

Evolution of lifestyles in Capnodiales

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Abstract: The Capnodiales, which includes fungi known as the sooty moulds, represents the second largest order in *Dothideomycetes*, encompassing morphologically and ecologically diverse fungi with different lifestyles and modes of nutrition. They include saprobes, plant and human pathogens, mycoparasites, rock-inhabiting fungi (RIF), lichenised, epi-, ecto- and endophytes. The aim of this study was to elucidate the lifestyles and evolutionary patterns of the *Capnodiales* as well as to reconsider their phylogeny by including numerous new collections of sooty moulds, and using four nuclear loci, LSU, ITS, *TEF-1a* and *RPB2*. Based on the phylogenetic results, combined with morphology and ecology, *Capnodiales s. lat.* is shown to be polyphyletic, representing seven different orders. The sooty moulds are restricted to *Capnodiales s. str.*, while *Mycosphaerellales* is resurrected, and five new orders including *Cladosporiales, Comminutisporales, Neophaeothecales, Phaeothecales* and *Racodiales* are introduced. Four families, three genera, 21 species and five combinations are introduced as new. Furthermore, ancestral reconstruction analysis revealed that the saprobic lifestyle is a primitive state in *Capnodiales s. lat.*, and that several transitions have occurred to evolve lichenised, plant and human parasitic, ectophytic (sooty blotch and flyspeck) and more recently epiphytic (sooty mould) lifestyles.

Key words: Capnodiales, Cladosporium, Mycosphaerella, Multigene phylogeny, Sooty moulds.

Taxonomic novelties: New orders: Cladosporiales Abdollahz. & Crous, Comminutisporales Abdollahz. & Crous, Neophaeothecales Abdollahz. & Crous, Phaeothecales Abdollahz. & Crous, Racodiales Abdollahz. & Crous.

New families: Comminutisporaceae Abdollahz. & Crous, Neoantennariellaceae Abdollahz. & Crous, Neophaeothecaceae Abdollahz. & Crous, Readerielliopsidaceae Abdollahz. & Crous.

New genera: Neoantennariella Abdollahz. & Crous, Neoasbolisia Abdollahz. & Crous, Neophaeotheca Abdollahz. & Crous.

New species: Capnodium alfenasii Abdollahz. & Crous, Capnodium blackwelliae Abdollahz. & Crous, Capnodium gamsii Abdollahz. & Crous, Capnodium neocoffeicola Abdollahz. & Crous, Capnodium paracoffeicola Abdollahz. & Crous, Chaetocapnodium summerellii Abdollahz. & Crous, Chaetocapnodium indonesiacum Abdollahz. & Crous, Chaetocapnodium insulare Abdollahz. & Crous, Chaetocapnodium tanzanicum Abdollahz. & Crous, Chaetocapnodium thailandense Abdollahz. & Crous, Leptoxyphium citri Abdollahz. & Crous, Neoantennariella phylicae Abdollahz. & Crous, Neoasbolisia phylicae Abdollahz. & Crous, Phaeoxyphiella australiana Abdollahz. & Crous, Phaeoxyphiella phylicae Abdollahz. & Crous, Scolecoxyphium blechni Abdollahz. & Crous, Scolecoxyphium blechnicola Abdollahz. & Crous, Scolecoxyphium leucadendri Abdollahz. & Crous, Scolecoxyphium phylicae Abdollahz. & Crous, Scorias camelliae Abdollahz. & Crous.

New combinations: Chaetocapnodium philippinense (Hongsanan & K.D. Hyde) Abdollahz. & Crous, Chaetocapnodium placitae (Cheewangkoon & Crous) Abdollahz. & Crous, Neophaeotheca salicorniae (Crous & Roets) Abdollahz. & Crous, Neophaeotheca triangularis (de Hoog & Beguin) Abdollahz. & Crous, Phragmocapnias plumeriae (Hongsanan & K.D. Hyde) Abdollahz. & Crous.

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INTRODUCTION

The Dothideomycetes represents a class of ecologically diverse and cosmopolitan fungi from aquatic to terrestrial ecosystems. Diverse lifestyles are found amongst the Dothideomycetes including epiphytes, endophytes, saprobes, plant and animal pathogens, mycoparasites, mycorrhizal, lichenised and rockinhabiting fungi (Schoch et al. 2009, Schoch & Grube 2015, Ametrano et al. 2019). The Dothideomycetes is divided into two subclasses, Pleosporomycetidae and Dothideomycetidae, and some incertae sedis lineages, accommodating more than 25 orders, 110 families and over 19000 species, thereby representing the largest class of Ascomycota (Schoch et al. 2009, Hyde et al. 2013, Jaklitsch et al. 2015, Schoch & Grube 2015, Van Nieuwenhuijzen et al. 2016, Bezerra et al. 2017, Videira et al. 2017, Wijayawardene et al. 2017). Morphologically they are mostly characterised by ascostromatic development and bitunicate asci with fissitunicate dehiscence (Schoch & Grube 2015).

The Capnodiales represent the second largest order in *Dothideomycetes* after the *Pleosporales*. The *Capnodiales* is included in the subclass *Dothideomycetidae* along with the *Dothideales* and *Myrangiales* (Crous *et al.* 2009). The taxonomic concept of this order was expanded from the original description by Luttrell (1955), based on a multigene phylogeny and the presence of ostiolar periphyses as a synapomorphic feature (Schoch *et al.* 2006). Taxa in this order lack pseudoparaphyses, but include several species with periphysoids and periphyses (Lumbsch & Lindemuth 2001).

As discussed by Schoch & Grube (2015), the Capnodiales was established based on the sooty moulds in three families, *Antennulariaceae*, *Capnodiaceae*, and *Coccodiniaceae*. However, phylogenetic analyses revealed that the sooty moulds are polyphyletic and include species residing in two different classes, *Dothideomycetes* and *Eurotiomycetes* (Crous *et al.* 2007a).

The Capnodiales now includes the epiphytic sooty moulds associated with honeydew produced by insects

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Table 1. Details of the sooty mould isolates included in this study. Type cultures and sequences generated in this study are in bold face.								
Family and Species	Voucher/Culture ¹	Substrate/Lifestyle ²	Country/ Location	Collector	GenBank accession numbers ³			
name					LSU	ITS	TEF-1α	RPB2
Capnodiaceae								
Capnodium alfenasii	CBS 146151 = CPC 22666	<i>Tabebuia</i> sp.	Brazil	A.C. Alfenas	MN74916	5 MN74923	3 MN82934	6 MN829260
	CBS 146152 = CPC 22667	<i>Tabebuia</i> sp.	Brazil	A.C. Alfenas	MN749166	6 MN749234	MN829347	7 MN829261
Ca. blackwelliae	CBS 133588 = CPC 14327	Myrtus communis	USA	P.W. Crous	MH878118	B MN74923	5 GU349054	4 GU371743
Ca. coartatum	MFLUCC10-0069	Psidium sp.	Thailand	P. Chomnunti	JN832614	-	-	-
	MFLUCC10-0070	-	Thailand	P. Chomnunti	JN832615	-	-	-
	CPC 17779	Alstonia scholaris	Thailand	K.D. Hyde	MN749167	7 MN74923	6 MN829348	8 MN829262
Ca. coffeae	CBS 147.52 = AFTOL-ID 939	Coffea robusta	Zaire	Deposited by J. Nicot/Isolated by A. Saccas	GU214400) DQ491515	5 DQ471089	9 KT216519
Ca. coffeicola	MFLUCC15-0206	Coffea sp.	Thailand	S. Hongsanan	KU358920	KU358921	-	-
Ca. gamsii	CBS 892.73	Sooty mould, on unknown leaf	Sri Lanka	W. Gams	GU301847	MN74923	GU34904	5 GU371736
	CBS 146153 = CPC 17765	Lagerstroemia speciosa	Thailand	K.D. Hyde	MN749168	3 MN749238	3 MN829349	9 MN829263
	MFLUCC10-0066	-	Thailand	S.K. Chandranath	JN832613	-	-	-
	CBS 146154 = CPC 20466 = MFLUCC12 0101	- Lagerstroemia floribunda	Thailand	S. Hongsanan	MN749169	9 MN749239) MN82935(0 MN829264
	CBS 146155 = CPC 20467 = MFLUCC12 0102	- Lagerstroemia floribunda	Thailand	S. Hongsanan	MN749170) MN749240) MN82935 [,]	1 MN829265
	CBS 146156 = CPC 20471 = MFLUCC12 0107	- Living leaf of unknown host	Thailand	S. Hongsanan	MN749171	I MN74924 [,]	MN829352	2 MN829266
Ca. neocoffeicola	CBS 139614 = MFLUCC14-0570	Coffea arabica	Thailand	S. Hongsanan	MN749172	2 MN749242	2 MN82935	3 MN829267
	CBS 139613 = MFLUCC14-0569	Coffea arabica	Thailand	S. Hongsanan	MN749173	3 MN749243	3 MN829354	4 MN829268
Ca. paracoffeicola	CBS 139616 = MFLUCC 14-0572	Coffea arabica	Thailand	S. Hongsanan	MN749174	1 MN74924	MN82935	5 MN829269
	CBS 139615 = MFLUCC14-0571	Coffea arabica	Thailand	S. Hongsanan	MN74917	5 MN74924	5 MN82935	6 MN829270
Chaetocapnodium indonesiacum	CBS 202.30	Camelia sinensis	Indonesia	Deposited by F.H. van Beyma/Isolated by Steinmann	GU301849	9 MH855113	3 GU349060) MN829273
Ch. insulare	CBS 146159 = CPC 19221	Phylica arborea	South Africa	M.J. Wingfield	MN749178	3 MN749248	8 MN829359	9 MN829274
	CBS 146160 = CPC 19223	Phylica arborea	South Africa	M.J. Wingfield	MN749179	9 MN749249	MN82936	0 MN829275
	CBS 146161 = CPC 19224	Phylica arborea	South Africa	M.J. Wingfield	MN749180) MN74925) MN82936 [,]	1 MN829276
Ch. philippinense	MFLUCC12-0110 = CPC 20474	Palm	Philippines	K.D. Hyde	KP744503	MN74925	MN829362	2 MN829277
Ch. placitae	CBS 124758 = CPC 13706	Eucalyptus placita	Australia	B.A. Summerell	GQ303299	9 GQ303268	3 MN82936	3 MN829278
Ch. siamensis	MFLUCC13-0778	Leaves of unidentified plant	Thailand	S. Hongsanan	KP744479	_	_	_
	CBS 139815 = MFLUCC13-0096	Leaves of unidentified plant	Thailand	S.C. Karunarathna	MN74918 ²	I MN749252	2 MN829364	4 MN829279
Ch. summerellii	CBS 146157 = CPC 13654	Eucalyptus placita	Australia	B.A. Summerell	MN749176	6 MN749240	6 MN829357	7 MN829271

Table 1. (Continued).

Family and Species name	Voucher/Culture ¹	Substrate/Lifestyle ²	Country/	Collector	GenBank accession numbers ³			
			Location		LSU ITS TEF-1a RPB2			
	CBS 146158 = CPC 17368	_	Laos	P. Pheng	MN749177 MN749247 MN829358 MN829272			
Ch. tanzanicum	CBS 145.79	Lichen	Tanzania	-	MN749182 MN749253 MN829365 MN829280			
Ch. thailandense	CBS 139619 = MFLUCC13-0787	_	Thailand	S.C. Karunarathna	MN749183 MN749254 MN829366 MN829281			
Conidiocarpus asiticus	MFLUCC10-0062	Coffea arabica	Thailand	J.K. Liu	JN832612 KU358924 – –			
Co. caucasicus	GUMH 937	Citrus sinensis	Iran	F. Byrami	KC833050 – – –			
Co. siamensis	MFLUCC10-0064	Mangifera indica	Thailand	R. Phokhomsak	JN832609 – – –			
Co. siamensis	MFLUCC10-0061	_	Thailand	P. Chomnunti	JN832607 KU358923 – –			
Co. siamensis	MFLUCC10-0063	Coffea arabica	Thailand	J.K. Liu	JN832608 KU358925 – –			
Conidiocarpus sp.	CPC 17778	Guave sp.	Thailand	K.D. Hyde	MN749185 MN749256 MN829368 MN829283			
	CPC 20463 = MFLUCC12-0098	<i>Malus</i> sp.	Thailand	W. Saowanee	MN749187 MN749258 MN829370 MN829285			
	CPC 20464 = MFLUCC12-0099	Mimusops elengi	Thailand	S. Hongsanan	MN749194 MN749265 MN829377 MN829292			
	CPC 20465 = MFLUCC12-0100	Mimusops elengi	Thailand	S. Hongsanan	MN749191 MN749262 MN829374 MN829289			
	CPC 20468 = MFLUCC12-0103	Mango	Thailand	Puttaluk	MN749193 MN749264 MN829376 MN829291			
	CPC 20472 = MFLUCC12-0108	Living leaf of unknown host	Thailand	S. Hongsanan	MN749188 MN749259 MN829371 MN829286			
	CPC 21380 = MFLUCC12-0404	Malus sp.	Thailand	K.D. Hyde	MN749186 MN749257 MN829369 MN829284			
	CBS 139818 = MFLUCC14-0874	Coffea arabica	Thailand	S. Hongsanan	MN749190 MN749261 MN829373 MN829288			
	CBS 139819 = MFLUCC14-0875	Coffea arabica	Thailand	S. Hongsanan	MN749192 MN749263 MN829375 MN829290			
	CBS 139820 = MFLUCC 14-0876	Coffea arabica	Thailand	S. Hongsanan	MN749184 MN749255 MN829367 MN829282			
	CBS 139821 = MFLUCC14-0877	Coffea arabica	Thailand	S. Hongsanan	MN749189 MN749260 MN829372 MN829287			
Heteroconium citharexyli	HM628775	Citharexylum ilicifolium	Ecuador	H. Sydow	HM628775 HM628776 – –			
Leptoxyphium cacuminum	MFLUCC10-0059	Gossypium herbaceum	Thailand	S.C. Karunarathna	JN832603 – – –			
	MFLUCC10-0049	Mimusops elengi	Thailand	P. Chomnunti	JN832602 – – –			
	MFLUCC10-0086	Ficus sp.	Thailand	K.D. Hyde	JN832604 – – –			
L. citri	CBS 451.66	Citrus sinensis	Spain	H.A. van der Aa	KF902094 MN749266 GU349039 GU371727			
	CBS 146162 = CPC 26196	-	-	V. Guarnaccia	MN749195 MN749267 MN829378 MN829294			
L. glochidion	IFRDCC 2651	Glochidion wrightii	China	H. Yang	KF982308 KF982307 – –			
L. kurandae	CBS 129530 = CPC 17274	<i>Eucalyptus</i> sp.	Australia	P.W. Crous & R.G. Shivas	JF951170 JF951150 MN829379 MN829295			
L. madagascariense	CBS 124766 = CPC 14623	Eucalyptus camaldulensis	Madagascar	M.J. Wingfield	MH874923 MH863407 MN829380 MN829296			
Leptoxyphium sp.	CPC 17767	Gossypium herbaceum	Thailand	K.D. Hyde	MN749203 MN749275 MN829388 MN829304			
	CPC 20470 = MFLUCC12-0106	Living leaf of unknown host	Thailand	S. Hongsanan	MN749200 MN749272 MN829385 MN829301			
					(continued on next page)			

Table 1. (Continued).

Family and Species	Voucher/Culture ¹	Substrate/Lifestyle ²	Country/	Collector	GenBank accession numbers ³			
name			Location		LSU	ITS	_TEF-1α	RPB2
	CPC 20473 = MFLUCC12-0109	Living leaf of unknown host	Thailand	S. Hongsanan	MN74919	7 MN749269	MN829382	2 MN829298
	CPC 20481 = MFLUCC12-0118	Living leaf of unknown host	Thailand	-	MN749202	I MN749273	MN829386	6 MN829302
	CPC 21382 = MFLUCC12-0406	Heliconia sp.	Thailand	S. Hongsanan	MN749199	9 MN749271	MN829384	4 MN829300
	CPC 21383 = MFLUCC12-0407	Ixora chinensis	Thailand	S. Hongsanan	MN749202	2 MN749274	MN829387	7 MN829303
	CBS 123.26 = ATCC 11925 = IMI 0893	363 Hibiscus tiliaceus	Indonesia	Deposited by M.B. Schwarz	GU21443() MH854862	GU349051	I GU371741
	CBS 382.87	Citrus aurantium	India	Deposited and isolated by N.D. Sharma	MN74920	5 MN749277	MN829390) MN829306
	CBS 135836	Insect gut	India	S. Kajale & M. Sonawane	MN749200	6 MN749278	MN829391	1 MN829307
	CBS 139617 = MFLUCC13-0781	-	Thailand	S. Hongsanan	MN74919(6 MN749268	MN829381	1 MN829297
	CBS 139618 = MFLUCC13-0783	-	Thailand	S. Hongsanan	MN749204	4 MN749276	MN829389	9 MN829305
	CBS 139620 = MFLUCC13-0786	-	Thailand	S.C. Karunarathna	MN74920	7 MN749279	MN829392	2 MN829308
	CBS 139812 = MFLUCC13-0078	Living leaf of unknown host	Thailand	S.C. Karunarathna	MN749208	3 MN749280	MN829393	3 MN829309
	CBS 139814 = MFLUCC13-0790	Living leaf of unknown host	Thailand	S.C. Karunarathna	MN749198	3 MN749270	MN829383	3 MN829299
Phragmocapnias betle	CPC 17762	Mimusops elengi (Bullet wood)	Thailand	K.D. Hyde	MN74922	I MN749293	MN829407	7 MN829323
	CPC 20476 = MFLUCC12-0112	Palm	Philippines	K.D. Hyde	MN749222	2 MN749294	MN829408	3 MN829324
	CPC 21379 = MFLUCC12-0403	<i>Malus</i> sp.	Thailand	K.D. Hyde	MN749223	3 MN749295	5 MN829409	9 MN829325
	MFLUCC10-0053	<i>lxora</i> sp.	Thailand	P. Chomnunti	JN832606	KU358922	_	_
Ph. plumeriae	MFLUCC15-0205	<i>Plumeria</i> sp.	Thailand	C. Singhapop	KU358918	KU358919	_	_
Polychaeton citri	CBS 116435	Citrus aurantium	Iran	R. Zare & W. Gams	GU21446	GU214649	MN829394	4 MN829310
Neoantennariellaceae								
Fumiglobus pieridicola	UBC F23788	Pieris japonica	Canada	Tanay Bose	KC833052	KF263961	_	_
Neoantennariella phylicae	CBS 146164 = CPC 19227	Phyllica arborea	South Africa	M.J. Wingfield	MN749209	9 MN749281	MN82939	5 MN829311
	CBS 146165 = CPC 19977	Phylica arborea	UK	P. Ryan	MN749213	3 MN749285	5 MN829399	9 MN829315
	CBS 146166 = CPC 19981	Phylica arborea	UK	P. Ryan	MN749212	2 MN749284	MN829398	8 MN829314
	CBS 146167 = CPC 19985	Phylica arborea	UK	P. Ryan	MN74921() MN749282	MN829396	6 MN829312
	CPC 19992	Phylica arborea	UK	P. Ryan	MN749214	4 MN749286	6 MN829400	0 MN829316
	CBS 146163 = CPC 19989	Phylica arborea	UK	P. Ryan	MN74921	I MN749283	MN829397	7 MN829313
Neoasbolisia phylicae	CBS 146168 = CPC 19982	Phylica arborea	UK	P. Ryan	MN74921	5 MN749287	MN829401	1 MN829317
Readerielliopsidaceae								
"Capnodium" salicinum	CBS 131.34 = AFTOL-ID 937	Bursaria spinosa	Indonesia	Deposited by E.E. Fisher	EU019269	AJ244240	DQ677889	9 KT216553
Phaeoxyphiella australiana	CBS 146169 = CPC 29527	<i>Agonis</i> sp.	Australia	P.W. Crous	MN749220) MN749292	MN829406	6 MN829322
Ph. phylicae	CBS 146171 = CPC 19979	Phylica arborea	UK	P. Ryan	MN749216	6 MN749288	8 MN829402	2 MN829318

Table 1. (Continued).

Family and Spacios	Voucher/Culture ¹	Substrate/Lifestyle ²	Country/ Location	Collector	GenBank accession numbers ³				
name							TFF_1α	RPR2	
							<u>, , , , , , , , , , , , , , , , , , , </u>		
	CBS 146172 = CPC 19984	Phylica arborea	UK	P. Ryan	MN/4921/	MN/49289	MN829403	MN829319	
	CBS 146173 = CPC 19987	Phylica arborea	UK	P. Ryan	MN749218	3 MN749290	MN829404	MN829320	
	CBS 146170 = CPC 19993	Phylica arborea	UK	P. Ryan	MN749219	MN749291	MN829405	MN829321	
Readerielliopsis fuscoporiae	CBS 139900 = CPC 24637	Fuscoporia wahlbergii	French Guiana	C. Decock	KR476755	KR476720	MN829410	MN829326	
R. guyanensis	CBS 117550 = MUCL 46082	Dead, decaying leaf, unidentified angiosperm in leaf litter	French Guiana	Deposited by C. Decock/Isolated by M.H. de Frahan	FJ493211	MH863023	MN829411	MN829327	
Scolecoxyphium blechni	CBS 146174 = CPC 19990	Blechnum palmiforme	UK	P. Ryan	MN749224	MN749296	MN829412	MN829328	
Sc. blechnicola	CBS 146175 = CPC 19991	Blechnum palmiforme	UK	P. Ryan	MN749225	5 MN749297	MN829413	MN829329	
Sc. leucadendri	CBS 146176 = CPC 18313	Leucadendron sp.	South Africa	P.W. Crous	MN749226	6 MN749298	MN829414	MN829330	
Sc. phylicae	CBS 146177 = CPC 19219	Phyllica arborea	South Africa	M.J. Wingfield	MN749227	MN749299	MN829415	MN829331	
	CBS 146178 = CPC 19225	Phyllica arborea	South Africa	M.J. Wingfield	MN749228	8 MN749300	MN829416	MN829332	
Scorias aphidis	CBS 325.33	Aphid	-	Deposited by L.H. Leonian	MH866910) GU214696	MN829417	KT216542	
Sc. camelliae	CBS 201.30	Camellia sinensis	Indonesia	Deposited by F.H. van Beyma/Isolated by Steinmann	MH866560) MH855112	MN829418	MN829333	
Sc. leucadendri	CBS 131318 = CPC 18312	Laucadendron muirii	South Africa	P.W. Crous	JQ044456	JQ044437	MN829419	MN829334	
	CPC 17088	Callistemon sp.	Australia	P.W. Crous	MN749229	MN749301	MN829420	MN829335	
Sc. mangiferae	MFLUCC15-0230	Mangifera indica	Thailand	S. Hongsanan	KT588603	KT588604	-	-	
Sc. spongiosa	MFLUCC10-0084	Entada sp.	Thailand	P. Chomnunti	JN832601	-	-	-	
Outgroup									
Elsinoe phaseoli	CBS 165.31 = AFTOL-ID 1855 = IMI 303278	Paseolus lunatus	Cuba	Deposited by A.E. Jenkins/Isolated by C. Aguian	[.] DQ678095	5 KX887263	DQ677935	KX887144	
Myriangium hispanicum	CBS 247.33	Acer monspessulanum	-	Deposited by J.B. Martínez/Isolated by H. Diddens	GU301854	KX887304	GU349055	GU371744	

¹ ATCC: American Type Culture Collection, Virginia, USA; CBS: Westerdijk Fungal Biodiversity Institute, Utrecht, the Netherlands; CPC: Culture collection of Pedro Crous, housed at Westerdijk Fungal Biodiversity Institute; IFRDCC: International Fungal Research & Development Centre Culture Collection, Chinese Academy of Forestry, Kunning, China; IMI: International Mycological Institute, CABI-Bioscience, Egham, Bakeham Lane, United Kingdom; MFLUCC: Mae Fah Luang University Culture Collection, Chinag Ria, Thailand; MUCL: Université Catholique de Louvain, Louvain-la-Neuve, Belgium.

² Lifestyle of all sooty mould strains coded as epiphyte.

³ ITS: internal transcribed spacers and intervening 5.8S nrDNA; LSU: partial 28S large subunit RNA gene; *TEF-1a*: partial translation elongation factor 1-alpha gene; *RPB2*: partial RNA polymerase II second largest subunit gene. **Bold** GenBank accession numbers for sequences generated in this study; – indicates unavailable sequence.

(Antennulariellaceae, Capnodiaceae, Euantennariaceae, Metacapnodiaceae), hyperparasites, rock-inhabiting fungi, ectophytes, saprobes, endophytes and pathogens associated with plants and humans (Cladosporiaceae, Cystocoleaceae, Dissoconiaceae, Extremaceae, Mycosphaerellaceae, Neodevriesiaceae, Schizothyriaceae, Phaeothecaceae (including Phaeotheca fissurella and Phaeotheca shathenatiana), Phaeothecoidiellaceae, Teratosphaeriaceae (including Piedraiaceae), Comminutispora, Phaeotheca (P. salicorniae and P. triangularis) and lichenised species (Cvstocoleaceae and Racodium) (Hughes 1976, Aptroot 2006, Crous et al. 2007a, 2009, 2016, 2018, Quaedvlieg et al. 2014, Hongsanan et al. 2017, Lücking et al. 2017, Videira et al. 2017).

During the course of the past decade, considerable attention has been paid to the phylogeny and systematics of genera and families in the Capnodiales. Presently the order accommodates fungi having highly diverse ecological niches, lifestyles and modes of nutrition (Crous et al. 2007a, 2009, Ruibal et al. 2009, Schoch et al. 2009, Hyde et al. 2013, Chomnunti et al. 2014, Quaedvlieg et al. 2014, Ismail et al. 2016, Hongsanan et al. 2017, Videira et al. 2017, Crous et al. 2018). Although the Capnodiales s. str. are epiphytic sooty moulds, the presently applied circumscription also includes ectophytes and plant pathogens. Previous studies have, however, not addressed this ecological divergence adequately. This is due to a limited sampling of sooty moulds, and a poorly resolved phylogenetic backbone mainly based on nuclear ribosomal RNA genes. The aim of this study was therefore to reconsider the phylogenetic backbone of the Capnodiales by including numerous new collections of sooty moulds, thus also providing a more robust phylogeny using four nuclear loci, LSU, ITS, TEF-1α and RPB2.

MATERIALS AND METHODS

Isolates

The sooty mould isolates studied here were obtained from the culture collection (CBS) of the Westerdijk Fungal Biodiversity Institute (WI), Utrecht, the Netherlands, and the working collection of Pedro Crous (CPC) housed at the WI (Table 1). Sequences of other strains were retrieved from GenBank (Tables 1, S1). Representative cultures of the new species described in this study were deposited in the CBS culture collection.

DNA extraction, PCR amplification and sequencing

Total genomic DNA was extracted from fresh mycelia grown on malt extract agar (MEA) using the Wizard® Genomic DNA Purification Kit (Promega Corporation, Fitchburg, Wisconsin, USA) following the manufacturer's protocols. The D1/D2 variable domains of the 28S nrDNA (LSU) and the ITS1, 5.8 and ITS2 region of ribosomal DNA and part of RNA polymerase II second largest subunit (*RPB2*) and the translation elongation factor 1alpha (*TEF-1a*) were amplified and sequenced using the following primer pairs: LROR/LR5 for LSU (Vilgalys & Hester 1990), ITS5/ITS4 for ITS (White *et al.* 1990), fRPB2-5F/fRPB2-7cR for *RPB2* (Liu *et al.* 1999), EF1-983F/EF1-2218R for *TEF*-

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1 α (Rehner & Buckley 2005). The PCR amplifications were performed in a total volume of 12.5 µL containing 1 µL genomic DNA, 1 × NH4 reaction buffer (Bioline, Luckenwalde, Germany), 0.2 µM of each primer, 200 µM dNTPs, 3 mM MgCl₂, and 0.5 U *Taq* DNA polymerase (Bioline). To improve amplification of *RPB2* in some difficult DNA templates 4 % Bovine Serum Albumin (BSA, New England BioLabs, #B9000S) was added to the reaction mixture.

PCR conditions for LSU, ITS and *TEF-1a* were: an initial denaturation step of 5 min at 95 °C followed by 35 cycles of 30 s at 95 °C, 45 s at 52 °C (ITS, LSU) or 55 °C (*TEF-1a*) and 1 min at 72 °C, and a final elongation step of 7 min at 72 °C. Touchdown PCR was performed for amplification of *RPB2* as follows: an initial denaturation at 95 °C for 5 min followed by 35 cycles of 30 s at 95 °C, 30 s at 60 °C (5–10 cycles)/56 °C (5–10 cycles)/52 °C (15–25 cycles) and 1 min at 72 °C, and a final elongation step of 7 min at 72 °C.

The PCR products were sequenced with both forward and reverse primers using an Applied Biosystems 3730xl DNA Analyzer (Thermo Fisher Scientific). The DNASTAR Lasergene SeqMan Pro v. 8.1.3. software was used to obtain consensus sequences. All new sequences were submitted to GenBank (Tables 1, S1).

Phylogenetic analyses

Generated sequences were aligned with sequences retrieved from GenBank (http://www.ncbi.nlm.nih.gov) using the online interface of MAFFT v. 7 (http://mafft.cbrc.jp/alignment/server/ index.html), and manually edited in MEGA v. 7.0.21. Maximum Likelihood (ML) and Bayesian analysis (BA) were implemented for phylogenetic inferences of both single locus and concatenated alignments on the CIPRES Science Gateway portal (https://www.phylo.org/; Miller et al. 2012) using RAxML-HPC BlackBox v. 8.2.10 (Stamatakis 2014) and MrBayes v. 3.2.6 (Huelsenbeck & Ronquist 2001, Ronquist & Huelsenbeck 2003), respectively. The ML analyses were performed using a GTR+GAMMA substitution model and four rate classes with 1 000 bootstrap iterations. For the Bayesian analyses the optimal nucleotide substitution models were determined for each locus using MrModelTest v. 2.3 (Nylander 2004). Bayesian analyses were computed under the optimal nucleotide substitution models with four simultaneous Markov Chain Monte Carlo chains, 10 M generations and a sampling frequency of 1000 generations, ending the run automatically when standard deviation of split frequencies dropped below 0.01. Burn-in was set to remove 25 % of the first sampled trees, after which the 50 % majority rule consensus trees and posterior probability (PP) values were calculated. The resulting trees were plotted using FigTree v. 1.4.3 (http://tree.bio.ed.ac.uk/software/figtree). Alignments and trees were deposited in TreeBASE (www.treebase.org; S25414) and taxonomic novelties in MycoBank (www.MycoBank.org; Crous et al. 2004).

Ancestral states were reconstructed using Mesquite v.3.6 (Maddison & Maddison 2018). Character history was inferred using the Bayesian tree (see above) as phylogenetic framework. Ancestral states were determined based on a maximum likelihood approach with a MK1 model of evolution. Characters states were defined as saprobe, epiphyte, parasite or lichen. The character state for taxa with an uncertain lifestyle was coded as "?".

Morphology

Isolates stored in liquid nitrogen or lyophilised were reactivated on 2 % malt extract agar (MEA) or oatmeal agar (OA). Colonies were sub-cultured onto MEA, OA, cornmeal agar (CMA), potato dextrose agar (PDA), and synthetic nutrient-poor agar (SNA) supplemented with pine needles at room temperature. Culture media were prepared as described by Crous *et al.* (2019a, b). Cultures were examined periodically for the development of reproductive structures. Slide preparations were made with clear lactic acid or Shear's mounting fluid. Morphological observations of fungal structures were made using a Nikon SMZ1000 dissecting microscope and a Zeiss Axioscope 2 compound microscope with differential interference contrast (DIC) illumination. Measurements and images were taken using a Nikon DS-Ri2 high definition colour digital camera. Measurements and descriptions of microscopic structures were made from cultures grown on SNA. A few strains that were sterile on SNA were described from other media (indicated in text). The mean, standard deviation, maximum and minimum values of at least 30 fungal structures were calculated where possible. Dimensions are presented as a range with extremes in parentheses. Growth rates were measured on MEA after 2 wk and colony characters were noted. Colony colours were rated according to the colour chart of Rayner (1970).

RESULTS

Phylogeny

Amplification of the partial sequences of LSU, ITS and $TEF-1\alpha$ was successful but *RPB2* proved difficult to amplify using normal



Fig. 1. Reduced phylogenetic tree inferred from a Bayesian analysis based on a concatenated alignment of LSU, *TEF-1a* and *RPB2*. Bayesian posterior probabilities (PP) and maximum likelihood bootstrap support values (ML-BS) are indicated at the nodes (PP/ML-BS). The scale bar represents the expected number of changes per site. The lineages in *Capnodiales s. I.* are indicated in different colours. The tree was rooted with *Venturia inaequalis* (CBS 594.70).





Fig. 2. Phylogenetic tree inferred from a RAxML search of a concatenated alignment of LSU, ITS, *TEF-1α* and *RPB2*. Maximum likelihood bootstrap support values (ML-BS) and Bayesian posterior probabilities (PP) are indicated at the nodes (ML-BS/PP). The scale bar represents the expected number of changes per site. Families and orders are highlighted in blocks of different colour and indicated to the right of the tree. The tree was rooted with *Myriangium hispanicum* (CBS 247.33) and *Elsinoe phaseoli* (AFTOL-ID-1855). ^T Ex-type, ^{ET} Ex-epitype.



Fig. 2. Continued.

PCR, and therefore a touchdown PCR program was used. For most of the isolates the combination of fRPB2-f5F and fRPB2-7cR primers (Liu *et al.* 1999) was more successful than the primer combination fRPB2-f5F2 and fRPB2-7cR (Sung *et al.* 2007). For a few isolates, we used the forward primer fRPB2-f5F2 instead of fRPB2-f5F.

Two datasets were analysed in this study. The first dataset consisted of combined LSU, TEF-1 α and RPB2, including 193 taxa representing three orders Capnodiales s. lat., Myrangiales, Dothideales, with Venturia inaegualis (CBS 594.70) as outgroup. After alignment the dataset contained a total of 3168 characters (LSU: 837, TEF-1α: 1176, RPB2: 1147), including alignment gaps. MrModelTest revealed that the general time-reversible model of evolution (Rodríguez et al. 1990), including estimation of invariable sites and assuming a discrete gamma distribution (GTR+I+G) with six rate categories (lsetnst = 6, rates = invgamma) and dirichlet (1,1,1,1)base frequencies is the best nucleotide substitution model for all loci (LSU, TEF-1a and RPB2). The Bayesian analyses of the concatenated alignments of three loci generated 7 292 trees from which 1822 trees were discarded as burn-in. The consensus tree and posterior probability values (PP) were calculated from the remaining 5470 trees. The average standard deviation of split frequencies was 0.009987 at the end of the run. The RAxML search of the dataset with 1767 distinct alignment patterns produced a best-scoring ML tree (InL = -66287.001595). The bootstrap values equal to or higher than 50 % were mapped on the Bayesian tree (Figs 1, S1). The same phylogenetic tree was obtained from both RAxML and Bayesian analyses. Capnodiales s. lat. was split into seven distinct clades representing seven orders. Sooty mould fungi constituted Capnodiales s. str., a single highly supported clade

(ML-BS = 90 %, PP = 1). Mycosphaerellales with high support in both analyses (ML-BS = 100 %, PP = 1) proved clearly distinct from Capnodiales s. str., and was thus resurrected here as a separate order containing eight families: Mycosphaerellaceae, Dissoconiaceae, Phaeothecoidiellaceae, Schizothyriaceae, Extremaceae, Cystocoleaceae, Neodevriesiaceae and Teratosphaeriaceae (Figs 1, S1). In the Bayesian analysis Mycosphaerellales grouped with Capnodiales s. str. in a well-supported clade (PP = 0.98), while in the RAxML analysis it was a sister group of Cladosporiaceae with low support (ML-BS < 50 %). Therefore, Cladosporiaceae was elevated to ordinal level, and Cladosporiales introduced. In both RAxML and Bayesian analyses four new orders were recognised: Phaeothecales, Neophaeothecales, Racodiales (for Racodium rupestre), and Comminutisporales (for Comminutispora agavaciensis).

The second dataset consisted of aligned sequences of four loci (LSU, ITS, TEF-1a and RPB2), and included 102 taxa belonging to Capnodiales s. str., and two species, Myriangium hispanicum (CBS 247.33) and Elsinoe phaseoli (AFTOL-ID-1855), as the outgroup taxa. The aligned dataset contained 3 603 characters (LSU: 849, ITS: 553, TEF-1a: 1035, RPB2: 1154), including alignment gaps. Results from MrModelTest indicated a GTR+I+G as the best fit model for the ITS sequence data, as was the case for three other loci (LSU, TEF-1 α and RPB2) in dataset 1. The RAxML search of the second dataset detected 1 463 distinct alignment patterns and yielded a tree with InL = -30052.650187 (Fig. 2). The Bayesian analyses generated 4 622 trees from which 1154 trees were discarded as burn-in. The consensus tree and posterior probability values (PP) were calculated from the remaining 3 468 trees. The average standard deviation of split frequencies was 0.009980 at the end of the run.



Fig. 3. Cladogram showing the ancestral state reconstruction and evolution of lifestyles over the tree. Maximum likelihood bootstrap support values (ML-BS) and Bayesian posterior probabilities (PP) are indicated at the nodes (ML-BS/PP).

Posterior probability values were mapped on the ML tree (Fig. 2). Three families, namely *Capnodiaceae*, *Neoantennariellaceae* and *Readerielliopsidaceae* were recognised in *Capnodiales* s. str. In *Capnodiaceae* seven morphologically and phylogenetically well-supported genera including *Capnodium*, *Chaetocapnodium*, *Conidiocarpus*, *Heteroconium*, *Leptoxyphium*, *Phragmocapnias* and *Polychaeton* were identified. Eleven new species were recognised in this family. *Readerielliopsidaceae* contained four genera (*Phaeoxyphiella*, *Readerielliopsis*, *Scolecoxyphium* and *Scorias*) and *Neoantennariellaceae* three genera (*Fumiglobus*, *Neoantennariella* and *Neoasbolisia*).

Ancestral state reconstruction revealed a saprobic lifestyle as the ancestral state of the Capnodiales s. lat. included in this study (see Neophaeothecales; Fig. 3), while whole genome sequences of a more diverse set of taxa also supported the ancestral state of Dothideomycetes to be saprobic (Haridas et al. 2020). In the dataset included in the present study the saprobic lifestyle emerged during the evolution of diverse taxa, with several reversals back to parasitism. All Capnodiales emerged from an ancestor that had an ectophytic lifestyle. The analyses also revealed that all Mycosphaerellales shared an ancestor that was saprobic. From this ancestor ectophytes and lichen associated fungi emerged, while the ancestors of several species in this group reverted to a parasitic lifestyle. The ancestor of all Cladosporiales and Comminutisporales were saprobes but some species in the *Cladosporiales* reverted back to parasitism. Results showed that the ancestor of the Racodiales was a lichen-associated fungus, and that of the Dothideales was a saprobe.

Taxonomy

Based on the phylogenetic analyses of the two datasets generated in this study, combined with the differences in morphology and ecology, the *Capnodiales* requires redefinition, and the *Mycosphaerellales* must be resurrected. Furthermore, five new orders, four new families, three new genera, 21 new species and five new combinations are introduced below.

Capnodiales Woron. Ann. Mycol. 23: 177. 1925.

Note: Treated below as Capnodiales s. str.

Cladosporiales Abdollahz. & Crous, *ord. nov.* MycoBank MB833140.

Etymology: Name refers to the genus Cladosporium.

Saprobic, endophytic, fungicolous, lichenicolous, human and plant pathogen. Ascomata pseudothecial, gregarious or scattered, immersed, black to red-brown, globose to subglobose, uniloculate, with 1(-3) short, periphysate ostiolar necks. Ostiole necks periphysoid. Hamathecium of hyaline, septate, subcylindrical pseudoparaphyses. Asci 8-spored, bitunicate, fissitunicate, sessile to short-stalked, obovoid to broadly ellipsoid or subcylindrical, straight to slightly curved. Ascospores bi- to multiseriate, or overlapping, hyaline, obovoid to ellipsoid-fusiform, with irregular luminar inclusions. Asexual morphs hyphomycetous. Conidiophores macronematous, mononematous, simple or branched, brown. Conidiogenous cells integrated, terminal and intercalary, sympodial or synchronous, mostly polyblastic, conidiogenous loci conspicuous, darkened-refractive or not. Conidia mostly in branched or unbranched acropetal chains, subhyaline to brown, smooth to verrucose or echinulate, ramoconidia present or not, dry, conidium secession schizolytic (adapted from Bensch et al. 2012).

Type genus: Cladosporium Link (sexual morph *Davidiella* Crous & U. Braun)

Family included: Cladosporiaceae Chalm. & R.G. Archibald (based on Cladosporium).

Comminutisporales Abdollahz. & Crous, *ord. nov.* MycoBank MB833141.

Etymology: Name refers to the genus Comminutispora.

Saprobic. Ascomata pseudothecial, immersed, uniloculate, separate. Asci bitunicate, 8-spored. Pseudoparaphyses absent, hamathecial tissue abundant, ostiolar canal periphysate. Ascospores muriformly septate, forming secondary ascospores



Fig. 4. Capnodium alfenasii. A, B. Colony (2-wk-old) on MEA. C. Conidiomata arising from mycelia or immature conidiomata on SNA. D. Conidioma on SNA. E. Ostiole surround by hyaline hyphae. F. Conidia produced in ellipsoidal central part of conidioma. G. Conidia. Scale bars: C = 25 µm; D, E = 20 µm; F, G = 10 µm.



Fig. 5. Capnodium blackwelliae. A, B. Colony (2-wk-old) on MEA. C-G. Conidiomata on SNA. H. Conidia. Scale bars: C-G = 25 µm; H = 10 µm.

within the ascus. *Hyphae* hyaline, becoming olivaceous, forming hyaline, aseptate endoconidia (adapted from Ramaley 1996).

Comminutisporaceae Abdollahz. & Crous, fam. nov. Myco-Bank MB833142.

Etymology: Name refers to the genus *Comminutispora*.

Saprobic. Ascomata pseudothecial, immersed, uniloculate, separate. Asci bitunicate, 8-spored. Pseudoparaphyses absent, hamathecial tissue abundant, ostiolar canal periphysate.

Ascospores muriformly septate, forming secondary ascospores within the ascus. *Hyphae* hyaline, becoming olivaceous, forming hyaline, aseptate endoconidia.

Type genus: Comminutispora A.W. Ramaley (asexual morph *Hyphospora* A.W. Ramaley).

Mycosphaerellales (Nannf.) P.F. Cannon, Ainsworth & Bisby's Dictionary of the Fungi Ed. 9. 2001.

Saprobic, ectophytic, lichenicolous and phytopathogenic. Ascomata immersed to semi-immersed within the pseudostroma or





Fig. 6. Capnodium gamsii. A, B. Colony (2-wk-old) on MEA. C, D. Conidiomata arising from mycelia or immature conidiomata on SNA. E–H. Conidiomata on SNA. I. Conidia. Scale bars: C–F = 25 µm; G, H = 50 µm; I = 10 µm.

clypeus or superficial, solitary, globose to subglobose with protruding central ostiole, dark brown to black, scattered or clustered, gregarious. *Peridium* thin- to thick-walled, of several layers of *textura angularis*, brown to black. *Hamathecium* present or absent, with cellular pseudoparaphyses, anastomosing, branching, sometimes aparaphysate. *Asci* bitunicate, fissitunicate, 8-spored, cylindrical to cylindrical-clavate, ovoid to saccate, sessile or stipitate, apically rounded with distinct or indistinct ocular chamber. *Ascospores* bi-to multi-seriate, ellipsoidal to obclavate, oblong to cylindrical, hyaline to subhyaline or pale yellowish, mostly 1-septate, constricted or not, smooth or roughwalled. Asexual morphs hyphomycetous or coelomycetous (see Videira *et al.* 2017 for more details about asexual morphs).

Type genus: Ramularia Unger (sexual morph *Mycosphaerella* Johanson).

Families included: Cystocoleaceae (based on Cystocoleus), Dissoconiaceae (based on Dissoconium), Extremaceae (based on Extremus), Mycosphaerellaceae (based on Mycosphaerella), Neodevriesiaceae (based on Neodevriesia), Phaeothecoidiellaceae (based on Phaeothecoidiella), Schizothyriaceae (based on Schizothyrium, asexual morph Zygophiala), Teratosphaeriaceae (based on Teratosphaeria, asexual morph Kirramyces).

Neophaeothecales Abdollahz. & Crous, *ord. nov.* MycoBank MB833143.

Etymology: Name refers to the genus Neophaeotheca.

Mycelium consisting of hyaline, smooth, septate, branched hyphae, that swell in terminal or intercalary cells, developing numerous endoconidia. *Endoconidia* brown, verruculose, globose to obovoid, muriformly septate, bursting open to release endoconidia that are red-brown, verruculose, aseptate, ellipsoid to subglobose or irregular.

Neophaeothecaceae Abdollahz. & Crous, *fam. nov.* MycoBank MB833144.

Etymology: Name refers to the genus Neophaeotheca.

Mycelium consisting of hyaline, smooth, septate, branched hyphae, that swell in terminal or intercalary cells, developing numerous endoconidia. *Endoconidia* brown, verruculose, globose to obovoid, muriformly septate, bursting open to release endoconidia that are red-brown, verruculose, aseptate, ellipsoid to subglobose or irregular.

Neophaeotheca Abdollahz. & Crous, *gen. nov.* MycoBank MB833145.

Etymology: Name refers to its morphological similarity with the genus *Phaeotheca*.

Mycelium consisting of hyaline, smooth, septate, branched hyphae, that swell in terminal or intercalary cells, developing numerous endoconidia. *Endoconidia* brown, verruculose, globose to obovoid, muriformly septate, bursting open to release endoconidia that are red-brown, verruculose, aseptate, ellipsoid to subglobose or irregular.

Type species: Neophaeotheca salicorniae (Crous & Roets) Abdollahz. & Crous

Neophaeotheca salicorniae (Crous & Roets) Abdollahz. & Crous, *comb. nov.* MycoBank MB833146.

Basionym: Phaeotheca salicorniae Crous & Roets, Persoonia 36: 365. 2016.

Neophaeotheca triangularis (de Hoog & Beguin) Abdollahz. & Crous, *comb. nov.* MycoBank MB833147.

Basionym: Phaeotheca triangularis de Hoog & Beguin, Antonie van Leeuwenhoek 71: 290. 1997.

Phaeothecales Abdollahz. & Crous, *ord. nov.* MycoBank MB833148.

Etymology: Name refers to the genus Phaeotheca.

Mycelium consisting of hyaline to brown, smooth-walled, septate, branched hyphae, terminal or intercalary cells becoming swollen, developing numerous endoconidia. *Endoconidia* brown, smooth to verruculose, thin- to thick-walled, globose to obovoid, aseptate to muriformly septate (from Crous *et al.* 2018).

Type genus: Phaeotheca Sigler, Tsuneda & J.W. Carmich.

Family included: Phaeothecaceae (based on Phaeotheca, see Crous et al. 2018).

Racodiales Abdollahz. & Crous, *ord. nov.* MycoBank MB833149.

Etymology: Name refers to the genus Racodium.

Thallus filamentous, of elongated, straight hyphae, longitudinally arranged, in close association with photobiont, not corticate, dark brown to black, forming wefts or circular patches, margin not delimited; *hyphae* 4–7 per photobiont filament, straight and parallel, unbranched, non-nodulose. *Ascomata* and *conidiomata* not known (from Smith *et al.* 2009).

Type genus: Racodium Fr.

Family included: Racodiaceae (based on Racodium).

Notes: The typification of *Racodium* Fr. (based on *R. rupestre* Pers.) was discussed by Hawksworth *et al.* (2011). Based on the

sequences included here, Racodium (Racodiaceae Link) represents an undescribed order.

Capnodiales s. str.

The genera delineated in Fig. 2 are treated alphabetically based on order.

Capnodiales Woron. Ann. Mycol. 23: 177. 1925.

Widespread in tropical and subtropical areas, occurring on honeydew excretions from insects, forming a black, sooty growth on green, healthy leaves, stems and bark. *Ascomata* superficial on mycelium, subglobose to globose, with or without setae, dark brown, with a central ostiole. *Pseudoparaphyses* absent. *Asci* bitunicate, saccate, with a short pedicel, lacking an ocular chamber. *Ascospores* multiseptate or muriform, hyaline to brown. *Asexual morphs* pycnidial coelomycetous or hyphomycetous (Hughes 1976, Crous *et al.* 2009, Chomnunti *et al.* 2011).

Type genus: Capnodium Mont.

Families included: Capnodiaceae (based on Capnodium), Neoantennariellaceae (based on Neoantennariella) and Readerielliopsidaceae (based on Readerielliopsis).

Capnodiaceae Höhn. ex Theiss., Verh. Zool.-Bot. Ges. Wien 66: 363. 1916.

Growing superficially on honeydew excretions from insects, having a black, sooty-like appearance on green leaves, stems and, bark; often co-occurring with other fungicolous taxa. *Mycelium* superficial on host surface, black, sooty-like, consisting



Fig. 7. Capnodium neocoffeicola. A, B. Colony (2-wk-old) on MEA. C. Conidiomata arising from mycelia or immature conidiomata on SNA. D. Conidiomata on SNA. E. Conidia produced in ellipsoidal central part of conidioma. F. Conidia. Scale bars: C, E = 20 μm; D = 25 μm; F = 10 μm.



of septate, branched, brown hyphae. Sexual morph: Ascomata formed in mycelial mass, subglobose to globose, setae present or lacking, dark brown, with central ostiole; peridium brown, thinwalled, cells of *textura angularis*. Pseudoparaphyses absent. Asci 8-spored, bitunicate, saccate, short pedicellate, generally lacking an ocular chamber. Ascospores bi- to tri-seriate, multiseptate or muriform, hyaline to brown. Asexual morphs: coelomycetous Conidiomata synnematous or pycnidial, globose to pyriform, mostly elongated, with or without necks, and with or without swelling, and central ostiole. Conidia hyaline, aseptate, ellipsoid; hyphomycetous. Conidiophores superficial, erect, brown, cylindrical, septate, proliferating percurrently at apex. Conidia brown, septate, ellipsoid or subcylindrical, solitary or in chains.

Type genus: Capnodium Mont.

Capnodium Mont., Ann. Sci. Nat. Bot. 11: 233. 1849.

Saprobic on sugary exudates from insects growing on the surface of leaves, fruits, stems and other non-plant objects. Thallus a loose or dense network of pale brown, superficial hyphae or a thick pseudoparenchymatous stromata, with sexual and asexual morphs often growing together. Ascomata superficial on mycelium, brown to dark brown or black, globose to ellipsoidal, shortstalked or sessile, ostiolate at maturity, scattered or in groups, lacking setae. Peridium comprising dark brown to pale brown. thick-walled cells forming a textura angularis. Asci 8-spored, bitunicate, clavate, ovoid or saccate, aparaphysate, apedicellate. Ascospores brown, oblong or ovoid and some reniform, transversely septate with or without one or more vertical septa. Conidiomata pycnidial, slender to flask-shaped, simple or branched, occur singly or in groups, sessile or with long or short stalk, sometimes on the same base or stalk, with or without conspicuous oval or ellipsoidal part, with short to long or without conspicuous neck, sometimes with two necks, dark brown. *Ostiole* at apex of pycnidia, hyphae continuing upwards to the tapered neck, terminating in an ostiole which is surrounded by obtusely rounded hyphal ends. *Conidia* small, ellipsoid, continuous, hyaline, aseptate (adapted from Chomnunti et al. 2011).

Type species: Capnodium citri Berk. & Desm.

Notes: The taxonomic history of *Capnodium* was discussed by Chomnunti *et al.* (2011). Index Fungorum lists 140 species names in *Capnodium*, while MycoBank lists 168 species names (accessed March 2019). DNA sequence data are available for only two recently published species; *Ca. coartatum* (LSU) and *Ca. coffeicola* (LSU/ITS). In the present study we sequenced LSU, ITS, *TEF-1a* and *RPB2* loci for 13 isolates. Phylogenetic analyses revealed that these isolates represent six species, five of which are described as new below.

Capnodium alfenasii Abdollahz. & Crous, sp. nov. MycoBank MB833150. Fig. 4.

Etymology: Named after Prof. Acelino Couto Alfenas, in recognition to his contributions to the study of Brazilian fungal biodiversity.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, flask-shaped, mostly simple and rarely branched, occur singly or in groups, medium to dark brown, synnematous, 113–243 µm long (av. = 187 µm, n = 20), mostly sessile or with short stalk ($25-46 \times 18-35 \mu$ m, av. = $40 \times 24 \mu$ m), oval or ellipsoidal central part ($60-124 \times 31-46 \mu$ m, av. = $93 \times 34 \mu$ m), neck ($38-118 \times 8.8-13.5 \mu$ m, av. = $70 \times 11 \mu$ m), wall comprising mostly cylindrical cells. *Ostiole* at apex of pycnidia, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to



Fig. 8. Capnodium paracoffeicola. A, B. Colony (2-wk-old) on MEA. C. Conidiomata arising from mycelia or immature conidiomata on SNA. D. Conidiomata on SNA. E. Ostiole surround by hyaline hyphae. F. Conidia. Scale bars: C-E = 25 µm; F = 10 µm.

ellipsoid, continuous, $(3.7-)3.9-4.8(-5.1) \times (1.4-)$ 1.7-2(-2.2) µm (av. = 4.4 × 1.8 µm, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium, with creamy exudates of pycnidia containing conidia, folded, edge sinuate, glaucous grey to olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 19 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Brazil**, Minas Gerais, Viçosa, on leaves of *Tabebuia* sp., 1993, A.C. Alfenas (**holotype** CBS H-24256, culture ex-type CBS 146151 = CPC 22666.

Additional material examined: Brazil, Minas Gerais, Viçosa, on leaves of Tabebuia sp., 1993, A.C. Alfenas, culture CBS 146152 = CPC 22667, CBS H-24262).

Notes: Phylogenetically *Ca. alfenasii* forms a distinct clade (Figs S1, 2), but morphologically it is difficult to distinguish from *Ca. gamsii*, despite having smaller conidia and a shorter central pycnidial body. *Ca. alfenasii* differs from *Ca. blackwelliae* in having longer pycnidia, from *Ca. neocoffeicola* in having smaller pycnidia, and from *Ca. paracoffeicola* in having smaller pycnidia and conidia.

Capnodium blackwelliae Abdollahz. & Crous, sp. nov. Myco-Bank MB833151. Fig. 5.

Etymology: Named after Prof. Meredith Blackwell, who organised the annual meeting of the Mycological Society of America at Baton Rouge, Louisiana in 2007, during which time this fungus was collected.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, slender or flask-shaped, simple or branched, occur singly or in groups, medium to dark brown, synnematous, 42–116 µm long (av. = 95 µm, n = 20), mostly sessile or with short stalk (25–31 × 17–22 µm, av. = 27 × 19 µm), with or without conspicuous oval or ellipsoidal central region (24–143 × 19–55 µm, av. = 70 × 38 µm), with or without neck (22–102 × 10–37 µm, av. = 38 × 12 µm); wall comprising mostly cylindrical cells. *Ostiole* at pycnidial apex, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, continuous, (3.6–) 3.9–4.3(–4.8) × (1.4–)1.6–1.9(–2) µm (av. = 4.2 × 1.7 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, with creamy conidial exudates from pycnidia;

surface folded, edge metallic, sinuate, greenish glaucous to olivaceous black after 2 wk in the dark at 25 °C. Colonies reaching 22 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **USA**, Louisiana, Baton Rouge, on living leaves of *Myrtus communis*, 3 Aug. 2007, P.W. Crous (**holotype** CBS H-24266, culture ex-type CBS 133588).

Notes: Phylogenetically *Ca. blackwelliae* is closely related to *Ca. gamsii* (Figs S1, 2), but morphologically differs from all other species by having the smallest pycnidial lengths (av. = 95 μ m long). Pycnidia in *Ca. blackwelliae* are variable in shape, and range from long and flask-shaped to short and cylindrical.

Capnodium coartatum Chomnunti & K.D. Hyde, Fungal Diversity 51: 117. 2011.

Material examined: **Thailand**, Chiang Rai, on living leaves of *Alstonia scholaris*, 13 Sep. 2009, K.D. Hyde, culture CPC 17779.

Notes: We examined isolate CPC 17779 and generated sequences of four loci, namely LSU, ITS, *TEF-1a* and *RPB2*. This isolate grouped with two isolates of *Ca. coartatum*, namely MFLUCC10-0069 (ex-type) and MFLUCC10-0070 (Figs S1, 2). There are only LSU sequence data available for both isolates, and they differ at two nucleotide positions. Isolate CPC 17779 is 100 % identical with isolate MFLUCC10-0070 based on LSU sequence data. Morphologically, pycnidia (115–203 µm high, n = 20) and conidia (3.6–4.6 × 1.7–2.5 µm; av. = 3.9 × 2.25 µm, n = 50) of CPC 17779 are both smaller than in *Ca. coartatum* as described by Chomnunti *et al.* (2011), although measurements in the latter were made from fungal structures *in vivo*.

Capnodium gamsii Abdollahz. & Crous, *sp. nov.* MycoBank MB833152. Fig. 6.

Etymology: Named in honour of Prof. K. Walter Gams, who was an avid collector of microfungi, and collected this species in Sri Lanka.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, mostly slender or flask-shaped, simple or branched, occur singly or in groups, medium to dark brown, synnematous, 97–350 µm long (av. = 185 µm, n = 20), mostly sessile or with short stalk (23–44 × 16–29 µm, av. = 32 × 21 µm), with or without conspicuous oval or ellipsoidal central part, 62–206 × 19–46 µm, av. = 165 × 40 µm, neck present or absent, 15–87 × 9–25 µm, av. = 40 × 16 µm; wall comprising mostly cylindrical cells. *Ostiole*



Fig. 9. Chaetocapnodium indonesiacum. A, B. Colony (2-wk-old) on MEA. C. Conidiomata on SNA. D. Conidia. Scale bars: C = 25 µm; D = 10 µm.





Fig. 10. Chaetocapnodium insulare. A, B. Colony (2-wk-old) on MEA. C, D. Septate hyphae with mucilaginous outer wall layer and immature conidiomata on SNA. E. Conidia inside conidioma. G. Conidia. H. Ascoma with 3-septate brown ascospores. Scale bars: C-F, H = 20 μm; G = 10 μm.

at pycnidial apex, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, continuous, (3.6-) 4-5.5(-8.1) × (1.6-)1.9-2.4(-2.9) µm (av. = 4.9 × 2.2 µm, n = 50).

Culture characteristics: Colonies, leathery, metallic, appressed, with fluffy aerial mycelium at the centre, with creamy exudates of pycnidia containing conidia, folded, edge sinuate, greenish grey to greenish black after 2 wk in the dark at 25 °C. Colonies reaching 20–22 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Sri Lanka**, Hakgala Botanic Gardens, on leaves of unknown plant, Jan. 1973, W. Gams (**holotype** CBS H-24296, culture ex-type CBS 892.73).

Additional materials examined: **Thailand**, Chiang Rai, on *Lagerstroemia speciosa*, 1 Jan. 2009, P.W. Crous, culture CPC 17765 = CBS 146153; Chiang Rai, on *Lagerstroemia floribunda*, 2009, P.W. Crous, cultures CPC 20466 = CBS 146154 and CPC 20467 = CBS 146155); unknown substrate, 2009, P.W. Crous, culture CPC 20471 = CBS 146156, CBS H-24263).

Notes: Capnodium gamsii forms a well-supported phylogenetic clade (Figs S1, 2). Morphologically it is distinguishable from other species in having more cylindrical pycnidia with a much longer (av. = $165 \times 40 \ \mu$ m, I/w ratio > 4) central region. In other species the average length of the central region is less than $100 \ \mu$ m (I/w ratio < 3).

Capnodium neocoffeicola Abdollahz. & Crous, sp. nov. MycoBank MB833153. Fig. 7.

Etymology: Name refers to the fact that it is related to *Ca. coffeicola*.

Mycelium superficial or immersed, hyaline to brown, branched; hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Conidiomata* pycnidial, superficial or immersed, flask-shaped, simple and erect, occur singly or in groups, medium to dark brown,

synnematous, 134–268 µm long (av. = 230 µm, n = 20), sessile or with short stalk (27–46 × 28–40 µm, av. = 38 × 30 µm), oval or ellipsoidal central part (74–115 × 38–52 µm, av. = 90 × 46 µm), neck (44–136 × 9–12 µm, av. = 110 × 11 µm), wall comprising mostly cylindrical cells. *Ostiole* at apex of pycnidial neck, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, continuous, $(3.7-)4-4.7(-5.2) \times (1.6-)$ 1.8-2(-2.3) µm (av. = 4.4 × 1.9 µm, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium, with abundant creamy exudates of pycnidia containing conidia, folded, edge sinuate, glaucous grey to pale greenish grey after 2 wk in the dark at 25 °C. Colonies reaching 19–22 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Thailand**, Chiang Rai, on living leaves of *Coffea arabica*, 20 Aug. 2014, S. Hongsanan (**holotype** CBS H-24267, culture ex-type CBS 139614 = MFLUCC 14-0570).

Additional material examined: **Thailand**, Chiang Rai, on living leaves of *Coffea arabica*, 20 Aug. 2014, S. Hongsanan, culture CBS 139613 = MFLUCC 14-0569.

Notes: In the phylogenetic tree, *Ca. neocoffeicola* is clearly a distinct species (Figs S1, 2). In terms of morphology, the smaller conidia can differentiate *Ca. neocoffeicola* from *Ca. paracoffeicola*, and the longer pycnidia (av. = 230 μ m) distinguishes it from other species examined in this study. In *Ca. paracoffeicola*, *Ca. neocoffeicola* and *Ca. coffeae* (CBS 147.52) the average pycnidial length is greater than 200 μ m, while in the other species (incl. *Ca. coffeicola*, 165–178 μ m) pycnidia are less than 200 μ m long.

Capnodium paracoffeicola Abdollahz. & Crous, sp. nov. MycoBank MB833154. Fig. 8.

Etymology: Name refers to the fact that it is related to *Ca. coffeicola*.



Fig. 11. Chaetocapnodium summerellii. A, B. Colony (2-wk-old) on MEA. C. Conidiomata on SNA. D. Conidioma with setae. E. Conidia inside conidiomata. F. Conidia. Scale bars: C, D = 25 µm; E = 20 µm; F = 10 µm.

Mycelium superficial or immersed, hyaline to brown, branched; hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, flask-shaped, simple or branched, occurring singly or in groups, medium to dark brown, synnematous, 223–337 µm long (av. = 266 µm, n = 20), sessile or with short stalk (32–87 × 17–28 µm, av. = 70 × 23 µm), oval or ellipsoidal central part (63–160 × 25–40 µm, av. = 90 × 35 µm), neck (82–173 × 8.8–13.9 µm, av. = 120 × 12 µm), wall comprising mostly cylindrical cells. *Ostiole* at apex of pycnidial neck, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, continuous, $(4.9-)5-6.5(-7.7) \times (1.8-)$ 1.9-2.3(-2.6) µm (av. = 6.4 × 2.15 µm, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, glaucous grey to pale greenish grey after 2 wk in the dark at 25 °C. Colonies reaching 17 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Thailand**, Chiang Rai, on living leaves of *Coffea arabica*, 20 Aug. 2014, S. Hongsanan (**holotype** CBS H-24268, culture ex-type CBS 139616 = MFLUCC 14-0572).



Fig. 12. Chaetocapnodium tanzanicum. A, B. Colony (2-wk-old) on MEA. C, D. Conidia inside conidiomata. E, F. Conidiomata with setae on SNA. G. Conidia. Scale bars: C, D, F = 25 µm; E = 50 µm; G = 10 µm.

Additional material examined: **Thailand**, Chiang Rai, on living leaves of *Coffea arabica*, 20 Aug. 2014, S. Hongsanan, culture CBS 139615 = MFLUCC 14-0571.

Notes: Phylogenetically this species constitutes a distinct clade (Figs S1, 2) that is characterised morphologically by larger conidia (av. = $6.4 \times 2.15 \mu m$). Average conidial lengths of the other species studied here are shorter than 5 μm . *Capnodium coffeicola* was recently described from leaves of *Coffea* sp. collected in Chiang Rai, Thailand (Hongsanan *et al.* 2015b). Morphologically it differs from *Ca. paracoffeicola* in its shorter conidiomata (165–178 μm long), and shorter stalks (19–24 long × 18–23 μm diam; Hongsanan *et al.* 2015b).

Chaetocapnodium Hongsanan & K.D. Hyde, Fungal Diversity 72: 68. 2015.

Type species: Chaetocapnodium siamensis Hongsanan & K.D. Hyde

Notes: Chaetocapnodium is a hitherto monotypic genus introduced based on the morphology of its sexual morph and supported by LSU sequence data (Liu *et al.* 2015). The phylogenies generated in the present study (Figs S1, 2), however, revealed that *Antennariella placitae* and *Phragmocapnias philippinensis* are congeneric with *Chaetocapnodium*. Nine other isolates clustered in six distinct clades representing six species, five of which are recognised as taxonomic novelties. Two new combinations are proposed and five new species described in *Chaetocapnodium*.

Chaetocapnodium indonesiacum Abdollahz. & Crous, sp. nov. MycoBank MB833156. Fig. 9.

Etymology: Name refers to Indonesia where this fungus was collected.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thinwalled, 1-2 cell layers of *textura angularis*, (20-) $25-35(-48) \times (16-)23-33(-40)$ µm. *Setae* not observed. *Ostiole* absent, or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, globose to subglobose, with minute guttules, smooth, thin-walled, (2-) $2.4-2.8 \times (1.8-)2.2-2.4(-2.6)$ µm, (av. = 2.5×2.2 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, smoke grey to pale mouse grey after 2 wk in the dark at 25 °C. Colonies reaching 43 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Indonesia**, Java Island, Bogor, Buitenzorg, on leaves of *Camellia sinensis*, 1930?, F.H. van Beyma (**holotype** CBS H-24269, culture ex-type CBS 202.30).

Notes: Phylogenetically *Ch. indonesiacum* clusters with *Ch. philippinense* (Figs S1, 2). Morphologically it is not possible to compare these two species, as the latter is known only from its sexual morph. They are genetically distinct in 1, 1, 9 and 40 bp in LSU, ITS, *TEF-1* α and *RPB2* loci. *Chaetocapnodium indonesiacum* can be distinguished from all other species by having the smallest pycnidia (25–35 × 23–33 µm). With the excepton of *Ch. indonesiacum* and *Ch. placitae*, all *Chaetocapnodium* species have conidiomatal pycnidia with setae. The radial growth rate of *Ch. indonesiacum* on MEA at 25 °C was more rapid (43 mm diam/2 wk) than that observed in all other species.



Fig. 13. Chaetocapnodium thailandense. A, B. Colony (2-wk-old) on MEA. C, D. Conidiomata with setae on SNA. E. Conidia inside conidioma. F. Conidia. Scale bars: C = 25 µm; D, E = 20 µm; F = 10 µm.

Chaetocapnodium insulare Abdollahz. & Crous, *sp. nov.* MycoBank MB833157. Fig. 10.

Etymology: Name reflects the fact that it was collected from an island.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches. meristogenous in development, pseudoparenchymatous, thinwalled, 1-2 cell layers of textura angularis, (28-) 35-55 × (22-)30-48 µm. Setae present, septate or aseptate, pale to dark brown, mostly around ostiole, $(7-)10-13(-19) \mu m$ long (av. = 12 µm, n = 30). Ostiole absent, or not well-developed, mostly releasing conidia by means of irregular rupture. Conidia hyaline, aseptate, globose to subglobose, with minute guttules, smooth, thin-walled, (2.8-)3.2-3.6(-4.4)х (2.6-)2.9-3.3(-3.7) µm, (av. = 3.4×3 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, with creamy conidial exudates, edge sinuate, smoke grey to greenish grey after 2 wk in the dark at 25 °C. Colonies reaching 14 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **South Africa**, Marion Island, Prince Edward Is., on *Phylica arborea*, 2011, M.J. Wingfield (**holotype** CBS H-24297, culture ex-type CPC 19221 = CBS 146159).

Additional material examined: **South Africa**, Marion Island, Prince Edward Is., on *P. arborea*, 2011, M.J. Wingfield, culture CPC 19223 = CBS 146160; on *P. arborea*, 2011, M.J. Wingfield, culture CPC 19224 = CBS 146161.

Notes: Based on the phylogenetic analyses *Ch. insulare* is related to *Ch. placitae* (Figs S1, 2), but morphologically it is distinct from all *Chaetocapnodium* species examined in this study by producing the largest conidia with average length and width greater than 3 (av. = $3.4 \times 3 \mu$ m). Moreover, pycnidia in *Ch. insulare* are setose with septate or aseptate setae while in *Ch. placitae* setae are absent. Radial growth rate on MEA at 25 °C is slower (14 mm diam/ 2 wk) than observed for all other species.

Chaetocapnodium philippinense (Hongsanan & K.D. Hyde) Abdollahz. & Crous, *comb. nov.* MycoBank MB833158. *Basionym: Phragmocapnias philippinensis* Hongsanan & K.D. Hyde, Fungal Diversity 72: 69. 2015.

Description: Liu et al. (2015).

Typus: **Philippines**, Laguna, Mount Makiling, on leaves of palm (*Arecaceae*), Feb. 2012, K.D. Hyde HSA14/1 (**holotype** MFLU 14-0748, ex-type culture MFLUCC 12-0110 = CPC 20474).



Fig. 14. Conidiocarpus, Phragmocapnias and Polychaeton conidiomata. A, B. Conidiocarpus conidiomata on SNA. C–E. Conidiocarpus conidiomata on OA. F. Polychaeton conidiomata on SNA. G, H. Polychaeton conidiomata on OA. I, J. Phragmocapnias conidiomata on SNA. Scale bars: A, C–E, G, H = 50 µm; F, J = 25 µm; I = 20 µm.



Fig. 15. Leptoxyphium citri. A, B. Colony (2-wk-old) on MEA. C, D. Conidiomata arising from mycelia on SNA. E. Mature funnel-shaped conidioma at apex with hyaline hyphae surrounding the ostiole. F. Proliferation through the fertile head of conidioma. G. Conidia. H–J. Synasexual morph 2-celled conidia. Scale bars: C = 40 μ m; D = 20 μ m; E = 25 μ m; F = 50 μ m; G–J = 10 μ m.

Chaetocapnodium placitae (Cheewangkoon & Crous) Abdollahz. & Crous, *comb. nov.* MycoBank MB833159. *Basionym: Antennariella placitae* Cheewangkoon & Crous, Persoonia 23: 57. 2009.

Description: Cheewangkoon et al. (2009).

Typus: **Australia**, New South Wales, Cessnock S 32°50′45″, E 151°17′07″, on *Eucalyptus placita*, 14 Oct. 2006, coll. B.A. Summerell, isol. P.W. Crous (**holotype** CBS H-20277, culture extype CPC 13706 = CBS 124785).

Chaetocapnodium siamensis Hongsanan & K.D. Hyde, Fungal Diversity 72: 69. 2015.

Asexual morph. Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thin-walled, 1-2 cell layers of textura angularis, (36-)45-70(-100) × (32-)40-65(-94) µm. Setae present, septate or aseptate, pale to dark brown, mostly around ostiole, (13.9-)20-26(-30) µm long (av. = 22.7 µm, n = 30). Ostiole absent, or not well-developed, mostly releasing conidia by means of irregular rupture. Conidia hyaline, septate or aseptate, globose to subglobose, with minute guttules, smooth, thin-walled, $(2.1-)2.4-2.8(-3) \times 2-2.4(-2.6) \mu m$ $(av. = 2.6 \times 2.3 \mu m, n = 50)$. For description of sexual morph, see Liu et al. (2015).

Material examined: **Thailand**, Chiang Rai, Bandu, on leaves of unknown plant host, 2013, S.C. Karunarathna, culture CBS 139815 = MFLUCC 13-0096.

Notes: Isolate CBS 139815 clustered (Figs S1, 2) with the extype of *Ch. siamensis* (MFLUCC13-0778, on an unidentified host plant, collected in Chiang Rai). Only LSU sequence data are available for *Ch. siamensis*, which differs from CBS 139815 at two nucleotide positions. We have characterised the asexual morph of CBS 139815, which was not described in the original description of *Ch. siamensis*.

Chaetocapnodium summerellii Abdollahz. & Crous, sp. nov. MycoBank MB833155. Fig. 11.

Etymology: Named in honour of Prof. Brett A. Summerell, Director Research & Chief Botanist at the Royal Botanic Garden Sydney, Australia, who is an active advocate for plant and fungal conservation.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thinwalled, 1-2 cell layers of *textura angularis*, (33-) $40-60(-67) \times (27-)40-55(-60)$ µm. *Setae* present, septate or aseptate, pale to dark brown, mostly around ostiole, (9-) 11-15(-19) µm long (av. = 14.3 µm, n = 30). *Ostiole* absent or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, globose to sub-globose, with minute guttules, smooth, thin-walled, (2-) $2.2-2.5(-2.6) \times 1.9-2.4$ µm, (av. = 2.3×2.1 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, with creamy conidial exudates in centre of colony; surface folded, edge sinuate, pale mouse grey to olivaceous black after 2 wk in the dark at 25 °C. Colonies reaching 29 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Australia**, New South Wales, on leaves of *Eucalyptus placita* (*Myrtaceae*), Oct. 2006, B.A. Summerell (**holotype** CBS H-24257, culture ex-type CPC 13654 = CBS 146157).



Fig. 16. Phragmocapnias bette. A. Ascoma with setae on SNA. B. Asci and ascospores. C. Conidia. D. Conidioma arising from mycelium on SNA. E. Conidioma on SNA. Scale bars: A, D = 25 µm; E = 20 µm; B, C = 10 µm.

Additional material examined: Laos, host unknown, 1 Jan. 2009, P. Pheng, culture CPC 17368 = CBS 146158, CBS H-24264.

Notes: Chaetocapnodium summerellii resembles Ch. thailandense and Ch. tanzanicum in morphology, but is phylogenetically distinct, forming a separate clade (Figs S1, 2). Furthermore, its radial growth rate on MEA at 25 °C (29 mm diam/2 wk) is slower than that of the latter two species (38 mm diam/2 wk).

Chaetocapnodium tanzanicum Abdollahz. & Crous, *sp. nov.* MycoBank MB833160. Fig. 12.

Etymology: Name refers to Tanzania where this fungus was collected.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thinwalled, 1-2 cell layers of *textura angularis*, (38–) $55-70(-138) \times (30-)45-65(-112) \mu m$. *Setae* present, septate or aseptate, pale to dark brown, mostly around ostiole, (10.5–) $12-17(-23) \mu m \log (av. = 15.6 \mu m, n = 30)$. *Ostiole* absent or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, globose to sub-globose, with minute guttules, smooth, thin-walled, (2.5–) $2.7-2.9(-3.2) \times (2.3-)2.5-2.8 \mu m$, (av. = $2.8 \times 2.6 \mu m$, n = 50).

Culture characteristics: Colonies, leathery, metallic, appressed, with fluffy aerial mycelium at the centre, with creamy exudates of pycnidia containing conidia, folded, edge sinuate, smoke grey to greenish black, edge sienna to cinnamon after 2 wk in the dark at 25 °C. Colonies reaching 38 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Tanzania**, on lichen, 1974, M. Dreyfuss (**holotype** CBS H-24270, culture ex-type CBS 145.79).

Notes: Chaetocapnodium tanzanicum is phylogenetically clearly distinct from other Chaetocapnodium spp. (Figs S1, 2). Morphologically it resembles *Ch. thailandense* and *Ch. summerellii*. It is distinguishable from *Ch. summerellii* by having a faster radial growth rate on MEA at 25 °C, and from *Ch. thailandense* by producing larger conidia.

Chaetocapnodium thailandense Abdollahz. & Crous, sp. nov. MycoBank MB833161. Fig. 13.

Etymology: Name refers to Thailand where this fungus was collected.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae mostly smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, globose to pyriform, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thinwalled, 1–2 cell layers of *textura angularis*, (32–) 40–70(–90) × (32–)40–60(–80) µm. *Setae* present, aseptate, pale to dark brown, mostly around ostiole, (8.7–) 13–19(–26.6) µm long (av. = 16.1 µm, n = 30). *Ostiole* absent, or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, 0(–1)-septate, globose to subglobose, with minute guttules, smooth, thin-walled, (2–) 2.2–2.6(–2.8) × (1.9–)2–2.7 µm (av. = 2.3 × 2.1 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, glaucous grey to pale olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 38 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Thailand**, Chiang Rai, host plant unknown, 2013, S.C. Karunarathna (holotype CBS H-24271, culture ex-type CBS 139619 = MFLUCC 13-0787).

Notes: Phylogenetically, *Ch. thailandense* constitutes a distinct lineage (Figs S1, 2). Morphologically *Ch. thailandense* resembles *Ch. tanzanicum* and *Ch. summerellii*. It is distinguished

from *Ch. tanzanicum* by having smaller conidia, and from *Ch. summerellii* by its faster radial growth rate on MEA at 25 °C.

Conidiocarpus Woron., Ann. Mycol. 24 (3/4): 250. 1926.

Saprobic on sugary exudates from insects, with dark mycelium forming a soot-like coating on the upper surface of leaves. Thallus composed of black, pelliculose, reticulately branched, dense, cylindrical, radiating, septate hyphae. Ascomata not observed. Conidiomata pycnidial, supported on black, long, narrow, cylindrical stalks composed of tightly compacted, anastomosed, synnematous cylindrical hyphae, lageniform with a brown oval or ellipsoid part, which produces a long neck and conidia. The pycnidium wall is composed of two or more layers, the outer one being more or less pseudoparenchymatous although the short cells tend to be arranged linearly, indicating their origin from longitudinally fused hypha. Ostiole surrounded by hyaline, subulate, hyphal extensions. Conidia small, ellipsoid, continuous, aseptate, hyaline, smooth-walled, arranged in a droplet at the apex of pycnidial neck (adapted from Hughes 1976, Chomnunti et al. 2011).

Type species: Conidiocarpus caucasicus Woron.

Notes: Conidiocarpus with the type species *Co. caucasicus* was introduced by Woronichin in Jaczewski (1917). However, Hughes (1976) stated that Batista & Ciferri (1963a) considered *Co. penzigii*, the second oldest species introduced in 1926, as the type species. *Conidiocarpus* has been reported as the asexual morph of *Phragmocapnias* (Hughes 1976). *Phragmocapnias betle* is the type species of *Phragmocapnias*. Following the ICN code based on the priority rule and one fungus = one name principles, Bose *et al.* (2014) chose *Conidiocarpus*.

In this study based on phylogenetic analyses, the type species of *Conidiocarpus* and *Phragmocapnias* clustered in two distinct clades representing two different genera (Figs S1, 2). As discussed by Hughes (1976), pycnidia of *Co. caucasicus* are elongated, $540-650 \mu m$ long including a stalk, swollen part and a neck. In morphological studies, we found that all of the species that grouped with *C. caucasicus* produced a *Conidiocarpus* pycnidial type, typified by having a long neck (Fig. 14).

Hughes (1976) mentioned that in cases where the sexualasexual connections have been confirmed, the pycnidial morphs of Phragmocapnias were the tall Conidiocarpus conidiomata that lacked necks (Figs 14, 16). Chomnunti et al. (2011) an epitype (MFLU09-0650, designated living culture MFLUCC10-0053) for the type species Phragmocapnias betle, and re-described this species based on the sexual morph. They did not observe the asexual morph. Based on Hughes (1976), asexual conidiomata are Conidiocarpus pycnidia (150-700 µm long) that lack a neck. To observe the pycnidial morph of Phragmocapnias betle, we used different culture media and were able to introduce the asexual morph on PDA, PCA and CMA. Conidiomata were pycnidia with short stalks, ellipsoidal swellings, and lacking necks. Therefore, following the views of Hughes (1976), we chose to resurrect Phragmocapnias for species with conidiocarpus-like pycnidia lacking necks.

A search of Index Fungorum and MycoBank (March 2019) revealed 12 names in *Conidiocarpus*, of which six species, *Co. asiaticus*, *Co. betle*, *Co. caucasicus*, *Co. philippinensis*, *Co. plumeriae* and *Co. siamensis* have DNA sequence data.

In our phylogenetic analyses based on four loci (LSU, ITS, TEF-1 α and RPB2), Conidiocarpus isolates clustered in two

subclades (Figs S1, 2). The ex-type isolates of Co. asiaticus and Co. siamensis together with Co. caucasicus and the nine isolates sequenced in this study clustered in the first subclade strongly supported in the RAxML analysis, but with no support from the Bayesian analyses. These species are morphologically different but phylogenetically unresolved, which may be due to missing data. Only LSU sequence data are available for the ex-type strains of Co. caucasicus (GUMH937) and Co. siamensis (MFLUCC10-0064), and LSU/ITS sequences for Co. asiaticus (MFLUCC10-0062). Two isolates CPC 20464 and CPC 20468 for which four genes sequenced in this study and an isolate belonging to Co. siamensis (MFLUCC10-0061) with LSU and ITS sequences clustered in the second sub-clade, representing a putatively new Conidiocarpus species supported by both RAxML and Bayesian analyses (ML-BS = 100 %, PP = 0.82). However, the identity of the other Conidiocarpus isolates included in this study can only be resolved once additional gene regions have been sequenced.

Heteroconium Petr., Sydowia 3: 264. 1949.

Type species: Heteroconium citharexyli Petr.

Descriptions: Hughes (2007), Cheewangkoon et al. (2012).

Notes: A search of Index Fungorum and MycoBank (March 2019) lists 25 and 28 names in *Heteroconium*, respectively. However, only LSU sequence data are available for the type species and no sequence data are available for other species in the genus. *Heterconium kleinzeense* was recently transferred to *Blastacervulus* (Crous *et al.* 2019a, b).

Leptoxyphium Speg., Physis, Rev. Soc. Arg. Cienc. Nat. 4 (17): 294. 1918.

Type species: Leptoxyphium graminum (Pat.) Speg.

Notes: A search in Index Fungorum and MycoBank (March 2019) listed 18 names in *Leptoxyphium*. However, sequence data are available only for the types of four species; *L. cacuminum* MFLUCC10-0059 (LSU), *L. glochidion* IFRDCC 2651 (LSU/ITS), *L. kurandae* CBS 129530 (LSU, ITS/*TEF-1a/ RPB2*) and *L. madagascariense* CBS 124766 (LSU, ITS/*TEF-1a/ RPB2*). *Leptoxyphium cacuminum* and *L. glochidion* are identical based on LSU sequences, but they are morphologically different. These species clustered in the same clade together with 14 isolates considered in this study (Figs S1, 2). Some variation. Isolates CBS 451.66 and CPC 26196 clustered in a distinct clade (Figs S1, 2) representing a new species described below.

Leptoxyphium citri Abdollahz. & Crous, *sp. nov.* MycoBank MB833163. Fig. 15.

Etymology: Name refers to *Citrus*, the host genus from which it was collected.

Mycelium superficial or immersed, grey to pale brown, branched, smooth to finely verruculose, thick-walled, septate, constricted at septa, with a mucilaginous outer wall layer, forming hyphal ropes. *Conidiomata* synnematous, simple or successively proliferating through the fertile head to produce another conidiogenous apex at a higher level, single or in groups, erect, straight to slightly flexuous; bulbous base medium to dark brown, cylindrical part dark olivaceous brown, 53–153 × 6–12 µm, expanding to a funnel-shaped hyphal apex, 20–40 µm high, 15–46 µm wide.



Fig. 17. Neoantennariella phylicae. A, B. Colony (2-wk-old) on MEA. C. Septate hyphae and immature conidioma on SNA. D–F. Intercalary, lateral and terminal conidiomata. G. Conidia. Scale bars: C–D = 20 μm; E–F = 25 μm; G = 10 μm.



Fig. 18. Neoasbolisia phylicae. A, B. Colony (2-wk-old) on MEA. C-E. Conidiomata on SNA. F. Conidia. Scale bars: C = 20 µm; D-E = 25 µm; F = 10 µm.

Conidiophores subcylindrical to subulate, septate, tightly aggregated in apical part of synnema, among synnematous hyphae that diverge close to apex. Conidiogenous cells integrated, terminal, phialidic, tapering to a truncate apex. Conidia broadly ellipsoid with rounded ends, aseptate, eguttulate, hyaline, smooth, $(3.9-)4.3-4.9(-5.3) \times (1.9-)2.1-2.4(-2.6)$ µm (av. = 4.7 × 2.2 µm, n = 50), aggregating in hyaline, slimy masses at apex of synnemata. Synasexual morph conidia arthric, single or in chains, frequently around the bulbous base of the synnemata, cylindrical to ellipsoid or ovoid, 1-septate, constricted at septum, smooth, pale to medium brown, $6-12 \times 2-5$ µm. *Culture characteristics*: Colonies, appressed, with fluffy aerial mycelium, with creamy conidial exudates, smoke grey to pale olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 39 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Spain**, on fruit of *Citrus sinensis*, Jan. 1966, H.A. van der Aa (**holotype** CBS H-14520, culture ex-type CBS 451.66).

Additional material examined: Italy, on Citrus sp., 2015, V. Guarnaccia, culture CPC 26196 = CBS 146162, CBS H-24265.

Notes: Phylogenetically, *L. citri* is closely related to *L. kurandae* (Figs S1, 2). These two species differ in 1, 4 and 10 bp in LSU,



TEF-1a and *RPB2* loci, respectively. Conidia of *L. citri* are smaller $(4.3-4.9 \times 2.1-2.4 \mu m)$ than those of *L. kurandae* $(6-7 \times 2-3 \mu m)$. It is interesting that both isolates of *L. citri* are known from *Citrus* collected in Europe.

Phragmocapnias Theiss. & Syd., Ann. Mycol. 15: 480. 1918.

Saprobic on sugary exudates from insects, dark mycelium forming a soot-like coating on the upper surface of leaves. Thallus composed of black, pelliculose, reticulately branched, dense, cylindrical, radiating, septate hyphae, Ascomata scattered, subglobose to broadly ellipsoidal, barely stalked, firmly attached to the basal hyphae, dark brown, thick-walled, ostiolate, with setae. Peridium consisting of pale to dark brown cells forming a textura angularis. Asci bitunicate, 8-spored, broadly clavate, with short pedicle. Ascospores cylindrical-clavate, hyaline, 4-septate and constricted at the septum (Chomnunti et al. 2011). Conidiomata pycnidial, similar to Conidiocarpus, but with a short stalk and oval or ellipsoid part and ostiole, lacking a neck. Ostiole surrounded by hyaline, subulate hyphal extensions. Conidia small, ellipsoid, continuous, aseptate, hyaline, smoothwalled, arranged in a droplet at the apex of pycnidium (Figs 14, 16).

Type species: Phragmocapnias betle (Syd. et al.) Theiss. & Syd.

Notes: Of the six *Conidiocarpus* species for which DNA sequence data are available, two species, *Co. betle* and *Co. plumeriae*, clustered in *Phragmocapnias* (Figs S1, 2). *Phragmocapnias betle* is consequently resurrected, and a new combination is introduced for *Conidiocarpus plumeriae*.

Phragmocapnias betle (Syd. *et al.*) Theiss. & Syd., Ann. Mycol. 15: 480. 1918. Fig. 16.

Descriptions: Hughes (1976), Chomnunti et al. (2011).

Notes: Chomnunti *et al.* (2011) provided a detailed description for *P. betle* based on the sexual morph, and designated an epitype for the species. The following description is provided for the asexual morph:

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Conidiomata* pycnidial, brown, comprised of cylindrical septate cells, (67-)90-120(-135) µm high (av. = 105 µm, n = 20), stalk brown to black, (27-) 50-70(-97) µm high (av. = 50 µm, n = 20), 12-31 diam (av. = 24 µm, n = 20), the oval, swollen part which produces conidia is brown, comprised of cylindrical, septate cells, (37-) 50-60(-69) µm high (av. = 48 µm, n = 20), (30-) 40-55(-75) µm diam (av. = 53 µm, n = 20). *Ostiole* surrounded by hyaline hyphae. *Conidia* oblong to ellipsoid, aseptate, hyaline, continuous, (4-)5-6(-7.9) × (1.4-)1.8-2.2(-2.8) µm (av. = 5.5 × 2 µm, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium, somewhat folded at the middle, pale olivaceous grey to pale greenish grey after 2 wk in the dark at 25 °C. Colonies reaching 32-34 mm diam on MEA after 2 wk in the dark at 25 °C.

Materials examined: **Philippines**, on living leaf of unidentified palm, 2009, K.D. Hyde, culture CPC 20476. **Thailand**, Chiang Rai, on living leaves of unknown plant, 1 Jan. 2009, P.W. Crous, culture CPC 17762; Chiang Rai, house of K.D. Hyde, living leaves of *Malus* sp., 2009, P.W. Crous, culture CPC 21379. *Phragmocapnias plumeriae* (Hongsanan & K.D. Hyde) Abdollahz. & Crous, *comb. nov.* MycoBank MB833164. *Basionym: Conidiocarpus plumeriae* Hongsanan & K.D. Hyde, Mycosphere 6: 820. 2015.

Description: Hongsanan et al. (2015b).

Notes: This species was recently described by Hongsanan *et al.* (2015b) based on its sexual morph. Phylogenetically, it is closely related to *P. betle* (Figs S1, 2).

Polychaeton (Pers.) Lév., In: Orbigny, Dict. Univ. Hist. Nat. 8: 493. 1846.

Mycelium superficial or immersed, hyaline to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. *Conidiomata* pycnidial, supported on a black mycelial network, narrow and cylindrical stalks composed of tightly compacted, anastomosed, synnematous cylindrical hyphae, lageniform with a brown oval or ellipsoid part which produces conidia and a long neck. The pycnidium wall is composed of two or more layers, the outer one being more or less pseudoparenchymatous although the short cells tend to be arranged linearly, indicating their origin from longitudinally fused hypha. *Ostiole* surrounded by hyaline, subulate, hyphal extensions. *Conidia* small, oblong to ellipsoid, continuous, aseptate, hyaline, smooth-walled, arranged in a droplet at the apex of pycnidium.

Type species: Polychaeton quercinum (Pers.) Kuntze

Notes: Polychaeton was introduced by Persoon as a sub-genus in Fumago, and raised to generic rank by Léveillé (1847). The taxonomy of Polychaeton was discussed by Hughes (1976) and Chomnunti et al. (2011). Hughes (1976) considered Po. citri and Po. guercinum suitable as generic types, and designated Po. quercinum as lectotype species of Polychaeton. Chomnunti et al. (2011) regarded Capnodium as sexual morph of Polychaeton, and chose Capnodium following the one fungus = one name concept. A search of Index Fungorum and MycoBank (March 2019) revealed several names in Polychaeton including Po. citri. In this study, we examined isolate CBS 116435 deposited in CBS as Po. citri (Pers.) Lév. from Iran on Citrus aurantium, isolated by Walter Gams. In the phylogenetic analyses (Figs S1, 2) this isolate clustered in a distinct clade close to Conidiocarpus and Phragmocapnias. Hughes (1976) mentioned that in Polychaeton pycnidia are supported on a stalk, have an ellipsoidal pycnidial cavity, with no conspicuous swelling, and terminate in a neck with hyaline hyphal extensions. On SNA isolate CBS 116435 produced pycnidia with a swollen body, a long neck, and a short stalk. On OA pycnidia tended to have much longer necks with no conspicuous swelling, while in Conidiocarpus pycnidia have a conspicuous swelling, a long neck, and are supported on a long stalk on both OA and SNA (Fig. 14). Therefore, we designate this clade as Polychaeton, although further studies are required to resolve the taxonomy of the various species described in the genus.

Neoantennariellaceae Abdollahz. & Crous, fam. nov. Myco-Bank MB833165.

Etymology: Name refers to the genus Neoantennariella.

Mycelium superficial or immersed, pale brown to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer.



Fig. 19. Phaeoxyphiella phylicae. A, B. Colony (2-wk-old) on MEA. C, D. Conidiomata with conidia on SNA. E. Transversely euseptate brown conidia. F, G. Spermogonia. H. Microconidia. Scale bars: C, E, F = 20 µm; D = 50 µm; G = 25 µm; H = 10 µm.



Fig. 20. Scolecoxyphium blechni. A, B. Colony (2-wk-old) on MEA. C-H. Irregularly cylindrical-oblong, straight or flexuous, simple or branched conidiomata on SNA. I. Conidia. Scale bars: C = 50 μm; D, F-H = 20 μm; E = 25 μm; I = 10 μm.

Pycnidia superficial or immersed, mostly globose or cylindrical, pale to dark brown, intercalary, lateral or terminal on erect hyphal branches. *Ostiole* absent, or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, ellipsoid to ovoid, with minute guttules, smooth, thin-walled.

Type genus: Neoantennariella Abdollahz. & Crous

Fumiglobus D.R. Reynolds & G.S. Gilbert, Cryptog. Mycol. 27: 252. 2006.

Type species: F. ficinus (Bat. *et al.*) D.R. Reynolds & G.S. Gilbert *Description:* Reynolds & Gilbert (2006).

Note: According to Index Fungorum and MycoBank (March 2019), *Fumiglobus* presently contains 10 names, of which LSU and ITS sequences data are only available for *F. pieridicola*.

Neoantennariella Abdollahz. & Crous, *gen. nov.* MycoBank MB833166.



Fig. 21. Scolecoxyphium leucadendri. A, B. Colony (2-wk-old) on MEA. C. Immature conidiomata. D–F. Irregularly cylindrical-oblong, straight or flexuous, simple or branched conidiomata on SNA. G. Conidia inside conidiomata. H. Conidia. Scale bars: C, E = 20 µm; D, F = 25 µm; G, H = 10 µm.

Etymology: Name reflects its morphological similarity to the genus *Antennariella*.

Mycelium superficial or immersed, pale brown to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, mostly globose or cylindrical, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches. *Ostiole* absent, or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, ellipsoid to ovoid, with minute guttules, smooth, thin-walled.

Type species: Neoantennariella phylicae Abdollahz. & Crous

Note: Morphologically similar to *Antennariella* (see Hughes 1976, fig. 11), but different in conidiomatal and conidium morphology.

Neoantennariella phylicae Abdollahz. & Crous, *sp. nov.* MycoBank MB833167. Fig. 17.

Etymology: Name reflects the host genus *Phylica* from which it was isolated.

Mycelium superficial or immersed, pale brown to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, mostly globose or cylindrical, pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, meristogenous in development, pseudoparenchymatous, thin-walled, 1–2 cell layers of *textura angularis*, $(40-)50-65(-85) \times (30-)50-60(-68) \ \mu m$ (av. = $45 \times 42 \ \mu m$, n = 20). *Ostiole* absent, or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, ellipsoid to ovoid, with minute guttules, smooth, thin-walled, $(3.9-)4.5-5.2(-5.9) \times 2.5-3.2(-3.6) \ \mu m$ (av. = $4.8 \times 2.9 \ \mu m$, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium; surface folded, edge sinuate, smoke grey to glaucous grey after 2 wk in the dark at 25 °C. Colonies reaching 17 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **UK**, Inaccessible Island, on *Phylica arborea*, 30 Sep. 2011, P. Ryan (**holotype** CBS H-24298, culture ex-type CPC 19989 = CBS 146163).

Additional materials examined: **South Africa**, Marion Island, Prince Edward Is., on *P. arborea*, 2011, M.J. Wingfield, culture CPC 19227 = CBS 146164. **UK**, Inaccessible Island, on *P. arborea*, 30 Sep. 2011, P. Ryan, cultures CPC 19977 = CBS 146165, CPC 19981 = CBS 146166, CPC 19985 = CBS 146167, CPC 19992.

Neoasbolisia Abdollahz. & Crous, *gen. nov.* MycoBank MB833168.

Etymology: Name reflects its morphological similarity to the genus *Asbolisia* Bat. & Cif.

Mycelium superficial or immersed, pale brown to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, mostly globose or pyriform, brown to dark brown, intercalary, meristogenous in development, pseudoparenchymatous, thin-walled. *Ostiole* absent or not well-developed, mostly releasing conidia by means of irregular rupture. *Conidia* hyaline, aseptate, oblong to ellipsoid, with minute guttules, smooth, thin-walled.

Type species: Neoasbolisia phylicae Abdollahz. & Crous

Note: Morphologically similar to *Asbolisia* Bat. & Cif. (*Nom. illegit.*, Art. 53.1), but as the latter is illegitimate, a new genus is introduced. *Neoasbolisia phylicae* Abdollahz. & Crous, *sp. nov.* MycoBank MB833169. Fig. 18.

Etymology: Name reflects the host genus *Phylica* from which it was isolated.

Mycelium superficial or immersed, pale brown to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, mostly globose or pyriform, brown to dark brown, intercalary, meristogenous in development, pseudoparenchymatous, thin-walled, 1-2 cell layers of textura angularis, (63-)80-140(-180) × (56-)70-100(-148) µm (av. = 110 × 90 µm, n = 20). Ostiole absent or not well-developed, mostly releasing conidia by means of irregular rupture. Conidia hyaline, aseptate, oblong to ellipsoid, with minute guttules, smooth, (4.5-)4.6-5.2(-5.9)thin-walled, (1.6-)х $1.8-2(-2.3) \ \mu m$ (av. = $5.2 \times 2 \ \mu m$, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium, edge sinuate, glaucous grey to olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 14 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **UK**, Inaccessible Island, on *Phylica arborea*, 30 Sep. 2011, P. Ryan (**holotype** CBS H-24299, culture ex-type CPC 19982 = CBS 146168).

Readerielliopsidaceae Abdollahz. & Crous, *fam. nov.* Myco-Bank MB833170.

Etymology: Name refers to the genus Readerielliopsis.

Mycelium superficial or immersed, hyaline to pale brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial, globose to pyriform, cylindrical to flask-shaped, short to long, straight to irregular, occurring singly or in groups, medium to dark brown, if synnematous, then with a hyaline to pale brown stalk forming a long neck. *Ostiole* absent, or present, with or without hyphal hairs. *Conidia* small, hyaline, smooth, aseptate, oblong to ellipsoid or obdeltoid, or pale to medium brown, transversely euseptate, filiform, fusoid-ellipsoidal.

Type genus: Readerielliopsis Crous & Decock

Fumagospora G. Arnaud, Ann. École Nat. Agric. Montpellier, Sér. 2 10: 326. 1911.

Type species: Fumagospora capnodioides G. Arnaud

As discussed by Hughes (1976) Fumagospora with hyaline and continuous conidia that become brown and transversely septate with one or more longitudinally septate cells, is the asexual morph of some Capnodium species (e.g. Capnodium salicinum). In this study Ca. salicinum (CBS 131.34) constitutes a distinct clade in Readerielliopsidaceae separate from other Capnodium species that grouped in Capnodium sensu Chomnunti et al. (2011). In a study on capnodiaceous sooty molds in Iran, phylogenetic analysis based on ITS sequence data placed isolate GUM 1315 with Fumagospora morphology (fig. 2, Khodaparast et al. 2020) close to Ca. salicinum CBS 131.34. Therefore, we designated Fumagospora as a generic name for this clade.

Phaeoxyphiella Bat. & Cif., Quad. Lab. Crittogam., Pavia 31: 145. 1963.

Type species: Ph. morototoni Bat. & Cif.

Notes: A search in Index Fungorum and MycoBank (March 2019) revealed seven names in *Phaeoxyphiella*. As discussed by Hughes (1976), Batista & Ciferri proposed this generic name for seven species, which fall into two groups based on conidial morphology. Four species, *Ph. bahiensis*, *Ph. fischeri*, *Ph. morototoni* and *Ph. walteri*, have long, fusoid or spindle-shaped, deeply pigmented, multiseptate phragmoconidia. Two species, *Ph. californica* and *Ph. rondeletiae*, have much shorter, oblong to



Fig. 22. Scolecoxyphium phylicae. A, B. Colony (2-wk-old) on MEA. C-F. Cylindrical-oblong, straight, simple conidiomata on SNA. G. Conidia. Scale bars: C = 25 µm; D-F = 20 µm; G = 10 µm.





Fig. 23. Scorias aphidis. A, B. Colony (2-wk-old) on MEA. C, D. Flask-shape conidiomata on SNA. E. Ostioles surround by hyaline hyphae. F, G. Conidia produced in ellipsoidal part of conidiomata. H. Conidia. Scale bars: D = 50 μm; E-F = 25 μm; G-H = 10 μm.

ellipsoidal, 3-septate phragmoconidia, which are at first hyaline, slowly becoming brown. The seventh species, *Ph. callitris*, has 5-septate conidia, but is a *nomen nudum* because it lacks a Latin diagnosis. There are no cultures or sequences available for any of these species. In this study, we examined five isolates which based on phylogenetic inference clustered in two distinct clades (Figs S1, 2).

Phaeoxyphiella australiana Abdollahz. & Crous, *sp. nov.* MycoBank MB833171.

Etymology: Name reflects the country where it was collected, Australia.

Culture sterile. *Phaeoxyphiella australiana*, differs from its closest phylogenetic neighbour, *Ph. phylicae*, by unique allelles in four loci based on alignments of the separate loci deposited in TreeBASE as Study S25414: *RPB2* positions 52(G), 55(G), 70(A), 91(C), 112(A), 136(G), 151(C), 178(A), 238(G), 259(G), 268(T), 298(T), 334(C), 346(A), 349(T), 367(T), 379(C), 382(T), 386(C), 388(G), 520(C), 529(T), 532(A), 535(C), 551(C), 586(C), 595(C), 607(T), 661(A), 667(C), 736(G), 805(A), 826(C), 862(C), 881(A), 882(T), 899(C), 904(C), 907(A), 937(C), 979(T), 988(A), 994(C), 1007(C), 1009(A), 1027(T), 1042(C), 1054(C), 1075(T); *TEF-1a* positions 60(C), 282(T), 283(T), 450(C), 489(T), 578(C), 596(C), 662(T), 674(G), 755(A), 788(T), 824(T), 848(C), 1011(A); ITS positions 67(C), 81(C), 149(T), 164(C), 412(A); LSU positions 414(T), 415(C), 609(C).

Typus: **Australia**, Western Australia, Denmark, Mount Lindesay Walk Trail, on *Agonis* sp., 19 Sep. 2015, P.W. Crous (**holotype** CBS H-24258, culture ex-type CPC 29527 = CBS 146169).

Note: Phaeoxyphiella australiana differs phylogenetically from *Ph. phylicae* in 3, 5, 14 and 50 bp in the LSU, ITS, *TEF-1* α and *RPB2* sequences.

Phaeoxyphiella phylicae Abdollahz. & Crous, *sp. nov.* Myco-Bank MB833172. Fig. 19.

Etymology: Name reflects *Phylica*, the host genus from which it was collected.

Mycelium superficial or immersed, hyaline to medium brown, branched, hyphae smooth to slightly verruculose, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, glabrous, sessile, obclavate, pyriform or conoidal, brown, membraneous, thinwalled, 105-200 × 60-130 µm. Ostiole present, simple, without hyphal extensions. Conidia pale to medium brown, transversely euseptate, not constricted at septa, 11-19-celled, filiform, fusoid-ellipsoidal, straight to somewhat curved, ends rounded, often with a truncate base, smooth, (43-) $67-80(-90) \times (5.9-)6-9(-9.8) \mu m$, (av. = $75 \times 7.7 \mu m$, n = 50). Spermatogonia superficial or immersed, globose to subglobose. pale to dark brown, mostly intercalary, lateral or terminal on erect hyphal branches, 49-69 × 43-62 µm. Ostiole absent, releasing microconidia by means of irregular rupture. Microconidia hyaline, aseptate, ellipsoid to ovoid, continuous, with minute guttules, smooth. thin-walled, (3.6-)4.2-4.9(-5.1)х (2-) $2.2-2.7(-2.9) \mu m$, (av. = $4.5 \times 2.4 \mu m$, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, glaucous grey to pale olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 18 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **UK**, Inaccessible Island, on *Phylica arborea*, 30 Sep. 2011, P. Ryan (**holotype** CBS H-24300, culture ex-type CPC 19993 = CBS 146170).

Additional material examined: UK, Inaccessible Island, on *P. arborea*, 30 Sep. 2011, P. Ryan cultures CPC 19979 = CBS 146171, CBS H-24259, CPC 19984 = CBS 146172, CPC 19987 = CBS 146173, CBS H-24260.



Fig. 24. Scorias camelliae. A, B. Colony (2-wk-old) on MEA. C. Flask-shape conidiomata on SNA. D. Ostiole surround by hyaline hyphae. E. Conidia. Scale bars: C-D = 20 µm; E = 10 µm.

Notes: Phaeoxyphiella phylicae differs from all seven species described by Batista & Ciferri (1963a) by producing 11–19-celled phragmoconidia. Conidial dimensions in *Ph. phylicae* are close to those of *Ph. walteri*, but conidia are 11–19-celled in *Ph. phylicae*, and 3–15-celled in *Ph. walteri*.

Readerielliopsis Crous & Decock, Persoonia 34: 195. 2015.

Type species: Readerielliopsis fuscoporiae Crous & Decock

Description: See Crous et al. (2015).

Note: Readerielliopsis includes two species, *R. fuscoporiae* (isolated from basidiomata of *Fuscoporia wahlbergii*) and *R. guyanensis* (isolated from the decaying leaf of an angiosperm).

Scolecoxyphium Cif. & Bat., Publicações Inst. Micol. Recife 47: 5. 1956.

Type species: Scolecoxyphium fraserae Cif. & Bat.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* irregularly cylindrical, straight or flexuous, short or long, sessile, with no swollen part to indicate the location of the pycnidial cavity; wall is composed of linearly arranged, fused hyphae. *Ostiole* present, without hyaline hyphal extensions; around the ostiole the hyphae are brown and obtusely rounded. *Conidia* ellipsoidal, hyaline and continuous (adapted from Hughes 1976).

Notes: A search in Index Fungorum and MycoBank (March 2019) revealed four names in *Scolecoxyphium*. This genus was established by Ciferri and Batista based on the type species, *S. fraserae* (Ciferri *et al.* 1956). Three additional species were introduced by Batista & Ciferri (1963a). No cultures or sequence data exist for any of these species. In the phylogenies generated here, a highly supported clade was resolved resembling *Scolecoxyphium* (Figs S1, 2). The taxa studied here differed from all previously described species of *Scolecoxyphium*, and are therefore described as new.

Scolecoxyphium blechni Abdollahz. & Crous, sp. nov. Myco-Bank MB833173. Fig. 20.

Etymology: Name reflects the host genus *Blechnum* from which it was isolated.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial or immersed, irregularly cylindrical-oblong, straight or flexuous, long, simple or

branched, occurring singly or in groups, medium to dark brown, synnematous, on SNA $80-225 \times 18-40 \ \mu m$ (av. = $160 \times 24 \ \mu m$, n = 20), on OA $180-420 \times 30-57 \ \mu m$ (av. = $225 \times 38 \ \mu m$, n = 20), sessile, without swollen part and neck; wall comprising mostly of cylindrical cells. *Ostiole* at apex of pycnidia, without hyaline hyphal extensions. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, continuous, $(3.3-)3.5-4(-4.4) \times 1.3-1.7 \ \mu m$ (av. = $3.8 \times 1.5 \ \mu m$, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, glaucous grey to pale olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 11 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **UK**, Inaccessible Island, on *Blechnum palmiforme*, 30 Sep. 2011, P. Ryan (**holotype** CBS H-24301, culture ex-type CPC 19990 = CBS 146174).

Notes: Phylogenetically (Figs S1, 2) S. blechni is closely related to S. blechnicola. Morphologically S. blechni resembles S. leucadendri, but pycnidia of S. leucadendri (av. = $500 \times 30 \ \mu$ m) on OA are much longer than those of S. blechni (av. = $225 \times 38 \ \mu$ m). Moreover, these two species differ in their geographical origin (S. blechni from UK, S. leucadendri from South Africa) and substrate (S. blechni on Blechnum palmiforme, S. leucadendri on Leucadendron sp.).

Scolecoxyphium blechnicola Abdollahz. & Crous, sp. nov. MycoBank MB833174.

Etymology: Name reflects the host genus *Blechnum* from which it was isolated.

Culture sterile. *Scolecoxyphium blechnicola* differs from its closely related species, *Scolecoxyphium blechni* by unique alleles in four loci based on alignments of the separate loci deposited in TreeBASE as Study S25414: *RPB2* positions 58(T), 82(T), 133(G), 355(T), 397(T), 403(C), 472(C), 475(C), 481(A), 551(T), 577(G), 580(A), 664(G), 667(C), 985(G), 994(C); *TEF-1a* positions 447(C), 683(G), 737(T), 761(C), 803(T), 878(T), 969(T), 1023(C); ITS positions 92(T), 93(C), 180(C), 515(T).

Typus: **UK**, Inaccessible Island, on *Blechnum palmiforme*, 30 Sep. 2011, P. Ryan (**holotype** CBS H-24261, culture ex-type CPC 19991 = CBS 146175).

Note: S. blechnicola differs from S. blechni by 4, 8 and 17 nucleotides in ITS, $TEF-1\alpha$ and RPB2 loci, respectively.

Scolecoxyphium leucadendri Abdollahz. & Crous, sp. nov. MycoBank MB833175. Fig. 21.



Etymology: Name reflects the host genus *Leucadendron* from which it was isolated.

Mycelium superficial or immersed, hyaline to brown, branched, consisting of smooth, thin-walled, septate hyphae, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, irregularly cylindrical-oblong, straight or flexuous, long, simple or branched, occurring singly or in groups, medium to dark brown, synnematous; on SNA 104-178 × 20-37 µm (av. = $150 \times 25 \mu m$, n = 20), on OA $350-700 \times 18-39 \mu m$ (av. = 500 × 30 µm, n = 20), sessile, without swelling part and neck, wall comprising mostly cylindrical cells. Ostiole at apex of pycnidia, without hyaline hyphal extensions. Conidia small, hyaline. aseptate. oblona to ellipsoid, continuous. $3-4 \times 1.3-1.6 \mu m$ (av. = $3.5 \times 1.5 \mu m$, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, smoke grey to olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 12 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **South Africa**, Western Cape Province, Hermanus, Fernkloof, on leaves of *Leucadendron* sp., 2 May 2010, P.W. Crous (**holotype** CBS H-24302, culture ex-type CPC 18313 = CBS 146176).

Notes: Scolecoxyphium leucadendri formed a distinct clade in the phylogenetic trees (Figs S1, 2), but is morphologically similar to *S. blechni*. The two species can be differentiated based on the pycnidial size on OA, geography and substrate.

Scolecoxyphium phylicae Abdollahz. & Crous, sp. nov. MycoBank MB833176. Fig. 22.

Etymology: Name reflects the host genus *Phylica* from which it was isolated.

Mycelium superficial or immersed, hyaline to brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. Pycnidia superficial or immersed, cylindrical-oblong, straight, short, simple, occur singly or in dark groups, medium to brown, synnematous, 40-65 × 27-37 µm (av. = 52 × 32 µm, n = 20), sessile, without swollen part and neck, wall comprising mostly cylindrical cells. Ostiole at apex of pycnidia, surrounded with brown and obtusely rounded hyphae, without hyaline hyphal extensions. Conidia small, hyaline, aseptate, oblong to ellipsoid, continuous, (3.6–) $4-4.3(-4.7) \times 1.2-1.7 \ \mu m$ (av. = $4.2 \times 1.4 \ \mu m$, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium, folded, edge sinuate, smoke grey to olivaceous grey with a pink pigment after 2 wk in the dark at 25 °C. Colonies reaching 15 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **South Africa**, Marion Island, Prince Edward Is., on *Phylica arborea*, 2011, M.J. Wingfield (**holotype** CBS H-24303, culture ex-type CPC 19219 = CBS 146177).

Additional material examined: **South Africa**, Marion Island, Prince Edward Is., on *P. arborea*, 2011, M.J. Wingfield, culture CPC 19225 = CBS 146178.

Notes: Genetically *S. phylicae* constitutes a separate clade within *Scolecoxyphium* (Figs S1, 2). It is morphologically distinguishable based on its short cylindrical pycnidia and longer conidia.

Scorias Fr., Syst. Mycol. (Lunde) 3(2): 269. 1832.

Type species: Scorias spongiosa (Schwein.) Fr.

Notes: A search in Index Fungorum and MycoBank (March 2019) revealed 13 names in *Scorias*. Chomnunti *et al.* (2011) reexamined the type species, *Sc. spongiosa*, and designated an epitype MFLU10-0013 (ex-epitype MFLUCC10-0084) from Thailand. Cultures and molecular data are presently available for three species: *Sc. leucadendri* (LSU, ITS, *TEF-1a* and *RPB2*), *Sc. mangiferae* (LSU and ITS) and *Sc. spongiosa* (LSU). In this study we examined and sequenced isolates CBS 201.30 and CBS 325.33, which are representatives of a new species described here.

Scorias aphidis Abdollahz. & Crous, sp. nov. MycoBank MB833177. Fig. 23.

Etymology: Name reflects the fact that it was isolated from an aphid.

Mycelium superficial or immersed, hyaline to pale brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial, flask-shape, simple, occurring singly or in groups, medium to dark brown, synnematous, 200–505 µm high (av. = 395 µm, n = 20), with a hyaline to pale brown stalk (50–108 × 32–74 µm, av. = 80 × 50 µm), conspicuous oval or ellipsoidal central part (90–260 × 42–135 µm, av. = 210 × 80 µm), and a long neck (50–185 × 15–60 µm, av. = 115 × 20 µm); wall comprising mostly cylindrical cells. *Ostiole* at pycnidial apex, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, oblong to ellipsoid, (4.9–) 5.5–6.5(–7.6) × (1.8–)2–2.6 µm (av. = 6.2 × 2.2 µm, n = 50).

Culture characteristics: Colonies leathery, appressed, with fluffy aerial mycelium and creamy exudates conidia; surface folded, edge sinuate, glaucous grey to olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 18 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Country unknown**, on aphid, 1933, dep. L.H. Leonian (**holotype** CBS H-24272, culture ex-type CBS 325.33).

Notes: Phylogenetically (Figs S1, 2) *Sc. aphidis* clustered in a clade containing *Sc. mangiferae*, *Sc. spongiosa* and *Sc. camelliae*, but differs from *Sc. mangiferae* in 8 bp in LSU and 7 bp in ITS, from *Sc. mangifera* in 7 bp in LSU and from *Sc. camelliae* in 9 bp in LSU, 8 bp in ITS, 28 bp in *TEF-1a* and 94 bp in *RPB2*. Morphologically, conidia of *Sc. aphidis* (av. = $6.2 \times 2.2 \mu$ m, on SNA) are larger than those of *Sc. spongiosa* (av. = $3.9 \times 1.9 \mu$ m on PDA) and *Sc. camelliae* (av. = $5.6 \times 2.4 \mu$ m on SNA), but similar to conidia of *Sc. mangiferae* (av. = $6.7 \times 2.5 \mu$ m *in vivo*). Moreover, pycnidia of *Sc. camelliae* (av. = 100μ m) are much shorter than those of *Sc. aphidis* (av. = 395μ m).

Scorias camelliae Abdollahz. & Crous, sp. nov. MycoBank MB833178. Fig. 24.

Etymology: Name reflects the host genus *Camillia* from which it was collected.

Mycelium superficial or immersed, hyaline to pale brown, branched, hyphae smooth, thin-walled, septate, constricted at septa, with a mucilaginous outer wall layer. *Pycnidia* superficial, flask-shape, simple, occurring singly or in groups, medium to dark brown, synnematous, $59-163 \mu m$ high (av. = 100 μm ,

n = 20), sessile or with short stalk (11–30 × 12–24 µm, av. = 15×17 µm), with conspicuous oval or ellipsoidal central part (37–85 × 16–57 µm, av. = 45×35 µm), and a long neck (19–77 × 6–27 µm, av. = 45×12 µm); wall comprising mostly cylindrical cells. *Ostiole* at apex of pycnidia, surrounded by hyaline hyphae. *Conidia* small, hyaline, aseptate, ellipsoid to ovoid, (4.4–)5.3–5.7(–7.4) × (1.9–)2.2–2.5(–2.9) µm (av. = 5.6×2.4 µm, n = 50).

Culture characteristics: Colonies, leathery, appressed, with fluffy aerial mycelium; surface folded, edge sinuate, dirty white to olivaceous grey after 2 wk in the dark at 25 °C. Colonies reaching 20 mm diam on MEA after 2 wk in the dark at 25 °C.

Typus: **Indonesia**, Java island, Bogor, Buitenzorg, on *Camellia sinensis* leaves, 1930, isol. Steinmann, dep. F.H. van Beyma (**holotype** CBS H-24273, culture ex-type CBS 201.30).

Notes: In the phylogenetic analyses (Fig. 2), *Sc. camelliae* clustered with *Sc. spongiosa* and *Sc. mangiferae*, but differs from *Sc. spongiosa* in 8 bp in LSU and from *Sc. mangiferae* in 10 bp in LSU and 1 bp in ITS. Morphologically, conidia of *Sc. camelliae* (av. = $5.6 \times 2.4 \mu m$ on SNA) are larger than those of *Sc. spongiosa* (av. = $3.9 \times 1.9 \mu m$ on PDA) and smaller than conidia of *Sc. mangiferae* (av. = $6.7 \times 2.5 \mu m$ *in vivo*).

DISCUSSION

The Capnodiales was originally established for three families of sooty moulds, namely Antennulariaceae, Capnodiaceae and Coccodiniaceae (Woronichin 1925). Schoch et al. (2006) transferred Mycosphaerellaceae and Piedraiaceae to the Capnodiales and recognised the *Cladosporiaceae* (= *Davidiellaceae*), thereby expanding the concept of the order. Subsequent phylogenetic studies further expanded the concept of Capnodiales, making it the second largest order of Dothideomycetes. These included fungi with a broad spectrum of morphology, life-styles and modes of nutrition, accommodating saprobes, plant and human pathogens, mycoparasites, lichenised and rock-inhabiting fungi, including epi-, ecto- and endophytes (Crous et al. 2009, Schoch et al. 2006, 2009, Schoch & Grube 2015), Based on this broad definition, we have included a collection of isolates representing 11 families: Capnodiaceae, Cladosporiaceae, Cystocoleaceae, Dissoconiaceae, Extremaceae, Mycosphaerellaceae, Neodevriesiaceae, Phaeothecaceae, Phaeothecoidiellaceae, Raco-Schizothyriaceae, Teratosphaeriaceae diaceae. (including Piedraiaceae) and two incertae sedis genera, Comminutispora and Phaeotheca.

Previous phylogenetic studies on *Capnodiales* all suffered from a limited sampling of true sooty moulds, an aspect that we have addressed in the present study. In addition, we have included two protein-coding genes (*TEF-1a* and *RPB2*) together with rDNA sequence data (LSU and ITS) to achieve a more stable and robustly supported phylogeny for this extremely diverse group of fungi.

Sooty moulds are presently classified in seven families, with some miscellaneous genera in either *Dothideomycetes* or *Eurotiomycetes* (Chomnunti *et al.* 2014). Dothideomycetous sooty moulds belong to four families, namely *Antennulariellaceae*, *Capnodiaceae*, *Euantennariaceae* and *Metacapnodiaceae*. Although many studies have focused on the taxonomy of sooty moulds (Hughes 1951, 1966, 1972, 1976, 1981, 2003, 2007, Yamamoto, 1954, Batista & Ciferri 1963a, b, Reynolds, 1975, 1979, 1986, 1998, Faull *et al.* 2002, Reynolds & Gilbert, 2005, 2006, Crous *et al.* 2007b, 2011a, b, Ruíz *et al.* 2008, Cheewangkoon *et al.* 2009, Chomnunti *et al.* 2011, 2014, Ren *et al.* 2012, Bose *et al.* 2014, Yang *et al.* 2014, Hongsanan *et al.* 2015a, b, Liu *et al.* 2015), cultures for inclusion in the present study were available only for *Capnodiaceae*.

In an attempt to explain the high levels of diversity in the *Capnodiales*, the resulting phylogenetic tree (LSU, *TEF-1a* and *RPB2*) revealed *Capnodiales s. lat.* as polyphyletic, representing seven orders. As a result, *Capnodiales s. str.* was redefined, the *Mycosphaerellales* was resurrected, and five new orders were introduced. These include the *Cladosporiales, Comminutisporales, Neophaeothecales, Phaeothecales* and the *Racodiales*.

All sooty mould isolates in both the RAxML and Bayesian analyses, constituted a well-supported monophyletic clade thus defining Capnodiales s. str. (ML-BS = 90 %, PP = 1). The monophyly of the sooty moulds was also supported by their unique morphology, ecology and mode of nutrition. Sooty moulds are epiphytes usually with dark-coloured and mucilaginous hyphae, occurring superficially on living plants. They are often associated with insects producing honeydew, and derive nutrients from the excretion of these insects, but they can occur also without insects and absorb other nutrients (Hughes 1976, Crous et al. 2009, Chomnunti et al. 2014). In the present study three families are recognised in Capnodiales s. str., namely Capnodiaceae, Neoantennariellaceae and Readerielliopsidaceae of which the latter two are newly described. Seven genera are recognised in Capnodiaceae: Capnodium, Cheatocapnodium, Conidiocarpus, Heterconium, Leptoxyphium, Phragmocapnias and Polychaeton. Based on morphology and phylogenetic analyses Phragmocapnias has been resurrected and a further 11 new species have been introduced in the family, which includes both hyphomycetous and coelomycetous asexual morphs.

The Neoantennariellaceae is introduced to accommodate *Fumiglobus* and two new monotypic genera, Neoantennariella and Neoasbolisia. All three genera produce pycnidial conidiomata. In *Readerielliopsidaceae* four coelomycetous genera, namely *Phaeoxyphiella*, *Readerielliopsis*, *Scolecoxyphium* and *Scorias*, and eight new species have been recognised.

Hawksworth et al. (1995) introduced Mycosphaerellaceae in the Dothideales, while Kirk et al. (2001) elevated this family to the order Mycosphaerellales, and Schoch et al. (2006) and Kirk et al. (2008) again placed it as a family in the Capnodiales. Despite high support values obtained in subsequent phylogenetic studies (Crous et al. 2009, Schoch et al. 2009, Suetrong et al. 2009, Hyde et al. 2013), the Mycosphaerellales was never resurrected, and the Capnodiales was applied in a broad sense beyond the original concept presented by Woronichin (1926) to accommodate the sooty moulds. In our multigene phylogeny (LSU, TEF-1 α and RPB2) using both RAxML and Bayesian analyses, we found Mycosphaerellales to represent a fully supported clade (ML-BS = 100 %, PP = 1) accommodating eight families, namely Cystocoleaceae, Dissoconiaceae, Extremaceae, Mycosphaerellaceae, Neodevriesiaceae, Phaeothecoidiellaceae, Schizothyriaceae and Teratosphaeriaceae. We have consequently resurrected Mycosphaerellales as a separate order and provided an amended description for it. Although the Mycosphaerellales includes species that are saprobes, ectophytes, plant pathogens and lichenised fungi, this order is mainly characterised by plant pathogenic fungi that are commonly

isolated as endophytes, being ecologically distinct from the sooty moulds, which are epiphytes. Xu *et al.* (2017) and Ismail *et al.* (2016) showed that the ectophytic sooty blotch and flyspeck fungi (*Dissoconiaceae*, *Phaeothecoidiellaceae*, *Schizothyriaceae*) have evolved from ancestral phytopathogenic relatives. Lichenisation has occurred once in *Cystocoleaceae*, while plant pathogens are found in *Mycosphaerellaceae*, *Neodevriesiaceae* and *Teratosphaeriaceae* and human pathogens have evolved in *Teratosphaeriaceae*. To better elucidate the general evolutionary pattern, a greater number of samples and genome-wide comparative analyses will be conducted in future studies.

Members of Cladosporiaceae are chiefly saprobic and endophytic, with a few species that are fungicolous, lichenicolous, or plant pathogenic. In a series of phylogenetic studies, members of Cladosporiaceae were resolved as a distinct clade apart from Capnodiaceae, Mycosphaerellaceae and allied families: Dissoconiaceae, Extremaceae, Neodevriesiaceae, Phaeothecoidiellaceae, Schizothyriaceae and Teratosphaeriaceae (Crous et al. 2009, Schoch et al. 2006, 2009, Suetrong et al. 2009, Bensch et al. 2012, Hyde et al. 2013, Van Nieuwenhuijzen et al. 2016, Videira et al. 2017). In the present study. Cladosporiaceae clustered apart from Mycosphaerellales and Capnodiales s. str. and formed a distinct clade sister to Comminutisopora agavaciensis. Therefore, Cladosporiaceae has been elevated to ordinal level as Cladosporiales. Morphologically, members of Cladosporiales are guite distinct from those of Mycosphaerellales, having long, solitary, flexuous conidiophores with chains of dry, pigmented conidia. Ecologically, conidia of Cladosporiales can rehydrate, germinate and grow within hours, while members of Mycosphaerellales are generally slow to reactivate, and far less hardy to extremes in temperature and moisture conditions.

Endoconidial taxa within Capnodiales s. lat. that belong to Comminutispora and Phaeotheca have received considerable attention (Sigler et al. 1981, Ramaley 1996, de Hoog et al. 1997, 1999, Zalar et al. 1999, Crous et al. 2009, 2016, 2018). These saprobic fungi (de Hoog et al. 1999, Crous et al. 2009) were found to occupy a basal position in the phylogenetic tree (Figs S1, 1, 3), representing three new orders, namely Comminutisporales, Neophaeothecales and Phaeothecales. Furthermore, our results have resolved Phaeotheca as polyphyletic, representing two distinct clades. Phaeotheca fissurella (Phaeothecales ord. nov.) clustered apart from P. salicornia and P. triangularis in a clade together with the lichen Racodium rupestre. Racodium rupestre clusters in a separate clade apart from another lichen species, Cystocoleus ebeneus (Cystocoleaceae, Mycosphaerellales), and represents a new order, Racodiales ord. nov. Phaeotheca salicornia and P. triangularis occupied the basal position in the phylogenetic tree (Figs S1, 1, 3) as one of the earliest lineages in Capnodiales s. lat., for which we introduced a new genus Neophaeotheca (Neophaeothecaceae fam. nov. and Neophaeothecales ord. nov.). Members of Phaeothecales, Racodiales and Neophaeothecales are commonly isolated under more dry, extreme conditions.

The present study has provided a more stable backbone for the phylogeny of sooty moulds and allied taxa formerly classified in what was circumscribed as "*Capnodiales s. lat.*" Many families are not yet represented in our phylogenetic analysis, pending further collections. Although our results revealed *Capnodiales s. lat.* as polyphyletic, including seven different orders, this remains a work in progress. Furthermore, phylogenetic ancestral reconstruction analysis has revealed the saprobic lifestyle to be a primitive state in *Capnodiales s. lat.* (see *Neophaeothecales*; Fig. 3), while Haridas *et al.* (2020) also showed the ancestral state of *Dothideomycetes* to be saprobic. Several transitions have occurred to evolve lichenised, epiphytic and plant and human pathogenic lifestyles (Hongsanan *et al.* 2016, Ametrano *et al.* 2019), with the sooty mould ecology apparently having evolved more recently. A more robust sampling of the unexplored or little-known clades of *Dothideomycetes*, and genome-wide comparative analyses will provide greater clarity on the evolutionary patterns of lifestyles and modes of nutrition, that has made it possible for communities of *Dothideomycetes* to adapt to changing environmental conditions.

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APPENDIX A. SUPPLEMENTARY DATA

Supplementary data to this article can be found online at https://doi.org/10.1016/j.simyco.2020.02.004.

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