Da. macnabbii* from Finland, Norway, Sweden, and the United Kingdom. It has also been reported from Denmark by Mathiesen (1991), whose descriptions and iconography agree well with our observations. The only two Swedish specimens previously identified as *Da. macnabbii* (S 250593 and UPS F-561899) were misidentified and belong to the *D. tortus* group.

Nomenclature & Taxonomy — The yellowish to brownish colour of the fruitbodies, as well as the abundantly branched hyphidia, the long loop-like clamp connections present at the bases of basidia, and the basidiospore dimensions, as already reported in the protologue (Reid 1974), easily distinguish this species. The holotype specimen is particularly well distinguished from the morphologically close *Da. macrosporum* thanks to its clearly ochraceous colour and the short basidiospores (10.5–13 µm as indicated by D.A. Reid). In addition, all sequenced samples accepted here as *Da. macnabbii* form a well-supported clade including Reid's type collection, clearly independent from the clade formed by all *Da. macrosporum* samples. Some species in the *D. tortus* group may be macro-morphologically very similar to both *Da. macnabbii* and *Da. macrosporum*, and a microscopical study is often needed to separate them. The cortical/marginal hyphae of *D. tortus* s.lat. are not finely and abundantly branched, their walls barely gelatinise in KOH (they may show a somewhat roughened surface), the hyphidia are less abundantly branched, basidia are smaller (particularly shorter), and clamp connections are much more abundant in all hyphae and not of the form of long or angulose medallions (short medallions may be present particularly on internal hyphae).

Additional specimens examined. FINLAND, Lapponia inarenensis, Inari, near Inari Siida, N68.910836° E27.016711°, on a woodplank, 15 Aug. 2018, J.C. Zamora & P. Posio, UPS F-940997. — NORWAY, Akerhus, Urdal, NE-side of Buråstjern, PN 121,014, on *Pinus sylvestris*, 24 May 1981, Anna-Elise Torkelsen 33/81 (O 160027); Akerhus, Nes, Kleiva, *Picea abies*, 13 Sept. 1965, Anna-Elise Torkelsen 113/5 (O 160060). — SWEDEN, Blekinge, Agerum, Fjärdsjömåla, på granstock, 03 Dec. 1946, S. Wilkand 328 (UPS F-729787); Öland, Böda, Lindreservatet, N57.319448° E17.030849°, on *Pinus sylvestris* wood, 03 Oct. 2017, J.C. Zamora (UPS F-940992); Småland (Inre), Femsjö, Prästgårdsskogen, på murken barträdssam, 17 Oct. 1939, Seth Lundell 1739 (UPS F-730059); Östergötland, Gryt, c. 1000 m Ö om Forhem, ad ramos dejectos decorticatos pini, 23 Apr. 1946, J.Ax. Nannfeldt 8308 (UPS F-729963); Södermanland, Nacka, Nackareservatet, N59.28955° E18.17560°, on *Pinus sylvestris* wood, 10 June 2018, J.C. Zamora, P. Posio & J. Nieminen (UPS F-940966); Värmland, Lekvattnet, N60.163134° E12.71224°, on *Pinus sylvestris* branch, 06 Oct. 2018, J.C. Zamora (UPS F-940986); Värmland, Lekvattnet, N60.172306° E12.699782°, on *Pinus sylvestris* branch, 07 Oct. 2018, J.C. Zamora (UPS F-940990, UPS F-940991); Västmanland, Sala, Skuggan, Fiskarbo, grönmålat virke (stöd åt hänggrön), 07 July 1946, R. Morander 658 (UPS F-729778, collection mixed with *D. tortus* and other species); Uppland, Älvkarleby, Långsandsörarna island, N60.65783° E17.55504°, on *Pinus sylvestris* branches, 17 Apr. 2018, J.C. Zamora, S. Ekman & M. Zuluaga (UPS F-940954, duplicate in H); Uppland, Uppsala-Näs, Lörsta skog, nära Skärfälten, på tallgren på marken, 24 May 1928, Seth Lundell 629 (UPS F-730091); Uppland, Uppsala, Norra Lunsen, N59.80680° E17.96804°, on coniferous wood, probably *Pinus sylvestris*, 20 Aug. 2017, J.C. Zamora (UPS F-940951, UPS F-940952); Uppland, Uppsala, Norby skog, and morschen *Pinus*-Åsten am Boden, 10 Aug. 1928, S. Lundell 631 (UPS F-729789); Uppland, Uppsala, Norra Lunsen, N59.79390° E17.68667°, on *Pinus sylvestris* wood, 19 Nov. 2017, J.C. Zamora (UPS F-940950); Uppland, Uppsala, Norra Lunsen, N59.790953° E16.687811°, on *Pinus sylvestris* branch, 30 Sept. 2018, J.C. Zamora, S. Ekman, Å. Kruys & D. Pfister (UPS F-940953); Uppland, Uppsala, Nästen, ovanför Läbyvads anhalt, på murket tallgårdsgel, 19 June 1931, S. Lundell 792 (UPS F-729790); Uppland, Uppsala, Stadsskogen

naturreservat, Vallstigen, N59.839763° E17.626805°, on highly decomposed, probably coniferous, wood, 09 Apr. 2017, J.C. Zamora (UPS F-940949); Uppland, Viksta, strax NV om 'Stentorget', ad ramos dejectos decorticatos pini, 25 May 1947, J.Ax. Nannfeldt 9141 (UPS F-729994, collection mixed with *D. tortus*); Hälsingland, Enånger, Borka, N61.54875° E17.04727°, on old *Pinus sylvestris* branch, 27 July 2018, J.C. Zamora (UPS F-940977); Medelpad, Borgsjö, Ånge, N62.54089° E15.70138°, on old exposed *Pinus sylvestris* branch, 27 July 2018, J.C. Zamora (UPS F-940980); Jämtland, Mörsil, Klukshäckren, N63.172614° E13.600665°, on *Pinus sylvestris* branches, 23 July 2018, J.C. Zamora (UPS F-940993, UPS F-940994, duplicate in H); Jämtland, Mörsil, c. 2 km W of Säillsjö, N63.236875° E13.634149°, on old and weathered *Pinus sylvestris* branches, 26 July 2018, J.C. Zamora (UPS F-940995, UPS F-940996).

***Dacryonaema macrosporum* J.C. Zamora & S. Ekman, sp. nov.** — MycoBank MB833099; Fig. 12

Etymology. The specific epithet '*macrosporum*' refers to the large basidiospores compared to those of the two other *Dacryonaema* species described so far.

Typus. FINLAND, Ostrobothnia ultima, Rovaniemi, N66.500201° E25.763870°, on *Pinus sylvestris* wood, 14 Aug. 2018, J.C. Zamora & P. Posio (UPS F-941000! holotype, isotypes in UPS F-941001! and H!).

Diagnosis. Morphologically similar to *Dacryonaema macnabbii*, but fruitbodies frequently darker (brown to dark reddish brown or blackish), cell cytoplasm with more distinct lipid bodies (higher lipid content), and basidiospores distinctly larger, (12.9–)14.7–18.6(–23) × (4.4–)5.2–6.6(–7.1) µm (on average > 14.5 × 5.3 µm).

Basidiocarps ± firmly gelatinous, yellowish brown or ochraceous brown when young but soon becoming brown to dark brown or blackish, darker when dried, xerotolerant, at first erumpent or pulvinate, later discoid, flattened to centrally depressed (pezizoid), sessile or shortly stalked, attached to the substrate by a central point and sometimes rooting, (0.4–)0.7–1.5(–1.7) mm diam. **Cortical/marginal hyphae** variable, some abundantly branched and anastomosing terminal cells, with walls that gelatinise in KOH, others slightly to clearly thick-walled and that may be partially swollen. **Internal hyphae** (2–)2.5–6 µm wide, slightly to clearly thick-walled, often roughened, embedded in a gelatinous matrix, and clampless, sometimes with scattered and obscure long clamps. **Hymenium** restricted to the upper surface of the fruitbodies, progressively forming numerous basidia intermixed with hyphidia. **Subhymenium** formed by thin- to slightly thick-walled hyphae, with common long and ± angulose, medallion clamp connections, including at the bases of basidia and hyphidia. **Hyphidia** abundantly branched, 2–4.5 µm thick in the upper half, terminal branches 1–2.5(–3) µm thick, often with simple septa throughout their length. **Young basidia** cylindrical to narrowly clavate, sometimes with a tortuous base. **Mature basidia** 2-spored (3-spored basidia may be rarely seen), 65–112 × 5.5–7.5(–9) µm. Sterigmata 16–30 µm long, often forming an obtuse angle or arising from slightly lateral position from the apex, rarely with an apical papilla. **Basidiospores** (12.9–)14.7–18.6(–23) × (4.4–)5.2–6.6(–7.1) µm, Q = (2.2–)2.4–3.3(–4), cylindrical-allantoid. **Basidia** and **basidiospore cytoplasm** hyaline or, sometimes, with a subtle cream hue, normally with a ± granular content in KOH due to the presence of lipid bodies, which may coalesce to form some distinct lipid drops. **Microconidia** from germinating basidiospores rarely seen, hyaline, narrowly ellipsoid to cylindrical, about 3–4.5 × 1–1.5 µm.

Ecology & Distribution — Insufficiently known; it seems to be much rarer than *Da. macnabbii* and *Da. rufum*. All studied specimens were growing on rather weathered and decomposed *Pinus sylvestris* wood, on branches lying on the ground, where it may grow intermixed with species in the *D. tortus* group (e.g., in Fig. 11a, the slightly paler fruitbody partially shown at the bottom of the picture is *D. tortus* s.lat. 3).

Nomenclature & Taxonomy — This species is morphologically distinguished from *Da. macnabbii* by its darker colour and,

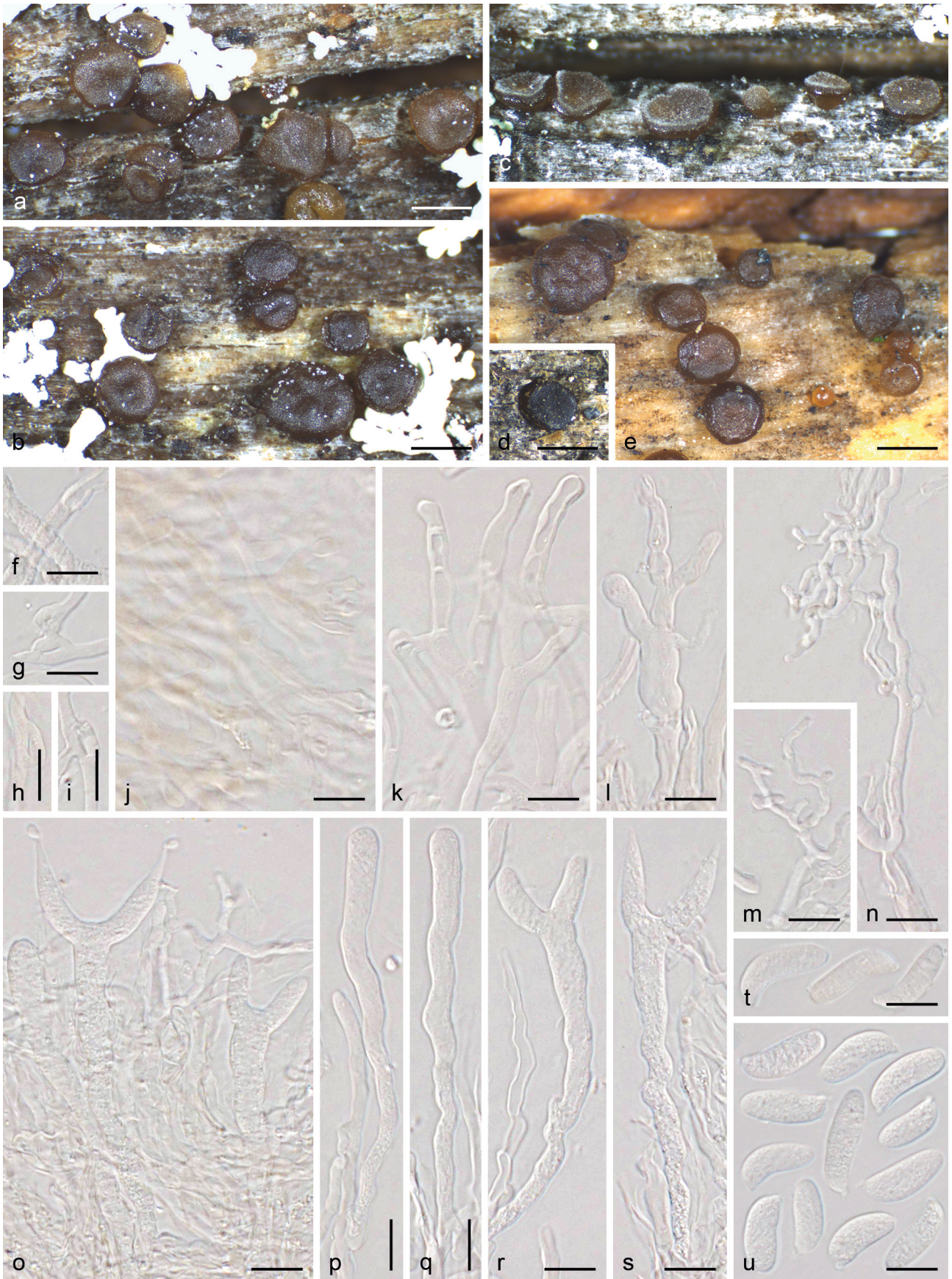


Fig. 12 *Dacrynaema macrosporum*. a–e. Fruitbodies in different degrees of development, all in fresh condition except d; d–u. Micromorphology; f. thick-walled, clampless internal hyphae; g–i. long medallion clamp connections; j. gelatinised and branched terminal cells of cortical hyphae; k. anastomoses; l. thickened and slightly dilated marginal hyphae; m–n. branched hyphidia; o. hymenium showing young basidia, mature basidia and hyphidia; p–q. young basidia; r–s. mature basidia in different degrees of development; t–u. basidiospores. a–c, f, h–j, l–s, u. from the holotype; g, k. from UPS F-941002.; d, t. from O 160045; e. from the isotype. — Scale bars a–e = 1 mm, f–u = 10 μ m.

in particular, by the larger basidiospores. However, since the description above is based on only c. 250 fruitbodies (c. 150 from the holotype and isotype) from six individuals, its intra-specific variation is insufficiently known, so the traits selected as diagnostic characters should be taken with some caution. Light-coloured specimens of *Da. macrosporum* and dark coloured specimens of *Da. macnabbii* may be very difficult to separate, unless they are fertile and a certain number of well-developed basidiospores could be measured. *Dacryonaema* fruitbodies are comparatively long-lived and environmental conditions may affect basidiospore size, particularly if the maturation is aborted when the fruitbodies suddenly dry up. In addition, basidia in *Da. macrosporum* tend to be slightly wider and the amount of visible lipids in the basidia and basidiospore cytoplasm is slightly higher. Unfortunately, those characters show a considerable overlap and, rarely, specimens of *Da. macnabbii* with unusually wide basidia and a considerable amount of lipid bodies occur. In any case, the two species are clearly separated by DNA sequence data. For the distinction between *Da. macrosporum* and species in the *D. tortus* group, see taxonomic comments under *Da. macnabbii*. The two isotypes are part of the same individual, while the holotype is a different individual, collected on a different piece of wood. However, the three specimens are to be considered as duplicates because they are part of the same gathering (ICN, Art. 8.2, Turland et al. 2018).

Additional specimens examined. FINLAND, Ostrobothnia ultima, Rovaniemi, N66.499030° E25.763717°, on *Pinus sylvestris* branches, 19 Aug. 2018, J.C. Zamora (UPS F-941002); Regio Kuusamoensis, Salla, Oulanka, Urriaavantie, N66.439727° E29.165568°, on *Pinus sylvestris* branch, 17 Aug. 2018, J.C. Zamora (UPS F-940998, UPS F-940999). – NORWAY, Hedmark, Kongsvinger, ved Sikåa mellom Møkeren og Varaldsjøen, on *Pinus sylvestris*, 27 May 1978, Anna-Elise Torkelsen 83/78 (O 160045).

Dacryonaema rufum (Fr.: Fr.) Nannf., Svensk Bot. Tidskr. 41: 336. 1947 — Fig. 13

≡ *Sphaeronaema rufum* Fr., Kongl. Vetensk. Acad. Handl. 1818 [ser. 3, vol. 6]: 357. 1818 [basion.]: Fr., Syst. Mycol. 2: 536. 1823.

≡ *Zythia rufa* (Fr.) Schwein., Trans. Amer. Philos. Soc., n.s. 4: 247. 1834 [1832].

≡ *Sphaeronaemella rufa* (Fr.) Sacc., Syll. Fung. 3: 618. 1884.

Typus. SWEDEN, [sine loc.], Scleromyceti Sueciae n. 27 (UPS F-117078) lectotype, designated here, MBT389286).

Basidiocarps firm-gelatinous to cartilaginous, xerotolerant, often consisting on a well-developed (but sometimes very short), reddish brown to dark reddish brown, narrowly conical stalk, 0.3–2(–2.5) mm long, 0.3–0.7(–0.8) mm wide at the base and tapering to around 0.2–0.4 mm wide near the apex. Tardily, an apical, concolorous or slightly paler, rounded, fertile head is developed, becoming pruinose and thus looking distinctly paler when fully mature, 0.3–0.8 mm diam. Fruitbodies sometimes orangish, amber-coloured or with a pinkish hue when growing in the dark or when developing and maturing quickly, distinctly shorter in the last case. *External hyphae of the stalk* typically thick-walled, strongly glued together and difficult to dissect; *cortical/marginal hyphae* variable, at least some with thin-walled, anastomosing and branched terminal cells, gelatinising in KOH, others thin- to thick-walled and mostly unbranched, sometimes a few slightly dilated thick-walled terminal cells can be seen; cytoplasmic contents hyaline or with some brownish pigments; walls hyaline to slightly yellowish brown or brownish. *Internal hyphae* 3–10 µm wide, thick-walled, claspless or with very scattered and obscure long clamp connections, slightly embedded in a cartilaginous matrix and tightly adhering. *Hymenium* amphigenous, producing numerous basidia intermixed with somewhat scattered hyphidia. *Subhymenium* formed by thin- to slightly thick-walled (rarely clearly thick-walled) hyphae, bearing long and

± angulose medallion clamp connections, including at the bases of basidia and hyphidia. *Hyphidia* distinct but sometimes sparse, branched, 2–4.5 µm thick in the upper half, terminal branches 1–2.5 µm thick, often with simple septa throughout their length. *Young basidia* cylindrical to narrowly clavate. *Mature basidia* 2-spored, 35–80(–85) × (2.5–)3.5–5.5(–6.5) µm, sterigmata 9–23 µm long, often forming an obtuse angle or arising from slightly lateral position from the apex, rarely with an apical papilla. *Basidiospores* (8.7–)10.1–13.4(–14.8) × (2.9–)3.2–4.3(–4.9), Q = (2.2–)2.5–3.9(–4.9), cylindrical-allantoid. *Basidia* and *basidiospore cytoplasm* hyaline, normally without visible lipid contents, rarely with scattered and minute lipid drops. *Microconidia* from germinating basidiospores hyaline, narrowly ellipsoid to cylindrical, about 3.5–4.5(–5) × 1–1.5 µm.

Ecology & Distribution — Rather common and widespread, known from the temperate and boreal biomes of North America and Europe but not recorded from Eastern Asia (Shirouzu et al. 2009, 2016). It is apparently restricted to coniferous wood, mainly *Pinus sylvestris* in Europe, but also occurs on other genera and species (e.g., *Pseudotsuga*) in North America. It very frequently occurs on anthropic wood, and is rather widespread on fences, roofs, benches, etc. The ecology of this species was thoroughly detailed by Nannfeldt (1947).

Nomenclature & Taxonomy — No specimens are cited, nor are illustrations or locality data present in the protologue of *Sphaeronaema rufum* (Fries 1818). However, in the later *Systema Mycologicum* 2, Fries (1823) cited the exiccata ‘Scleromyceti Sueciae n. 27’. According to the current Art. F.3.9 of the ICN, and since the name is sanctioned, those specimens are available for lectotype designation by being elements from the context of the sanctioning work. Two duplicates of *Scleromyceti Sueciae* n. 27 with the original label are kept in UPS. One is mounted on an envelope joined with other samples of the *Scleromyceti Sueciae* (first 100 numbers, UPS F-117472), and the other is mounted on a free envelope (UPS F-117078). The two specimens are conspecific, well-developed and typical, despite all fruitbodies being young and sterile. We chose UPS F-117078 as the lectotype in order to facilitate future revisions. Since the selected lectotype is taxonomically unambiguous, the designation of an epitype is unjustifiable (ICN, Art. 9.9), regardless of, e.g., the lack of molecular data from this sample. We have studied two further specimens collected by E. Fries. One (S F250639) was collected in Sweden without other locotypic indication, and is labelled as ‘Original!’ by a later hand than that of Fries. Both ‘*Sphaeronaemella rufa* (Fr.) Sacc.’ and ‘*Sphaeria rufa* Fr.’ were written with the same ink and by the same hand, so the label was likely written after Saccardo (1884) made the combination in *Sphaeronaemella*. The other specimen (UPS F-117079) was collected in Femsjö, but the label has no date and Femsjö is not cited as a collecting locality neither in the protologue nor in the sanctioning work. As a result, none of these specimens can be considered as original material.

The initially conical and typically dark reddish brown fruitbodies of *Da. rufum* have a unique morphology among all *Dacrymycetes*, being easily characterised in the field whether in its sterile or fertile state. Indeed, this species is not likely to be confused with any other *Dacrymycetes*, but some ascomycetes producing dark peritheca or pycnidia with long necks might appear similar to the naked eye. In fact, Fries (1818, 1823) placed various ascomycetes in the same genus and described the fruitbodies as perithecia. When fertile, *Da. rufum* is more difficult to confuse with any other species, although perhaps some confusion with calicioid fungi could arise. A careful examination under the microscope or even with a hand-lens would quickly dispel any doubt. Clamp connections in this species have been largely overlooked, even in the detailed study of Nannfeldt (1947), perhaps because they are normally restricted to the sub-

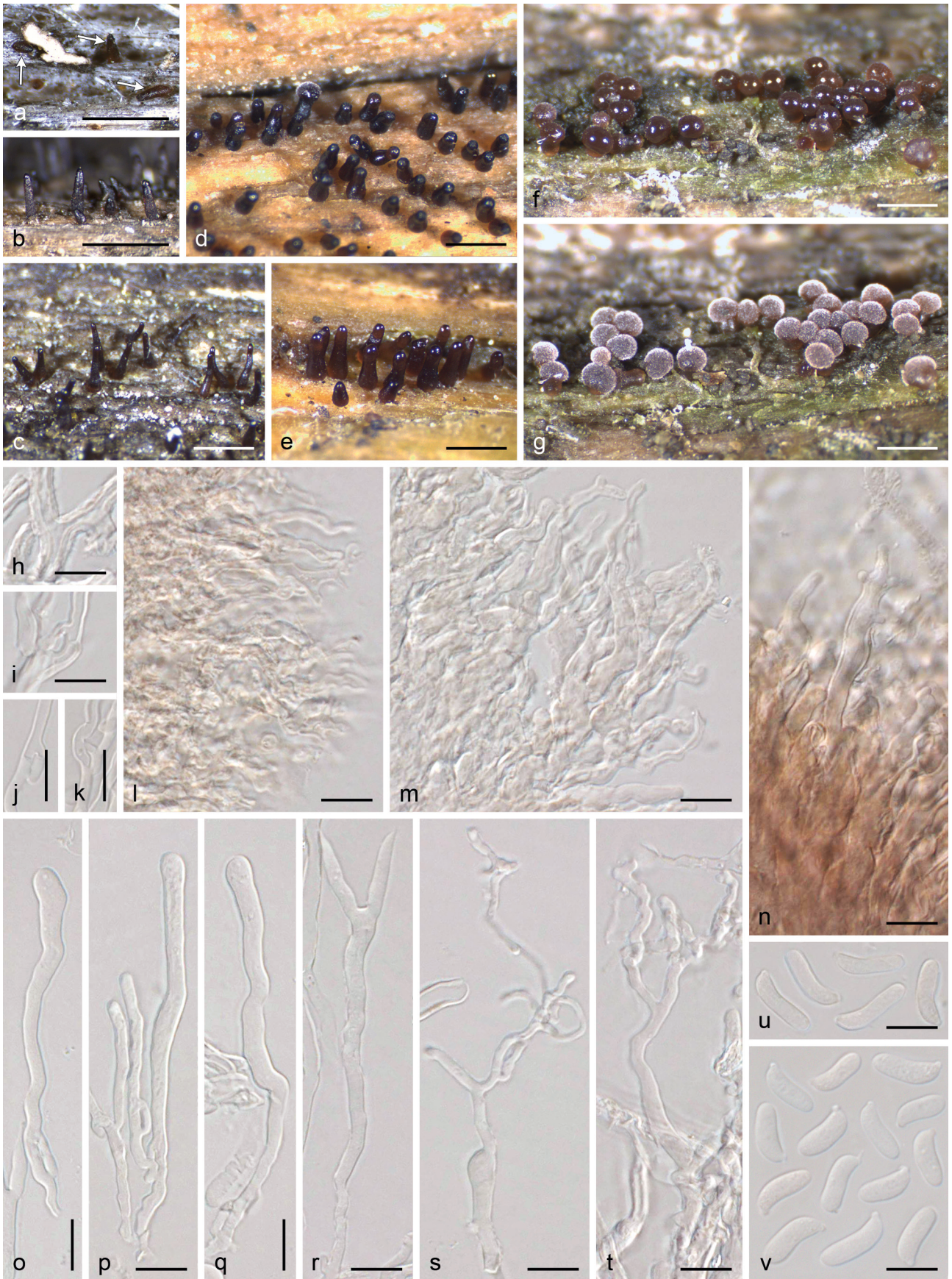


Fig. 13 *Dacryonaema rufum*. a–g. Fruitbodies in different degrees of development, all in fresh condition except a–c; h–v. Micromorphology; h. thick-walled, clampless internal hyphae; i–k. long medallion clamp connections; l–m. slightly gelatinised and branched terminal cells of marginal hyphae; n. thick-walled and slightly dilated marginal hyphae; o–q. young basidia; r. mature basidium; s–t. branched hyphidia; u–v. basidiospores. a. From the lectotype; b. from TRTC 48002 (in UPS); c. from UPS F-941012; d–s, v. from UPS F-941005; t–u. from UPS F-941006. — Scale bars: a–g = 1 mm, h–v = 10 µm.

hymenium and the bases of basidia and hyphidia, and because it is difficult to dissect *Da. rufum* specimens due to their hard texture. However, they were noted and accurately illustrated by Brough & Bandoni (1975), but this was later considered as an exception in some North American specimens (e.g., Læssøe & Petersen 2014). All fertile specimens of *Da. rufum* studied by us have conspicuous clamp connections at the bases of basidia, regardless of their geographical origin. The branched hyphidia were also not noted by Nannfeldt (1947), who stated: “no sterile elements have been seen in the hymenium”. However, although they may be scarce and difficult to find in some samples, and several slides may have to be prepared to locate them, they are present in all fertile specimens studied by us. Brough & Bandoni (1975) also found and illustrated branched hyphidia in the hymenium of this species.

Additional specimens examined (due to the high number of specimens studied, for European material we only cite those specimens used for morphological and molecular analyses, and typification, but c. 50 additional specimens, mainly from the Nordic Countries and preserved in O, S, and UPS, were revised as well). CANADA, Nova Scotia, Kings Co., Kentville, coniferous fence post, 17 Nov. 1954, K.A. Harrison (UPS s.n.); Ontario, Thunder Bay Distr., Black Sturgeon Lake, on old wood, 17 June 1977, A. Carter (TRTC 48002 in UPS). – FINLAND, Tavastia australis, Tammela, Mustiala, Salois (Saloinen), around P.A. Karsten’s house, on a wood plank, 11 May 2018, J.C. Zamora & P. Posio (UPS F-941012). – NORWAY, Hordaland, Voss, Mølster-tunet, på gjerde, LN 583-585,246-248, 23 Sept. 1990, Anna-Elise Torkelsen 389/90 (O 295442); Troms, Storfjord, Skibotndalen, Lullesletta, DB 77,88, on *Pinus sylvestris*, 19 Aug. 1992, Anna-Elise Torkelsen 78/92 (O 295528). – SWEDEN, Skåne, Örkelljunga, Storeös mosse, on a wooden fence, 08 June 1977, Rolf Santesson 27333 (UPS F-730180); Uppland, Solna, Bergshamra, N59.38671° E18.02956°, on an old, fallen *Pinus sylvestris* tree, 01 Apr. 2018, J.C. Zamora & P. Posio (UPS F-941004, UPS F-941005); *ibid.*, 08 July 2018, J.C. Zamora (UPS F-941006); Uppland, Uppsala, Norra Hagunda, Vänge, close to the parking at south of Fiby urskog naturreservat, N59.881589° E17.354279°, on a wood bench, 07 Apr. 2017, J.C. Zamora, M. Westberg, M. Svensson & S. Ekman (UPS F-941003); Jämtland, Åre, Mörsil, Klukshäckren, N63.172614° E13.600665°, on *Pinus sylvestris* branches, 23 July 2017, J.C. Zamora (UPS F-941011). – UNITED STATES OF AMERICA, Michigan, Mackinac Co., Big Knob Campground on Lake Michigan, N46.0333° W85.5833°, in a *Thuja occidentalis* swamp, 20 Aug. 1977, R. Santesson 27444 (S F250641); Oregon, Benton County, Mary’s Peak, 914 m, on decorticated branch of douglas fir, 26 Nov. 1983, B.R. Kropp (UBC F12372, UBC F12373).

Family **Unilacrymaceae** Shirouzu, Tokum. & Oberw., Mycologia 105: 1120. 2013

Type. *Unilacryma unispora* (L.S. Olive) Shirouzu, Tokum. & Oberw., Mycologia 105: 1120. 2013.

Basidiocarps moderately to clearly gelatinous, but not of soft consistency, xerotolerant, pulvinate to depressed pulvinate when mature, with an almost amphigenous hymenium. *Cortical/marginal hyphae* normally unbranched, ± thick-walled, with terminal cells cylindrical to slightly tapering, not conspicuously swollen, with frequent clamp connections but also with some simple septa; anastomoses sparse. *Context hyphae* slightly to clearly thick-walled, with ± rounded and short clamp connections visible at several septa, but some loop-like clamp connections and simple septa also often present. *Subhymenium* composed by thin to slightly thick-walled, clamped hyphae. *Basidia* 1- or 2-spored, bearing a basal clamp connection; when 2-spored apex often U-shaped, but basidia with an apical protuberance or with sterigmata forming an acute angle also occur. *Hyphidia* sparsely to abundantly branched, rarely anastomosing among them, with a basal clamp connection. Recently discharged and still aseptate basidiospores presumably uninucleate. *Basidiospores* subglobose to narrowly ovoid or ellipsoid, soon with one transverse septum and later becoming muriform due to the formation of additional longitudinal, oblique and sometimes transverse septa, thin-walled, hyaline or with a very subtle yellowish tinge. *Spore print* whitish. *Microconidia*

ellipsoid. *Cell cytoplasm* without conspicuous carotenoids under the light microscope, at most, some basidia and basidiospores with a very subtle creamish tinge. Parietal diffuse yellowish brown pigments sometimes seen at some cortical/marginal hyphae. Basidia and basidiospores with conspicuous lipid bodies, becoming ≥ 2 µm lipid drops at least in some old cells.

Included genera — *Unilacryma* Shirouzu, Tokum. & Oberw., Mycologia 105: 1120. 2013 (type).

Key to species

- Basidia 1-spored, 5–7.5 µm width; basidiospores (10.2–) 11.2–13.5(–15.7) µm width, Q = (1.1–)1.2–1.4(–1.6), most of them ovoid to broadly ellipsoid *U. unispora*
- Basidia 2-spored, 6.5–8.5(–9) µm width; basidiospores (7.2–)8.2–10.4(–12.4) µm width, Q = (1.2–)1.4–1.9(–2.2), most of them narrowly ovoid to ellipsoid *U. bispora*

Unilacryma bispora J.C. Zamora & Ekman, *sp. nov.* — Myco-Bank MB833100; Fig. 14

Etymology. The specific epithet ‘*bispora*’ refers to the 2-spored basidia, in contrast to the 1-spored basidia of the only other described species in the genus, *U. unispora*.

Typus. SWEDEN, Uppland, Uppsala, Norra Lunsen, N59.79390° E17.68667°, on an old *Pinus sylvestris* branch, 19 Nov. 2017, J.C. Zamora (UPS F-941257! holotype, isotypes in UPS F-941258! and H!).

Diagnosis. Macromorphology similar to *Unilacryma unispora*, but basidia developing two sterigmata and thus producing two basidiospores, which are normally narrowly ovoid to ellipsoid, (7.2–)8.2–10.4(–12.4) µm wide, Q = (1.2–)1.4–1.9(–2.2), instead of ovoid to broadly ellipsoid.

Basidiocarps ± gelatinous, amber-coloured to brownish, xerotolerant, isolated, gregarious or crowded, initially erumpent, pulvinate, tardily becoming depressed-pulvinate, without a conspicuous margin, 0.3–1(–1.5) mm diam. *Cortical/marginal hyphae* slightly to clearly thick-walled, with cylindrical or attenuate terminal cells, (3–)3.5–7 µm wide, unbranched or rarely sparingly branched, with sparse anastomoses and both clamped and clampless septa. *Internal hyphae* 2.5–6 µm wide, slightly to clearly thick-walled, embedded in a gelatinous matrix, with clamp connections but also frequently with some simple septa. *Hymenium* almost amphigenous, progressively forming numerous basidia intermixed with some hyphidia. *Subhymenium* with thin to slightly thick-walled, clamped hyphae. *Hyphidia* hyaline, sparsely to heavily branched, 2.5–4(–4.5) µm thick in the upper half, terminal branches (1–)1.5–2.5 µm thick, often with clamped septa throughout their length. *Young basidia* narrowly claviform to subcylindric, tapering towards the base, 6.5–8.5(–9) × 65–91 µm when sterigmata start to grow. *Mature basidia* developing two sterigmata 19–33 µm long, often forming an obtuse angle, sometimes acute or with an apical papilla at the basidium apex. *Basidiospores* (10.4–)12.4–18.2(–19.7) × (7.2–)8.2–10.4(–12.4) µm, Q = (1.2–)1.4–1.9(–2.2), narrowly ovoid to ellipsoid, sometimes subfusiform, rarely ovoid or subcylindric, with a conspicuous 1.2–1.6 µm long hilar appendix, becoming transversely 1-septate and tardily developing additional oblique, longitudinal and transverse septa, then becoming muriform, with a single hardly visible nucleus when recently discharged and still aseptate. *Basidium* and *basidiospore cytoplasm* hyaline or with a very pale yellowish tinge, with numerous small lipid bodies that slowly coalesce in dead state to form conspicuous lipid drops. Basidiospores germinating by the production of hyphae or microconidia. *Microconidia* narrowly ellipsoid to cylindric, about 3–4(–5) × 1.2–1.8 µm, hyaline.

Ecology & Distribution — *Unilacryma bispora* is a very common species that has only been observed fruiting on decorticated, old and frequently exposed wood of *Pinus sylvestris*,

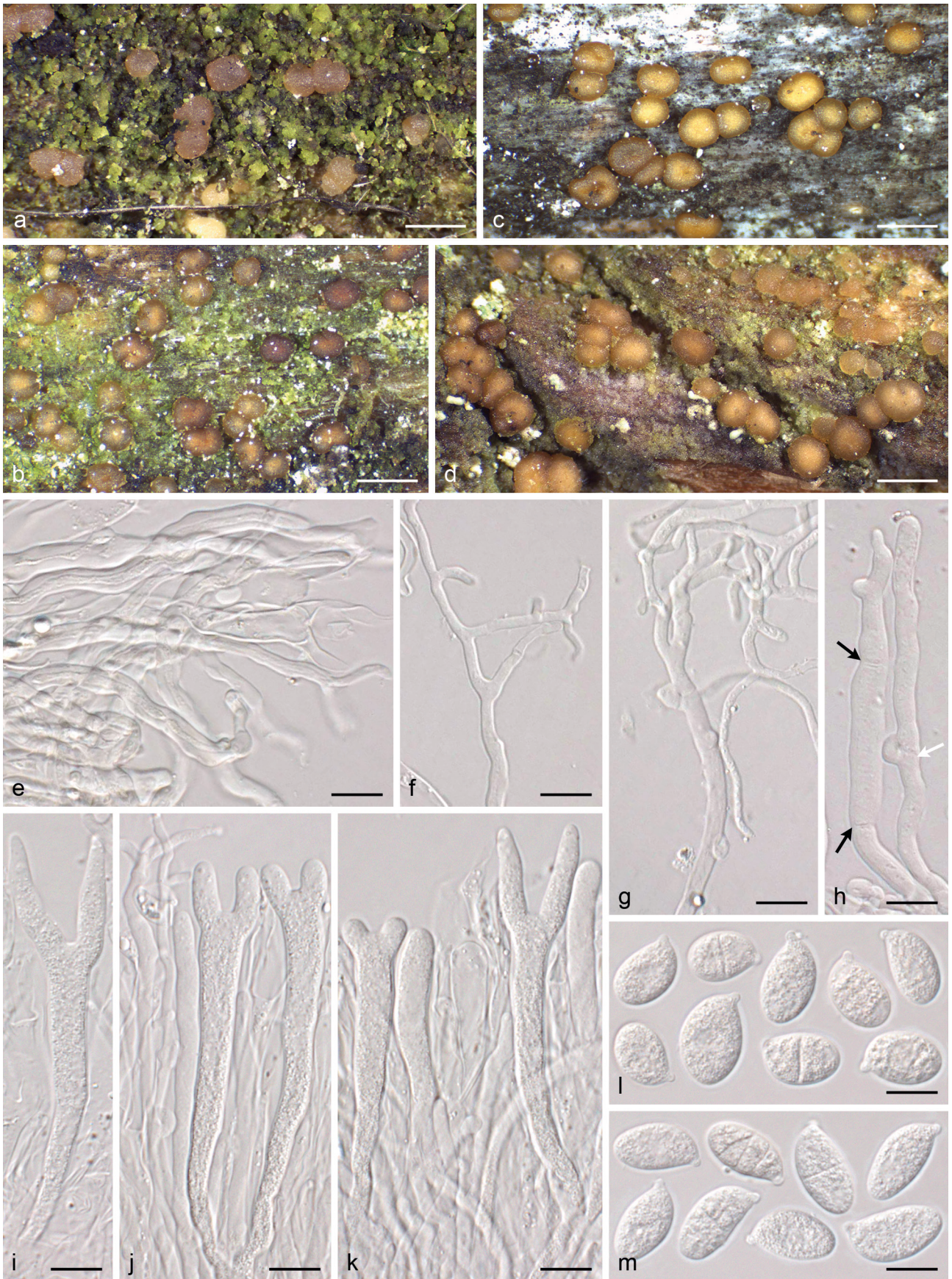


Fig. 14 *Unilacryma bispora*. a–d. Fruitbodies in different degrees of development, all in fresh condition; e–m. Micromorphology; e. poorly differentiated marginal hyphae; f–g. branched hyphidia; h. terminal cells of marginal hyphae, note the presence of both claspless (black arrows) and clamped (white arrow) septa; i. basidium; j–k. hymenium showing young and mature basidia and hyphidia; l–m. basidiospores. a–b, e–g, i–l. From the holotype; c. from UPS F-941274; d, h. from UPS F-941259; m. from the isotype. — Scale bars: a–d = 1 mm, e–m = 10 μ m.

especially on branches still attached to living trees. It occurs in both temperate and boreal forests, but it seems to be particularly common in the boreal biome, reaching the northern boreal zone. It is currently known from Finland, Norway, Russia, and Sweden. In a study on the fungal communities inhabiting *Picea abies* stumps (Kubart et al. 2016), environmental sequences probably related to this species were found in three Swedish nature reserves (Altarliden, Västerbotten; Fänstjärnskogen, Värmland; Ugglehöjden, Närke). The representative ITS sequence deposited in GenBank (KM494136) is identical to some of the ITS sequences of *U. bispora* generated in this study. Based on the results from the species delimitation analyses and considering the observed distribution in the field, we consider it very likely that those environmental sequences indicate that the species has a wider ecology than we have observed.

Nomenclature & Taxonomy — *Unilacryma bispora* is readily distinguished by the combination of small, pulvinate and often amber-coloured basidiocarps, 2-spored basidia, ovoid to ellipsoid basidiospores that become muriform when mature, branched hyphidia, and absence of obvious carotenoids under the light microscope. It is similar to *U. unispورا* except for the slightly broader, bisterigmate basidia, the slightly narrower and more elongate basidiospores, and the more northern distribution. The basidiospore shape is nevertheless somewhat plastic, and some specimens have predominantly ovoid basidiospores (e.g., UPS F-941254), while others show mostly ellipsoid ones (e.g., UPS F-941255). There may be little overlap among the length/width ratio of the basidiospores in certain specimens, but since most of them show intermediate shapes, and we have not detected any other differences, this should be regarded as part of the intraspecific variation. The holotype and the isotypes were collected on different, unconnected pine branches and are different individuals of the same gathering (ICN, Art. 8.2).

Following the literature and published identification keys (e.g., Torkelsen 1997), it might be confused with *Dacrymyces ovisporus*, a species with somewhat similar basidia and basidiospores, but with clearly larger, yellow-orange basidiocarps (Fig. 1f), and with very obvious carotenoid contents in the cytoplasm of all cells, including the basidiospores (Fig. 5s). The comments by Oberwinkler (2014) on '*Dacrymyces ovisporus*', indicating a macromorphology identical to that of *Unilacryma unispورا*, and a very close micromorphology, might mean that the specimens studied by Oberwinkler actually belong to *U. bispora*. In its most commonly accepted sense, *Dacrymyces ovisporus* is additionally distinguished by often staining the substrate yellow (Torkelsen 1997), by hyphidia that are simple or rarely with 1–2 short branches (Fig. 2l), and by both the basidiospore shape and septation, which is more similar to that of *U. unispورا* (thus, more rounded and becoming muriform earlier), this last character being well represented in the original plate of the protologue (Brefeld 1888). Further details on *D. ovisporus* will be provided in a forthcoming study. The two species are distantly related according to our phylogeny, belonging to two different families (Fig. 8–9).

Additional specimens examined. FINLAND, Nylandia, Espoo, near Nuuskio National Park, on a decorticated *Pinus sylvestris* branch, 12 May 2018, J.C. Zamora & P. Posio (UPS F-941270); Tavastia australis, Tammela, near Saaren Kansanpuisto, on *Pinus sylvestris* branch, 11 May 2018, J.C. Zamora & P. Posio (UPS F-941271); Ostrobothnia ultima, Rovaniemi, N66.49917° E25.767530°, on *Pinus sylvestris* branches, 14 Aug. 2018, J.C. Zamora (UPS F-941272); Regio Kuusamoensis, Salla, Oulanka, Urriavantie, N66.439727° E29.165568°, on *Pinus sylvestris* branch, 17 Aug. 2018, J.C. Zamora (UPS F-941274, UPS F-941275); Lapponia sompiensis, Pelkosenniemi, Saunavaara, N67.049234° E27.524627°, on *Pinus sylvestris* branches, 16 Aug. 2018, J.C. Zamora & P. Posio (UPS F-941273). — NORWAY, Nord-Trøndelag, Meråker, on *Pinus sylvestris* branches, 24 July 2018, J.C. Zamora, M. Svensson & M. Westberg (UPS F-941276). — RUSSIA, Karelia onegensis, Mdvezh' yegorsk regio, ad fontis fluminis Vitshka (UTM WQ3), on the roadside, hard, old wood, 21 June 1942, Matti Laurila (H 6020982).

— SWEDEN, Öland, Böda, Bökakusten västra naturreservat, Byrums Sandvik, N57.236677° E16.963497°, on *Pinus sylvestris* wood, 02 Oct. 2017, J.C. Zamora (UPS F-941268, UPS F-941269, duplicate in H); Södermanland, Stockholm, Skärholmen, N59.301129° E17.934630°, on *Pinus sylvestris* wood, 06 Apr. 2019, J.C. Zamora, P. Posio & J. Nieminen (UPS F-941263); Värmland, Lekvattnet, N60.172306° E12.699782°, on *Pinus sylvestris* branch, 07 Oct. 2018, J.C. Zamora (UPS F-941267); Värmland, Ransäter, Munkfors V, N59.814609° E13.390181°, on *Pinus sylvestris* branch, 08 June 2019, J.C. Zamora (UPS F-941467); Uppland, Älvkarleby, Långsandsörarna island, N60.65783° E17.55504°, on *Pinus sylvestris* branches, 17 Apr. 2018, J.C. Zamora, S. Ekman & M. Zuluaga (UPS F-941262); Uppland, Solna, Haga, coniferous wood, 01 July 2018, J.C. Zamora & I. Olariaga (UPS F-941261); Uppland, Uppsala, Norra Lunsen, N59.80680° E17.69804°, on old *Pinus sylvestris* wood, 20 Aug. 2017, J.C. Zamora (UPS F-941255, UPS F-941256); Uppland, Uppsala, Lunsen, on *Pinus sylvestris* branch, 08 Apr. 2018, J.C. Zamora (UPS F-941259); Uppland, Uppsala, Norra Lunsen, N59.790953° E17.687811°, on *Pinus sylvestris* branch, 30 Sept. 2018, J.C. Zamora, S. Ekman, A. Kruys & D. Pfister (UPS F-941260); Uppland, Uppsala, Stadsskogen naturreservat, N59.84234° E17.62527°, on highly decomposed, coniferous wood, 13 Aug. 2017, J.C. Zamora (UPS F-941254); Jämtland, Åre, Handöl, N63.245084° E12.444898°, on *Pinus sylvestris* branches, 24 July 2018, J.C. Zamora (UPS F-941265); Jämtland, Åre, N63.23271° E12.42884°, on *Pinus sylvestris* wood, 25 July 2018, J.C. Zamora (UPS F-941266); Jämtland, Mörsil, Klukshäckren, N63.172614° E13.600665°, on *Pinus sylvestris* branches, 23 July 2018, J.C. Zamora (UPS F-941264, duplicate in H); Norrbotten, Övertorneå, Luppiberget, N66.313259° E23.608372°, 13 Aug. 2019, J.C. Zamora & P. Posio (UPS F-945010).

Unilacryma unispورا (L.S. Olive) Shirouzu, Tokum. & Oberw., Mycologia 105: 1120. 2013 s.lat. — Fig. 15

≡ *Platyglœa unispورا* L.S. Olive, J. Elisha Mitchell Sci. Soc. 74: 41. 1958 [basion.].

— *Achroomyces unisporus* (L.S. Olive) Wojewoda, Mala Fl. Grzybow 2: 205. 1981, nom. inval. (Art. 6.10, Art. 41.8 does not apply).

— *Dacrymyces unisporus* (L.S. Olive) K. Wells, Mycologia 86: 31. 1994, nom. inval. (see above).

Typus. UNITED STATES OF AMERICA, North Carolina, Orange, [Chapel Hill], Battle Park, 11 Dec. 1943, L.S. Olive s.n. (NCU-F-0026842! holotype).

Basidiocarps ± gelatinous, amber-coloured to brownish, xerophilic, isolated, gregarious or crowded, initially erumpent, pulvinate, tardily becoming depressed-pulvinate, without a conspicuous margin, 0.3–1.0(–1.2) mm diam. **Cortical/marginal hyphae** slightly to clearly thick-walled, with cylindrical or attenuate terminal cells, 4–7 µm wide, unbranched or rarely sparingly branched, with sparse anastomoses and both clamped and clampless septa. **Internal hyphae** 3–7 µm wide, slightly to clearly thick-walled, embedded in a gelatinous matrix, with clamp connections but also frequently with some simple septa. **Hymenium** almost amphigenous, progressively forming numerous basidia intermixed with some hyphidia. **Subhymenium** with thin to slightly thick-walled, clamped hyphae. **Hyphidia** hyaline, sparsely to heavily branched, (2.5–)3–4.5 µm thick in the upper half, terminal branches (1–)1.5–3 µm thick, often with clamped septa throughout their length. **Young basidia** initials cylindrical to narrowly claviform, 5–7.5 µm thick when close to maturity. **Mature basidia** developing a single thick sterigma (very rarely two irregularly formed sterigma-like structures may be seen), about 18–32 µm long, bearing a single terminal and asymmetrically attached basidiospore. Total basidium length reaching 70–130 µm. **Basidiospores** (11.9–)14–17.4(–19.5) × (10.2–)11.2–13.5(–15.7) µm, Q = (1.1–)1.2–1.4(–1.6), ovoid to broadly ellipsoid, rarely globose or ellipsoid, with a conspicuous 1.0–1.5 µm long hilar appendix, becoming transversely 1-septate and developing additional oblique, longitudinal and transverse septa, then becoming muriform, with a single hardly visible nucleus when still aseptate. **Basidium** and **basidiospore cytoplasm** hyaline or with a very pale yellowish tinge, with numerous small lipid bodies that slowly coalesce in dead state to form conspicuous lipid drops. Basidiospores germinating by the production of hyphae or microconidia. **Microconidia** rarely

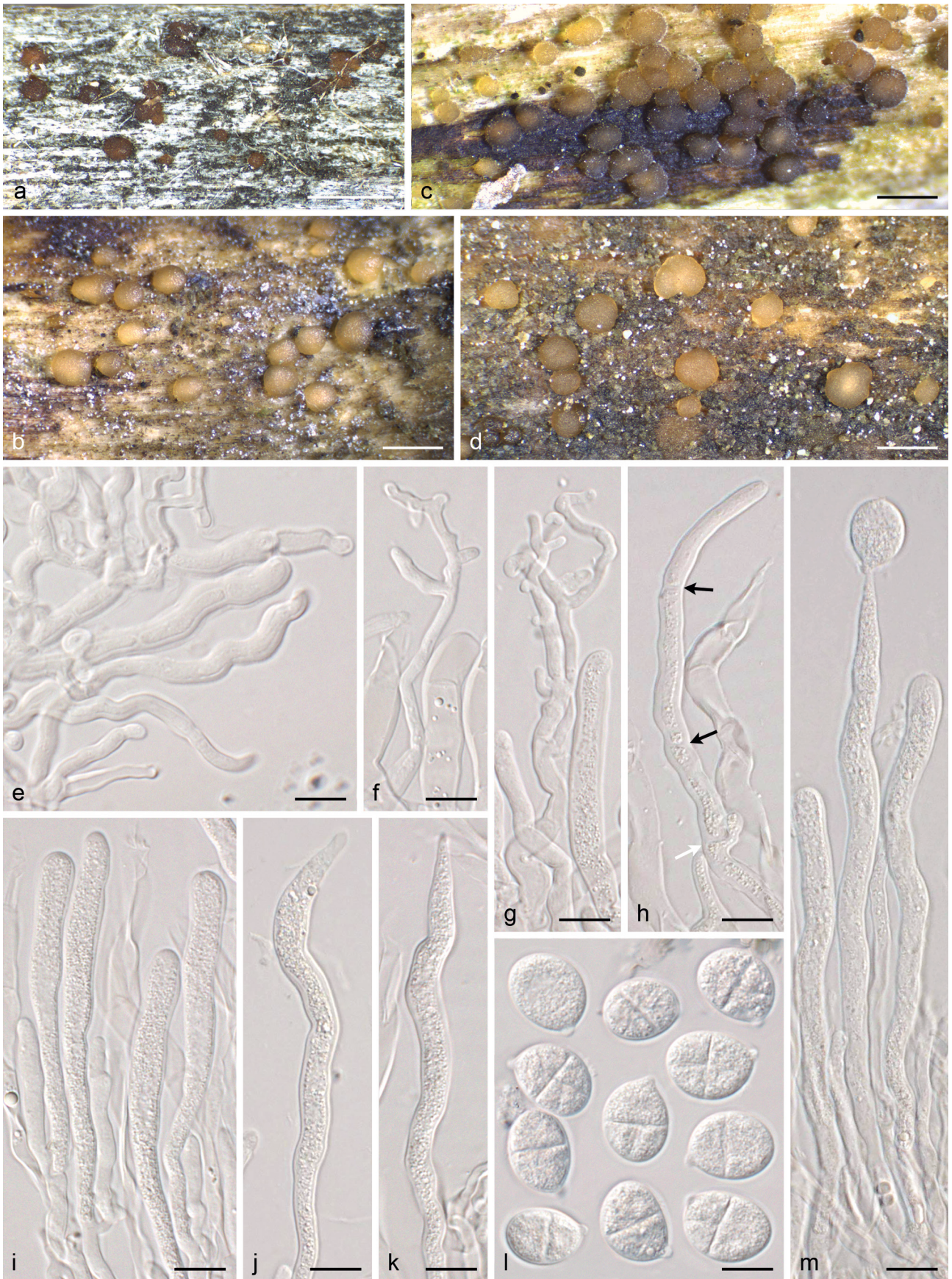


Fig. 15 *Unilacryma unispora* s.lat. a–d. Fruitbodies in different degrees of development, all in fresh condition except a; e–m. Micromorphology; e. marginal hyphae; f–g. branched hyphidia; h. terminal cells of marginal hyphae, note the presence of both clampless (black arrows) and clamped (white arrow) septa; i. young basidia; j–k. mature basidia; l. basidiospores; m. two young basidia and one mature basidium bearing a single sterigma with a terminal basidiospore. a. From the holotype; b, e–g, j–k. from UPS F-941279; c. from UPS F-941286; d. from UPS F-941284; h–i, m. from UPS F-941283; l. from UPS F-941280. — Scale bars: a–d = 1 mm, e–m = 10 μ m.

observed, narrowly ellipsoid to cylindrical, about 3–4 × 1.5 µm, hyaline.

Ecology & Distribution — Very common, frequently growing on decorticated, old and rather exposed wood of *Juniperus* spp. and *Pinus* spp., and particularly common on dead branches still attached to trees. In areas where the distributions of both *Unilacryma* species overlap, they may grow together on the same branch and even intermixed. In Europe, *U. unispora* has been previously reported only from The Netherlands by Dam & Dam (2014). In a wide sense, it is broadly distributed in the Holarctic temperate biome (North America, East Asia, and Europe), and extends into the southern boreal zone, but it has not been found in the middle and northern boreal forests of Sweden and Finland where *U. bispora* is common.

Nomenclature & Taxonomy — When describing *Platygløea unispora*, Olive (1944) failed to include the required Latin diagnosis, resulting in an invalid name (ICN, Art. 39.1). The taxon was validated by Olive (1958), who provided the required diagnosis as well as a holotype. Olive (1944) is often incorrectly cited as the place of publication of the basionym, which renders the intended new combinations in *Achroomyces* and *Dacrymyces* invalid, none of the requirements of Art. 41.8 being fulfilled.

In a broad taxonomic sense, *U. unispora* is well distinguished by 1-spored basidia and subglobose to ovoid basidiospores that become transversely septate and finally muriform with age. The macromorphology is nearly identical to that of *U. bispora*, so a micromorphological study of the samples is always needed to identify them. The European specimens agree morphologically very well with the description of Olive (1944) and additional descriptions and iconography of North American specimens (e.g., Wells 1994), as well as with the North American specimens we studied, including the holotype. Unfortunately, we were not able to obtain DNA sequence data of the studied North American specimens, not even after using specific primers to amplify short fragments of the ITS. The description of the Japanese specimens by Shirouzu et al. (2009) also agrees relatively well, although they reported basidia and basidiospores to be on average slightly smaller, marginal hyphae as thin-walled, and branched hyphidia as absent. In addition, the samples from North Europe and East Asia form two different clades (Fig. 10). Nevertheless, with the current morphological and genetic data, it is unclear whether these clades represent two distinct species. Therefore, we assume here a conservative approach and treat the North American, East Asian, and European specimens as different populations of a single species.

Additional specimens examined. FINLAND, Alandia, Kökar, Hamno, N59.947100° E20.875280°, on *Juniperus communis* branch, 22 July 2019, J.C. Zamora & P. Posio (UPS F-945056); Regio aboensis, Nauvo, N60.192401° E21.912373°, on *Pinus sylvestris* branch, 21 July 2019, J.C. Zamora & P. Posio (UPS F-945073); Nylandia, Espoo, near Nuukio National park, on *Pinus sylvestris* branch, 12 May 2018, J.C. Zamora & P. Posio (UPS F-941295); Nylandia, Helsinki, Veräjämäki, old, rather moist forest depression, on a fallen branch (*Pinus sylvestris*, decay stage 3, 3 cm diam), N60.22390° E24.97607°, elev. 29 m, 25 May 2010, O. Miettinen 14064 (H 6012675); Tavastia australis, Tammela, near Saaren Kansapuis-to, on *Pinus sylvestris* branch, 11 May 2018, J.C. Zamora & P. Posio (UPS F-941298); Tavastia australis, Tammela, Mustiala, N60.816501° E23.764646°, on *Pinus sylvestris* branches, 11 May 2018, J.C. Zamora & P. Posio (UPS F-941296); Tavastia australis, Tammela, Mustiala, Salois (Saloinen), around P.A. Karsten's house, on *Pinus sylvestris* branch, 11 May 2018, J.C. Zamora & P. Posio (UPS F-941297). — NORWAY, Hedmark, Kongsvinger, Austmarka, N60.106530° E12.358369°, on *Pinus sylvestris* branch, 06 Oct. 2018, J.C. Zamora & P. Posio (UPS F-941293); Hedmark, Kongsvinger, Kubergskogen, N60.236862° E12.426987°, on *Pinus sylvestris* branch, 06 Oct. 2018, J.C. Zamora & P. Posio (UPS F-941294); Troms, Storfjord, Skibotn, N69.389745° E20.296217°, on *Pinus sylvestris* wood, 16 Aug. 2019, J.C. Zamora & P. Posio (UPS F-945011). — SWEDEN, Öland, Böda, Bökadusten västra naturreservat, Byrums Sandvik, N57.236677° E16.963497°, on *Pinus sylvestris* branches, 02 Oct. 2017, J.C. Zamora (UPS F-941291, duplicate in H);

Öland, Böda, Lindreservatet, N57.319448° E17.030849°, on *Pinus sylvestris* wood, 03 Oct. 2017, J.C. Zamora (UPS F-941292); Södermanland, Nacka, Nackareservatet, N59.28955° E18.17560°, on *Pinus sylvestris* branches, 10 June 2018, J.C. Zamora, P. Posio & J. Nieminen (UPS F-941288); Värmland, Lekvattnet, N60.172306° E12.699782°, on *Pinus sylvestris* branch, 07 Oct. 2018, J.C. Zamora (UPS F-941290); Värmland, Ransäter, Munkfors V, N59.814677° E13.391138°, on *Pinus sylvestris* branches, 08 June 2019, J.C. Zamora (UPS F-941466); Uppland, Älvkarleby, Långsandsörarna island, N60.65783° E17.55504°, on *Pinus sylvestris* branches, 17 Apr. 2018, J.C. Zamora, S. Ekman & M. Zuluaga (UPS F-941283); Uppland, Funbo, c. 400 m north of Fjällnora farm, N59.840762° E17.913327°, on *Pinus sylvestris* branch, 14 Nov. 2017, J.C. Zamora & S. Ekman (UPS F-941284); Uppland, Jumkil, Granslätt, N59.95312° E17.27913°, on old pieces of a rabbit cage, made of unidentified (presumably coniferous) wood, 12 May 2017, J.C. Zamora, S. Ekman & M. Westberg (UPS F-941277); Uppland, Solna, Bergshamra, N59.38358° E18.03756°, on *Pinus sylvestris* branch, 23 June 2018, J.C. Zamora (UPS F-941286); Uppland, Solna, Hagaparken, N59.365065° E18.032583°, on unidentified weathered wood, probably *Pinus sylvestris*, 28 Apr. 2019, J.C. Zamora (UPS F-941287); Uppland, Uppsala, Norra Lunsen, N59.80680° E17.69804°, on *Pinus sylvestris* branches, 20 Aug. 2017, J.C. Zamora (UPS F-941279, duplicate in H); Uppland, Uppsala, Norra Lunsen naturreservat, N59.79390° E17.68667°, on *Pinus sylvestris* branches, 19 Nov. 2017, J.C. Zamora (UPS F-941280); Uppland, Uppsala, Norra Lunsen naturreservat, on *Pinus sylvestris* branches, 08 Apr. 2018, J.C. Zamora (UPS F-941281); Uppland, Uppsala, Norra Lunsen naturreservat, N59.790953° E16.687811°, on *Pinus sylvestris* branches, 30 Sept. 2018, J.C. Zamora (UPS F-941282); Uppland, Uppsala, Stadsskogen naturreservat, N59.84234° E17.62527°, on highly decomposed *Pinus sylvestris* wood, 13 Aug. 2017, J.C. Zamora (UPS F-941278); Hälsingland, Gnarp, N62.037699° E17.154672°, on exposed *Pinus sylvestris* branch, 27 July 2018, J.C. Zamora (UPS F-941289). — UNITED STATES OF AMERICA, North Carolina, Orange, University Lake area, near Chapel Hill, *Juniperus virginiana*, 14 Apr. 1982, L.S. Olive NC82-40 (NCU F-0026843).

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