The genus Mycosphaerella and its anamorphs

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Abstract: More than 2000 names have been described in *Mycosphaerella* and *Sphaerella* (*Dothideales*). Based on various morphological features depicted by earlier workers, six sections are recognized in the genus. About 27 anamorph genera have been linked to *Mycosphaerella*, 23 of which are recognized here. Based on phylogenetic analyses of the ITS-1, 5.8S and ITS-2 DNA sequence data of 46 species of *Mycosphaerella* species (58 strains), one large cluster was distinguished containing primarily taxa with cercosporoid anamorphs, as well as another smaller cluster for species with *Dissoconium* anamorphs.

Key words: Cercospora, Mycosphaerella, Dissoconium, ITS rDNA, phylogeny.

Introduction

Mycosphaerella Johanson (Mycosphaerellaceae, Dothideales) (including species described in Sphaerella Ces. & De Not.) is one of the largest genera of ascomycetes. Many species are economically important plant pathogens, causing leaf spots and stem cankers (Park & Keane, 1984). Some saprophytic species grow as endophytes, and fruit only later on decaying parts of their hosts. The genus includes about 2000 described species, most of which have been distinguished based on the host (Corlett, 1991, 1995). The host has also played a dominant role in distinguishing anamorph species linked to Mycosphaerella (Chupp, 1954).

Kendrick & DiCosmo (1979) calculated that Mycosphaerella had been linked to more than 27 different anamorph genera. Von Arx (1983) treated several generic names that he regarded as anamorphs, or potential anamorphs, of Mycosphaerella, many based on the different morphs colonizing the same substratum. Sutton & Hennebert (1994) and Braun (1995a), who studied the different types of conidiogenesis occurring in these genera, found that many had been prematurely reduced to synonymy by von Arx (1983), or had teleomorph affinities elsewhere. Twenty-three genera were eventually accepted as anamorphs of Mycosphaerella by Sutton & Hennebert (1994). Asteromella Pass. & Thüm. is now commonly accepted as a spermatial anamorph that occurs in most species of Mycosphaerella (Crous & Wingfield, 1996); these may also have been described as anamorphs in genera such as Ascochyta Lib., Asteroma DC. and Phoma Sacc.

In contrast, genera such as Stigmina Sacc. sensu Sutton & Pascoe (1989), Polythrincium Kunze and Lecanosticta Syd., though sometimes cited as anamorphs of Mycosphaerella, have teleomorphs in other genera (Sivanesan, 1984; Barr, 1996; Crous, 1998, 1999; Crous & Corlett, 1998), while new genera such as Cercostigmina and Xenostigmina have been erected for stigmina-like species with Mycosphaerella teleomorphs (Braun, 1993; Crous, 1998; Crous & Corlett, 1998). In addition, Mycovellosiella and Thedgonia have recently been linked to Mycosphaerella (Crous et al., 1998; Kaiser & Crous, 1998), as well as Phaeophleospora (= Kirramyces J.

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Walker, B. Sutton & Pascoe) (Crous & Wingfield, 1997b; Crous et al., 1997; Crous, 1998), Colletogloeopsis (Crous & Wingfield, 1997a), Uwebraunia (Crous & Wingfield, 1996), Clypeispora (Ramaley, 1991) and Sonderhenia (Park & Keane, 1984; Swart & Walker, 1988). Some of the genera reported as possible anamorphs of Mycosphaerella by Sutton & Hennebert (1994) are based on older literature citations reporting a species of Mycosphaerella occurring in close proximity to an anamorph [e.g. Toxosporium camptospermum Maubl. and Mycosphaerella abietis (Rostr.) Lindau fide Grove (1937) (= Delphinella abietis (Rostr.) E. Müll.) fide Müller & von Arx, 1962]; these remain to be confirmed.

A significant problem in the taxonomy of Mycosphaerella is the degree of host specificity of the various species. Most anamorph species are still defined based on the host and thus assumed to be host-specific, restricted at least to a family of phanerogamic plants (Chupp, 1954; Braun, 1995a). However, the tenability of many species is called into question because some taxa, including M. punctiformis, the type species of Mycosphaerella, have been shown to be polyphagous (von Arx, 1949). The cooccurrence of morphologically different anamorphs on the same host adds valuable information to support arguments against conspecificity of morphologically similar teleomorphs. Morphologically identical species on different hosts, which also conform in ecology and anamorphs, might be regarded as conspecific. However, this would still have to be confirmed by cross inoculation trials and molecular studies.

Because of their simple ascomatal morphology, it is unclear whether Mycosphaerella forms a monophyletic group or whether the simple morphology is derived, in which case Mycosphaerella would be paraphyletic or even polyphyletic. If the group turns out to be para- or polyphyletic, then monophyletic groups could possibly be characterized by the anamorphs.

In the past, several segregates from Mycosphaerella have been proposed, based on anamorphs. Examples include: Cercosphaerella Klebahn (with Cercospora anamorphs), Ovosphaerella Laibach (with Ovularia), Ramisphaerella Laibach (with Ramularia), Septisphaerella Laibach (with Septoria), Ramularisphaerella Klebahn (with Ramularia) and Septorisphaerella Klebahn (with Septoria). These genera, described by Klebahn (1918) and Laibach (1922) respectively, have never been widely used. Moreover, most of these generic names were not validly published, and are therefore unavailable for potential future segregation of Mycosphaerella.

It also remains to be seen whether the anamorph genera as currently circumscribed are monophyletic; preliminary studies on the ultrastructure of conidiogenesis of Septoria sensu lato (Verkley, 1997; Verkley & Priest, this volume) have shown that percurrently and sympodially proliferating cells can occur within a single strain, as is the case with other, unrelated fungi. Some Mycosphaerella species produce a Septoria anamorph in the spring (under humid conditions) that develops into a Phloeospora Wallr. conidioma in the summer on the same leaves. This suggests that the supposed differences between pycnidial and acervular conidiomata may be taxonomically insignificant. Septoria heraclei (Lib.) Desm. is a typical example. Potebnia (1908) was the first author to describe and discuss this phenomenon in detail.

Recent molecular data support the hypothesis that convergent evolution could have occurred in Mycosphaerella (Crous et al., 1999). Thus, although Uwebraunia and Dissoconium are two morphologically similar anamorphs, they differ in ecology, and may not both be allied with Mycosphaerella. Based on these data (Crous et al., 1999; Stewart et al., 1999) and the preceding discussion, 23 anamorph genera are presently accepted to have Mycosphaerella teleomorphs (Table 1). Using rDNA sequence analysis of the ITS-1 and ITS-2 regions, representatives of some of these genera have been compared in the present study (Table 2). The objectives were to determine whether Mycosphaerella is para- or polyphyletic, and whether the morphological characters used to separate the various anamorph genera are phylogenetically informative, or whether some generic concepts require revision.

Materials and methods

DNA AMPLIFICATION AND SEQUENCING

Genomic DNA was isolated from fungal mycelium (Table 1) collected directly from malt extract agar plates (MEA) (Biolab, Midrand, South Africa), using a modification of the isolation protocol of Lee & Taylor (1990). DNA quantification was done by UV spectroscopy using a Beckman Du Series 7500 Spectrophotometer. Template DNA was amplified in a 25 µl PCR reaction mix containing 2 units of Taq polymerase (Boehringer Mannheim, Germany), PCR reaction buffer supplied by the manufacturer, an additional 1.25 mM MgCl2, and 500 µM each of dATP, dCTP, dGTP, and dTTP, with 60 pmol ITS-1 and ITS-4 primers (White et al., 1990). The reaction profile was as follows: initial denaturation at 96°C for 2 min, followed by 30 cycles of denaturation at 94°C for 30 s, annealing at 55°C for 30 s, extension at 75°C for 2 min, and final extension at 75°C for 7 min in a Rapidcycler (Idaho Technology Idaho, U.S.A.). A negative control using water instead of template

Table 1. Anamorph genera linked to Mycosphaerella1.

Conidia								
Anamorph genus	Conidiomata ²	Proliferation ³	Colour ⁴	Septation	Loci ⁵	Arrangement ⁶	Mycelium ⁷	Reference
Cercospora Fres.	F	S	Н	0-multi	T,D,R	S	I	Pons & Sutton (1988)
Cercosporella Sacc.	F	S	Н	multi	T,R	S	I	Deighton (1973)
Cercostigmina U. Braun	F,S	P	P	multi	I	S	I	Braun (1993)
Cladosporium Link	F,Sol	S	P	0-5	T,D,R, P	S,C	I	von Arx (1949); David (1997)
Clypeispora Ramaley	P	M	H	0	I	S	I	Ramaley (1991)
Colletogloeopsis Crous & M.J. Wingf.	A	P,S	P	0-1	I	S	I,E	Crous & Wingfield (1997a)
Dissoconium de Hoog, Oorschot & Hijwegen	F/Sol	S	P	1	I	S	I,E	Crous et al. (1999)
Fusicladiella Höhn.	F/S	M	P	1	D	S	I	von Arx (1983)
Miuraea Hara	F,Sol	S	H,P	muriform, multi	I	S	I,E	von Arx (1983)
Mycovellosiella Rangel	F,Sol	S	P	0-multi	T,D,R	S,C	I,E	Deighton (1973); Crous et al. (1998)
Passalora Fr.	F,S	S	P	0-multi	T,D,R	S	1	Braun (1995b)
Phaeoisariopsis Ferraris	F,Syn	S	P	1-multi	T,D,R	S	I	von Arx (1983)
Phaeophleospora Rangel	P	P	P	0-multi	I	S	I	Crous & Wingfield (1997b)
Phloeospora Wallr.	A	S	H	multi	I	S	I	Sivanesan (1984)
Pseudocercospora Speg.	F,S,Sol,Syn	S	P	1-multi	I	S,C	I,E	Deighton (1976)
Pseudocercosporella Deighton	F,S,Sol	S	Н	l–multi	Ι	S,C	I,E	Braun (1998)
Ramularia Unger	F,S,Sol	S	H	0-5	T,D,R	S,C	I,E	von Arx (1983); Braun (1998)
Septoria Sacc.	P	S	Н	I-multi	1	S	I	von Arx (1983)
S <i>onderhenia</i> H.J. Swart & J. Walker	P	P	P	0-5	I	S	I	Swart & Walker (1988)
Stenella Syd.	F,Sol	S	P	0-multi	T,D,R	S,C	I,E	Sivanesan (1984)
Thedgonia B. Sutton	F,Sol	S,P	H	1-multi	I	C	I,E	Braun (1995a); Kaiser & Crous (1998)
/webraunia Crous & M.J. Wingf.	F/Sol	S	P	1	I	S	I,E	Crous & Wingfield (1996)
Xenostigmina Crous	F,S,Sol	S,P	Р	muriform, multi	I	S	I,E	Crous & Corlett (1998)

Asteromella Pass. & Thüm. spermatial states have also been described in Ascochyta Lib., Asteroma DC., Phyllosticta Pers. and Phoma Sacc.
 Fasciculate (F), sporodochial (S), solitary (Sol), pycnidial (P), acervular (A) or synnematous (Syn).
 Sympodial (S), percurrent (P), monoblastic, determinate (M).
 Hyaline (H), pigmented (P).
 Thickened (T), darkened (D), refractive (R), protruding (P), inconspicuous (I).
 Internal (I), external (E).

DNA was included in each experiment. PCR products were separated on a 0.8 % (wt/v) agarose gels (Promega, Madison, Wisconsin) stained with ethidium bromide and visualized under UV illumination. PCR products were purified by using a QIAquick PCR Purification Kit (Qiagen GmbH, Germany). The purified PCR products were sequenced using an ABI Prism 377 DNA Sequencer (Perkin-Elmer, Norwalk, Connecticut), with an ABI PRISMTM Dye Terminator Cycle sequencing Ready Reaction Kit (Perkin Elmer, Warrington, UK).

PHYLOGENETIC ANALYSIS

The nucleotide sequences of the 5.8S rRNA gene and the flanking internal transcribed spacers (ITS-1 and ITS-2) were assembled using Sequence Navigator TM version 1.0.1. (Perkin Elmer, Applied Biosystems, Inc., Foster City, CA). Alignments of the sequence files were made using CLUSTAL W (Thompson et al., 1994) and improved by eye where necessary. Alignment gaps were coded as missing data. Phylogenetic analyses were performed with PAUP* (Phylogenetic Analysis Using Parsimony) version 4.0b2a (Swofford, 1999). Maximum parsimony analysis was conducted for 284 parsimony-informative characters in the alignment using heuristic search option with 1000 random sequence input orders for exact solution. A strict consensus dendrogram was constructed with the equally most parsimonious trees. Clade stability was assessed by 1000 parsimony bootstrap replications. Tree scores, including tree length, consistency index, retention index, rescaled consistency index and homoplasy index (Cl, Rl, RC and HI), were also calculated. Botryosphaeria dothidea (Gen-Bank accession no. AF195774) was used as outgroup. Two species of Ramulispora Miura were also included. The latter genus has been linked to teleomorphs in Tapesia (Pers.) Fuckel (Dyer et al., 1996). Sequence data of isolates were deposited in GenBank (AF222826-222850).

Results

MORPHOLOGY OF TELEOMORPHS

Mycosphaerella is well-defined by erumpent to superficial, small, simple, globose, ostiolate ascomata, bitunicate, fasciculate asci, the complete absence of hamathecial filaments, and 1-septate, hyaline ascospores that lack appendages, but can have a mucous sheath (Taylor & Crous, 1998).

In contrast to the vast number of *Mycosphaerella* species described, only a few taxa can be distinguished among the saprobic species. Based on a preliminary examination of 500 type specimens by A. Aptroot, many synonyms of polyphagous saprobic species were identified, e.g. 121 for *M. punctiformis*, 140 for *M. tassiana*, 65 for *M. subradians* (Fr.) J. Schröter and 43 for *M. longissima* (Fuckel) Lindau. These species were often described merely on the basis of the host. These synonymies still have to be

confirmed with cultural studies and molecular techniques.

Based on the monographic revision of Mycosphaerella to date (A. Aptroot, in prep.), the following sections (modified from Barr, 1972) are recognized:

- (i) Section Mycosphaerella, characterized by cylindrical asci and mostly uniseriate, thin-walled, often small, inequilateral ascospores that are constricted at the septum, with rounded apices. Anamorphs: Typically Ramularia with Asteromella spermatial anamorphs. Representative species: The common polyphagous M. punctiformis (Pers.: Fr.) Starb.
- (ii) Section Tassiana M.E. Barr, characterized by pyriform asci and irregularly arranged, thick-walled, nearly equilateral ascospores that are often large and constricted at the septum, relatively broad with rounded ends. Anamorph: Typically Cladosporium. Representative species: The common polyphagous species M. tassiana (De Not.) Johanson (with large ascospores) and M. longissima (Fuckel) Lindau (with small ascospores). Further research is required to determine if the teleomorphs of Cladosporium subgen. Heterosporium (David, 1997) could be accommodated here.
- (iii) Section Caterva M.E. Barr, characterized by cylindrical asci and irregularly arranged, thin-walled, inequilateral, often medium-sized ascospores that are rarely constricted at the septum, with more or less pointed ends. Asteroma and Asteromella spermatial anamorphs are typical. Representative species: The common polyphagous M. subradians (Fr.) J. Schröter.
- (iv) Section Longispora M.E. Barr, characterized by cylindrical asci with aggregated, thin-walled, long, slender, mostly equilateral ascospores that are rarely constricted at the septum, characteristically with rounded apices and pointed bases. Anamorphs: Phloeospora or Septoria sensu lato. Representative species: M. eryngii (Fr.) Oudem. (with short spores), M. latebrosa (Cooke) J. Schröt. (with longer spores) and M. populi (Auersw.) J. Schröt. (with the longest spores in the genus). Sphaerulina Sacc., which differs only by additional ascospore septa, may be a synonym.
- (v) Section Fusispora M.E. Barr, characterized by pyriform asci and irregularly arranged, thin-walled, mostly equilateral, fusiform ascospores that are rarely constricted at the septum, pointed at both ends. Anamorphs have not been proven. Representative species: The common M. lineolata (Roberge) J. Schröt. on members of the Poaceae.
- (vi) Section Plaga M.E. Barr (including Section Macula M.E. Barr) including endophytic species sporulating on leaf spots, many of which are descri-

bed as plant pathogens. This section is characterized by obovoid to ellipsoidal or cylindrical asci, small to medium sized, fusiform to obovoid ascopores, with rounded ends. The majority of the described species originate from warm-temperate and tropical areas. Anamorphs: Colletogloeopsis, Mycovellosiella, Phaeophleospora, Pseudocercospora, Pseudocercosporella, Sonderhenia, Stenella Syd., Uwebraunia and possibly others. Representative species: M. suttoniae Crous & M.J. Wingf. and others listed by Crous (1998) on Eucalyptus species.

ANAMORPH RELATIONSHIPS

Of the 27 anamorph genera linked to Mycosphaerella by Kendrick & DiCosmo (1979), 23 are recognized in the present study (Table 1). These anamorph genera have been separated into more 'natural' or recognizable units based on features such as mycelium (presence or absence of superficial mycelium, and texture thereof), conidiophores (arrangement, branching, pigmentation), conidiogenous cells (placement, proliferation, scar type) and conidia (formation, shape, septation, wall and pigmentation).

MOLECULAR PHYLOGENY

Phylogenetic analysis of the 5.8S, ITS-1 and ITS-2 rDNA of 46 teleomorph and/or anamorph species in the Mycosphaerella complex, as well as two species of Ramulispora (Tapesia teleomorphs) revealed three clades. The major Mycosphaerella clade included most of the anamorphs studied, except Dissoconium (clade 2). However, this clade was only weakly supported by bootstrap values (53%), and should still be considered as part of the major Mycosphaerella clade. As expected, species of Ramulispora (Tapesia clade 3) grouped outside Mycosphaerella. Contrary to expectations, Mycosphaerella proved to be monophyletic. In most cases, isolates representing the same anamorph genus clustered together. In several instances, however, representatives of the same anamorph genus were found in more than one clade, suggesting that some forms may have evolved more than once or be plesiomorphic in the complex (Fig. 1).

Discussion

Species of Mycosphaerella are either saprobes or plant pathogens, usually causing leaf spots. In addition, some species have been isolated as endophytes (Crous, 1998), and such taxa might sporulate either before or after shedding of leaves. Crous (1998) reported that as many as four species could occur in the same lesion on leaves of Eucalyptus species, further suggesting that some may be primary and

others secondary pathogens. There may be a gradient of parasitism or succession of Mycosphaerella spp. on some hosts, i.e. lesions caused by primary pathogens could later be colonized by secondary pathogens, and possibly even later by saprobic taxa or endophytes. However, no species of Mycosphaerella has yet been recovered in detailed studies of foliicolous fungi occurring as endophytes in hosts in the Proteaceae known to have Mycosphaerella pathogens (Swart et al., 2000; J. E. Taylor et al., unpublished). It can be assumed that many plant pathogenic species have a latent phase, and that during this phase they could be isolated together with true endophytes. More research needs to be focused on the different sections of Mycosphaerella, however, to further clarify these issues, and determine if the species included differ in general

Field observations suggest that a major division of Mycosphaerella into two groups with different life styles (parasitic vs. saprophytic) would be artificial. There appears to be a strong correlation between the life styles of Mycosphaerella spp. and different climatic zones. In arctic and boreal regions, species appear to be saprobic, fruiting on overwintered leaves and stems (with species of the section Tassiana dominant). In the tropics, species of Mycosphaerella tend to be predominantly plant pathogens, sporulating on leaf spots on leaves that are often still attached to the host (with species of the section Plaga dominant). In temperate and subtropical regions, both groups occur, often sporulating on leaf spots or whole leaves that may or may not be dead (with species of the sections Caterva and Mycosphaerella dominant).

ANAMORPH RELATIONSHIPS

In Mycosphaerella, the morphology of the teleomorphs is conserved, and the morphologically informative morph tends to be the anamorph. A single teleomorph morphology can be linked to more than one anamorph morphology. This may reflect our inability to distinguish the teleomorphs of two biological species with distinct anamorphs, or a failure to recognize a species of Mycosphaerella with synanamorphs. Although the latter has rarely been considered in the literature on Mycosphaerella, it has been noted in cultures of Mycosphaerella suttoniae Crous & M.J. Wingf. (Crous & Wingfield, 1997b). Conflicting reports of different anamorphs for a single teleomorph may also be based on uncorroborated links observed in nature. Given the fact that several species regularly occur on the same lesion, any links based on association only must be treated with suspicion. Taxonomists working with Mycosphaerella have not typically included cultural experiments, a situation that

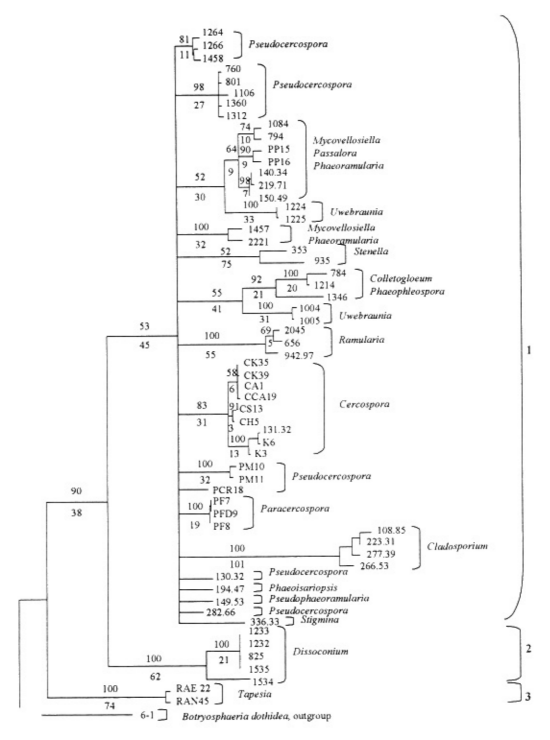


Fig. 1. The majority rule consensus tree of 80 equally most parsimonious trees derived from the alignment of the 5.8S rDNA and ITS-1 and ITS-2 regions of 58 Mycosphaerella strains using parsimony analysis with heuristic search and 1000 random sequence input orders. The robustness of the tree was assessed with 1000 bootstrap replicates. The tree was rooted with the outgroup Botryosphaeria dothidea (GenBank Accession No. AF195774). Tree length = 1367, CI = 0.407, RI = 0.680, RC = 0.277, HI = 0.593. The bootstrap values and lengths are indicated above and below the tree branches, respectively.

should be remedied in order to prove the links between species of *Mycosphaerella* and their anamorphs.

Considerable attention has been given to conidiomatal structure and mode of conidiogenesis (von Arx, 1983; Sutton & Hennebert, 1994) in this group of fungi, and significant overlap has been observed between pycnidia, acervuli and subepidermal sporodochia (Braun, 1995a). Nag Raj (1993) discussed the lack of distinguishing characters between pycnidia and acervuli, while Crous (1998) commented on sporodochia that appeared acervular when immature, which were noted by Braun (1995a) as subepidermal sporodochia. Sutton et al. (1996) examining immature material of Cylindrosporium ribis J.J. Davis, concluded that the conidiomata were acervular, and established the name Phloeosporella ribis (J.J. Davis) B. Sutton, Crous & Shamoun. Braun (1998), however, interpreted the conidiomata as subepidermal sporodochia, and established the name Pseudocercosporella ribis (J.J. Davis) U. Braun. This single criterion, and how it is interpreted, is all that distinguishes these two genera. It is clear that distinctions between some of the coelomycete and hyphomycete anamorph genera associated with Mycosphaerella are debatable.

Deighton recognized genera of cercosporoid fungi based on conidiogenesis and conidium release, as it is inferred from the dehiscence scar (Deighton, 1973, 1976, 1987, 1990). David (1993) used SEM to show differences in conidial scar structures between Mvcosphaerella anamorphs such as Cladosporium, Cercospora and Stenella. The stability of this feature in some genera, however, is questionable. Genera that have been distinguished by different modes of proliferation of the conidiogenous locus sometimes include species that are variable for this character, as reported in Septoria/Stagonospora (Sacc.) Sacc. by Verkley (1997; see also Verkley & Priest, this volume). Crous (1998) also reported that marginal thickening along the conidial rim, the character that separates Paracercospora from Pseudocercospora, disappears in culture, while conidiogenous cells on older herbarium specimens of Pseudocercospora species sometimes again develop such marginal thickening. For the majority of genera, however, scar features appear to be a usable, stable feature (Braun, 1995a).

MOLECULAR PHYLOGENY

Based on the ITS-1 and ITS-2 DNA phylogeny obtained here for the various Mycosphaerella spp. and their anamorphs (Fig. 1), three clusters were identified. These include a major monophyletic Mycosphaerella cluster (with subclusters for cercosporoid and coelomycete anamorphs), and a second cluster with Dissoconium anamorphs. A third cluster represents Ramulispora spp. with Mollisia (Fr.) P. Karst. (= Tapesia) teleomorphs. The Ramulispora spp. were included in this study because they are still often incorrectly treated as anamorphs of Mycosphaerella (under the name Pseudocercosporella). Although not related to Mycosphaerella, their taxonomy must still be resolved, and for the time being they will be referred to as the Tapesia cluster.

Based on the ITS data, it is still too early to say whether *Mycosphaerella* is mono- or polyphyletic. Although the *Dissoconium* anamorphs cluster outside the main monophyletic *Mycosphaerella* cluster, this smaller cluster is not strongly supported by bootstrap values. This situation will be reassessed in future studies incorporating more isolates (those included here being primarily cercosporoid), and also by analyzing other areas of the genome. Of note, however, are the subclusters coinciding with the different anamorph genera, which will be discussed below.

Clade 1. Mycosphaerella species with cercosporoid and coelomycete anamorphs

Pseudocercospora, Paracercospora and aggregates

Pseudocercospora is morphologically variable, and accommodates all cercosporoid anamorphs with pigmented conidiophores and conidia with inconspicuous conidial scars. Braun (1995a) showed that all the other characters are variable. These include the presence or absence of secondary mycelium, conidiophores that can be solitary to fasciculate, arranged in loose or rich fascicles or dense sporodochial conidiomata, which can also be superficial or subepidermal. Conidia are primarily formed singly, rarely in chains; they are scolecosporous, multiseptate, subhyaline to brown, mostly have transverse eusepta, and rarely have longitudinal or oblique septa. Conidiogenous cells are polyblastic, proliferating sympodially, or rarely percurrently by inconspicuous, regular annellations. Scars are mostly inconspicious and flat, or can occur on small outgrowths that appear as reduced denticles.

Several genera are morphologically similar to Pseudocercospora. Semipseudocercospora J.-M. Yen includes species with conidiogenous cells that are conspicuously denticulate, non-geniculate, and give rise to transversely euseptate conidia with basal cells that constrict to narrow hila. Similarily, species of Denticularia Deighton also have denticulate conidiogenous cells that give rise to 0-1(-3)-septate, fusiform-ellipsoidal, catenate conidia. Pantospora Cif. species have dense, synnematous fascicles, and produce dictyospores or scolecospores.

Deighton (1976) originally distinguished Cercoseptoria Deighton from Pseudocercospora based on the narrowly acicular conidia that taper only slightly at the truncate base in species of Cercoseptoria. This separation was later found to be impractical, and Cercoseptoria was accepted as a synonym of Pseudocercospora (Deighton, 1987). Paracercospora Deighton (1979) was distinguished from Pseudocercospora based on the narrow thickening along the rim of the scars on the conidiogenous cells and the conidial hila. Crous (1998), in a study of Pseudocercospora basiramifera in culture, found that this thickening was not a stable feature. Furthermore, based on ITS-1 & ITS-2 rDNA sequence analysis, Stewart et al. (1999) were

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unable to distinguish species of *Pseudocercospora* from those of *Paracercospora*, and therefore reduced *Paracercospora* to synonymy under *Pseudocercospora*. Following the concepts of Deighton (1976, 1979), *Pseudocercospora natalensis* would have been placed in *Cercoseptoria*, and *P. basiramifera* and *Paracercospora fijiensis* in *Paracercospora*.

Pseudophaeoramularia U. Braun represents species intermediate between Pseudocercospora and Phaeoramularia Muntañola, with unthickened, but slightly darkened, refractive hila, and catenate conidia. Although separate in the final analysis, in some trees Pseudophaeoramularia angolensis clustered with Pseudocercospora, suggesting that the concept of Pseudocercospora could be further expanded to incorporate taxa with catenate conidia, and unthickened, but slightly darkened, refractive hila. Cercospora angolensis was placed in Phaeoramularia by Kirk (1986), who observed conidia to be catenate. Braun (1999b) found conidial hila to be unthickened, and he therefore placed the species in Pseudophaeoramularia. Analysis of additional species of the former genus would help clarify this subgroup. For the present, however, it seems best to retain Pseudophaeoramularia separate from Pseudocercospora.

Many different groups of cercosporoid fungi are now accommodated in Pseudocercospora. In 1991 R.K. Verma (pers. comm.), who studied over 300 different species of Pseudocercospora, suggested that seven groups could be recognized in the genus. Three groups were characterized by having well-developed stromata and branched conidiophores, that could be (i) arranged in dense fascicles, (ii) be longer and in loose fascicles, or (iii) in dense, synnematous conidiomata, with sympodially, or rarely percurrently proliferating conidiogenous cells. A further two groups were distinguished based on their poorly developed or absent stromata, and (iv) denticulate conidiogenous cells, or (v) percurrent proliferation of the conidiogenous cells. The final two groups were distinguished by having no or poorly developed stromata, with conidia that could be catenate, with slightly thickened hila, and (vi) not forming rope-like structures on leaf surfaces, or (vii) forming them in abundance. Based on current taxonomic concepts, groups 6 and 7 can be accommodated in Mycovellosiella, because of the smooth superficial hyphae and thickened conidial loci. Group 5 was described as Cercostigmina U. Braun (1993), although some species have well-developed stromata, and rough conidia. Braun et al. (1999) recently introduced Scolecostigmina U. Braun to accommodate species with sporodochial conidiomata, firm stromata, verruculose conidiophores, coarse annellations, and multi-septate scolecospores that are smooth to rough, and thick-walled. Group 4 could possibly be accommodated in *Denticularia* or *Semi-pseudocercospora*, as discussed above, which essentially leaves three groups that are distinguished based on whether their conidiomata consist of short, densely arranged conidiophores, longer, loosely arranged conidiophores, or loosely synnematous conidiophores. The type species of *Pseudocercospora*, *P. vitis* (Lév.) Speg., frequently forms loosely synnematous conidiomata, and it is therefore debatable whether conidiophore arrangement can provide useful characteristics for further subdivision of the genus.

Braun (1998) proposed a formal division of Pseudocercospora into five sections, viz. sect. Pseudocercospora (conidiophores very long, in dense, loosely synnematous conidiomata), section Cercocladospora (G.P. Agarwal & S.M. Singh) U. Braun (conidiophores solitary or in loose fascicles, non-sporodochial, conidia scolecosporous, pluriseptate, superficial mycelium formed or lacking), section Cercoseptoria (Petr.) U. Braun (conidiophores in dense sporodochial conidiomata, conidia scolecosporous, pluriseptate, superficial mycelium usually absent, rarely present), section Helicomina (L.S. Olive) U. Braun (conidia strongly curved to helicoid), and section Brachypseudocercospora U. Braun (conidia solitary, rarely in chains, non-scolecosporous, (0-)1-3(-4)-septate), but he emphasized that these 'units' should only be regarded as morphological groups, which are formally treated as sections, independent of their taxonomic value.

Based on ITS-1 and ITS-2 sequence data obtained for species of Pseudocercospora in the present study (STE-U 760, 801, 1106, 1264, 1266, 1312, 1360, 1458, PCR 18, PM 10, 11, CBS 130.32, 282.66), several subclades could be distinguished within the cercosporoid clade (Fig. 1). Isolates from Eucalyptus species occurred in two subclades, while those from banana clustered separately, suggesting that coevolution with the host may play a role, or species on the same host are simply more closely related. In earlier analyses (Crous et al., 1999; Stewart et al., 1999), species of Paracercospora grouped with Pseudocercospora, although in our final analysis in this study, they again grouped separately. However, Ps. basiramifera (STE-U 1266), which has a marginal thickening along its conidial rim, clustered with species of Pseudocercospora.

In some trees, *Phaeoisariopsis griseola* (synnematous conidiophores with slightly thickened conidial loci) (CBS 194.47), and *Stigmina platani* (muriformly distoseptate conidia, and conidiogenous cells with irregular percurrent proliferations) (CBS 336.33) also clustered with species of *Pseudocercospora*, although the final analysis showed them to be distinct, but closely related. Braun (1993) and Crous (1998) speculated that species of *Stigmina* would not have *Mycosphaerella* teleomorphs, and that those species with reported *Mycosphaerella* teleomorphs would be better accommodated in *Cercostigmina* or *Xenostigmina*. The fact that *S. platani* groups in the *Mycosphaerella* cluster refutes these arguments. Additional genera in this complex that could possibly also have teleomorphs in *Mycosphaerella* include *Scolecostigmina* (Braun *et al.*, 1999) and *Dictyorostrella* U. Braun (Braun, 1999a).

Another Pseudocercospora subgroup that contains species from Eucalyptus leaves includes P. crystallina, P. colombiensis, P. heimii, P. irregulariramosa, and P. heimioides (STE-U 801, 760, 1360, 1312). All species in this cluster (except P. colombiensis) form red crystals in culture. Although the ascospore germination patterns differ among these species (Crous, 1998), the DNA data obtained here suggest that these taxa are closely related, and with the exception of P. colombiensis, may represent different varieties of P. heimii. Isolates representing the P. heimii-complex have also recently been obtained from eucalypts collected in Hawaii and Brazil, and this species may thus have a very wide geographical distribution. Further collections and pathogenicity trials are required to clarify this issue. Species in this clade have light brown, smooth to finely verruculose, obclavate to subcylindrical conidia, with internal and external mycelium, small, dense fascicles situated on brown stromata, and conidiophores that are frequently reduced to conidiogenous cells.

Presently no single character or set of morphological characters separates the subgroups of *Pseudocercospora* that have emerged from DNA sequence data. Analysis of more species is required to provide an expanded view of these groupings. What these data have shown is that several of the morphological features discussed above, used to help group species of *Pseudocercospora*, are not indicative of phylogeny.

Cercospora and Passalora

Species of Cercospora have conspicuously thickened, darkened conidial scars and hyaline to subhyaline, acicular, cylindrical-filiform, multiseptate conidia. These species formed a distinct subclade in Mycosphaerella (Fig. 1). Species with slightly thickened conidial scars and subglobose, ellipsoidal—ovoid, broadly obclavate—fusiform, sparsely septate, usually pigmented conidia are accommodated in Passalora. Several taxa that have been traditionally treated in Cercospora have subsequently been reallocated to Passalora, while Phaeoisariopsis is seen as the synnematous analogue of Passalora (Braun, 1995b).

Passalora, Mycovellosiella and Phaeoramularia

Phaeoramularia comprises Ramularia-like species with pigmented conidia, thickened, darkened and refractive loci, and lacking superficial mycelium. Mycovellosiella is similar, but differs in having superficial mycelium that frequently climbs leaf hairs.

Three species of Mycovellosiella, namely M. bellynckii (CBS 150.49), M. vaginae (CBS 140.34) and M. eucalypti (STE-U 1457), were included in the present study. The former two species clustered with Phaeoramularia dissiliens (CBS 219.77), Passalora arachidicola (PP16) and Passalora personata (PP15), suggesting that these genera are very similar, while M. eucalypti clustered with Phaeoramularia saururi (STE-U 2221). The type specimen of Phaeoramularia was recently examined by U. Braun, who found it to represent a Mycovellosiella species.

Braun (in Braun & Melnik, 1997) assigned Fusicladium levieri Magnus (= F. kaki Hori & Yoshino) to Phaeoramularia but monographic studies of Fusicladium and allied genera recently completed suggest that F. levieri has to be retained in Fusicladium (U. Braun et al., unpublished). Braun (in Braun & Melnik, 1997) reduced Hormocladium Höhn., based on F. levieri, to synonymy with Phaeoramularia. However, Hormocladium is an older name and would have priority unless the name Phaeoramularia is conserved. This species has catenate conidia similar to Phaeoramularia species, but SEM studies reveal that scars and conidial hila are fusicladium-like. Some species of Fusicladium produce conidia singly, in chains, or both, depending on the specimen examined. Catenation of conidia can thus not be used to separate these species into different genera. Hence, Hormocladium, which has catenate conidia, has to be considered a synonym of Fusicladium and is no longer considered a synonym of Phaeoramularia. This complex will be fully addressed in a separate molecular study that is presently underway.

Mycosphaerella africana (STE-U 794) and M. keniensis (STE-U 1084) have no known anamorphs, but cluster together in a clade with Mycovellosiella, Passalora and Phaeoramularia anamorphs. This suggests that they would probably have the same type of anamorph if they exist, or that they may have lost the ability to form anamorphs, and are holomorphic teleomorphs.

Stenella

Mycosphaerella parkii (anamorph: Stenella parkii) (STE-U 353) clustered with M. marksii (STE-U 935), for which no anamorph has yet been described. A recent examination of several cultures of M. marksii

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has shown, however, that hyphae can be verruculose, suggesting that the anamorph, if it exists, will probably be a species of *Stenella*. *Stenella* species are characterized by solitary or catenate conidía with thickened, darkened, refractive conidíal hila and verrucose, superficial hyphae. The formation of verrucose, superficial hyphae separates this genus from *Mycovellosiella* and *Phaeoramularia*. The relationships among these anamorph genera also deserve further investigation.

Ramularia

Three species with Ramularia anamorphs were included in this study, namely Mycosphaerella fragariae (anamorph R. grevilleana) (STE-U 656), M. punctiformis (Ramularia sp.) (CBS 942.97) and R. collocygni (STE-U 2428). While the first two species have typical Mycosphaerella teleomorphs, R. collo-cygni was considered an unusual species of Ramularia (Braun, 1998) because of its curled conidiophores, and conidia with somewhat eccentrically positioned scars. R. collo-cygni was initially considered distinct from Ramularia, and typified the genus Ophiocladium Cavara (as O. hordei Cavara) (Sutton & Waller, 1988). Results of our study show that not only does R. collo-cygni cluster closely with the other two typical Ramularia species, but the teleomorph of R. collo-cygni, if it exists, would likely be a species of Mycosphaerella, Mycosphaerella punctiformis is also the type of the genus Mycosphaerella, and is appropriately included in the main Mycosphaerella clade. The conidia of M. fragariae and M. punctiformis are catenate, but R. collo-cygni belongs to a group of Ramularia species with solitary conidia. The present molecular data support Braun's (1998) taxonomic treatment of Ramularia, based on the premise that separation into two genera based on conidial catenation is untenable.

Colletogloeum and Phaeophleospora

The genera Colletogloeum and Phaeophleospora respectively represent acervular and pycnidial anamorphs of Mycosphaerella. Both genera include species with verruculose, brown conidia situated on brown conidiogenous cells that proliferate percurrently in both genera, but also sympodially in Colletogloeum. Furthermore, Colletogloeum species mostly form 0(-1)-septate conidia, whereas species of Phaeophleospora form aseptate to multiseptate scolecospores. These two genera form one subclade within the cercosporoid clade, together with Uwebraunia. As more species of Phaeophleospora are added in future analyses, this grouping may begin to diverge from

Uwebraunia. For the present, however, Phaeophleospora and Colletogloeum remain easily recognizable anamorph genera.

Clade 2. Mycosphaerella species with Dissoconium anamorphs

The genus Dissoconium was originally described for D. aciculare, a hyperparasite of an Erysiphe species (de Hoog et al., 1983). A further species, D. dekkeri, was later described (de Hoog et al., 1991). Crous & Wingfield (1996) erected the genus Uwebraunia for species thought to be pathogenic to Eucalyptus. However, later comparisons showed that one species, U. lateralis (STE-U 825, 1232, 1233), was morphologically similar and had identical ITS sequences to isolates of D. dekkeri (STE-U 1535) (Crous et al., 1999). This was the first report of a Mycosphaerella teleomorph for a Dissoconium species. Based on sequence data (Crous et al., 1999), it also appeared that Dissoconium represented a separate group in Mycosphaerella. Presently, Dissoconium is accepted for species thought to be hyperparasitic, with wide host ranges. This contrasts with the more host-specific, plant-pathogenic species that are accommodated in Uwebraunia. This also suggests that convergent evolution of morphology has occurred, and thus U. ellipsoidea (STE-U 1224, 1225) and U. juvenis (STE-U 1004, 1005) with similar morphologies reside in separate subclades in the larger Mycosphaerella clade.

Clade 3. Mycosphaerella species with Cladosporium anamorphs

Cladosporium is distinguished from other genera in this complex by its species with pigmented, catenate conidia with thickened, darkened, characteristically protuberant conidial scars and hila. Furthermore, the species usually lack abundant superficial mycelium, and conidia are usually in branched chains, being subglobose, ellipsoidal—ovoid to subcylindrical, and sparsely septate.

Two species of Mycosphaerella are found in this clade, namely M. tassiana (CBS 108.85, 223.31) (anamorph: Cladosporium herbarum) and M. asteroma (CBS 266.53), which has no known anamorph. In some analyses, Cladosporium clustered outside the larger Mycosphaerella clade, although this separation was never strongly supported by bootstrap values. In the final analysis, however, the Cladosporium clade grouped within the main Mycosphaerella clade. These data are preliminary, however, and many more taxa must be added to determine whether all species of Cladosporium are allied with Mycosphaerella, or only the saprobes.

Clade 4: Tapesia species with Ramulispora anamorphs

Several Pseudocercosporella-like species occur on roots and basal parts of small-grain crops. These species are presently included in the anamorph genus Ramulispora, which is confined to species occurring on grasses and cereals (von Arx, 1983) and have discomycetous, Tapesia teleomorphs (Robbertse et al., 1995; Dyer et al., 1996). Although morphologically similar to Pseudocercosporella, species of this genus typically have teleomorphs in Mycosphaerella (Braun, 1995a). In the present analysis, Ramulispora clusters well outside the Mycosphaerella cluster.

Future research

The observations presented in this study suggest that the ITS-1, 5.8S and ITS-2 regions, which contain approximately 300 informative sites in the alignment (data not shown), are suitable for distinguishing species and the various anamorph genera in *Mycosphaerella*. However, other more conserved areas of the genome such as 18S and 28S rDNA should also be analyzed to see if they reveal additional distinct groups within *Mycosphaerella*, or provide better bootstrap support for otherwise weakly supported clades. Other questions are whether saprobic and pathogenic taxa might emerge as distinct genera, and whether any tendency towards coevolution with hosts might exist.

Thoughts and observations presented here represent only a preliminary step towards solving questions of host specificity and determining whether Mycosphaerella is monophyletic. Presently, the majority of the anamorph genera discussed appear to be correctly allied with Mycosphaerella. The distinctions between the anamorph genera in this complex are currently based on small and frequently inconspicuous differences and these need clarification. For the present, the classification system followed by most taxonomists interested in anamorphs of Mycosphaerella remains valid. Delimitation of some anamorph genera in the major Mycosphaerella clade must now be studied in more detail. An unexpected result of this study is that the very large genus Mycosphaerella appears to be an essentially monophyletic group. This holds true despite the many anamorph genera that occur in the genus.

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Table 2. Fungal isolates included for ITS sequence analysis

Accession no.	Teleomorph	Anamorph	Origin
CAL	Mycosphaerella state unknown	Cercospora apii Fresen.	ATCC 12246
K6	Mycosphaerella state unknown	Cercospora beticola Sacc.	Beta, Italy
K3	Mycosphaerella state unknown	Cercospora beticola Sacc.	Chenopodium, Italy
CCA19	Mycosphaerella state unknown	Cercospora canescens Ellis & G. Martin	ATCC 32779 (Mungbean, Taiwan)
CH5	Mycosphaerella state unknown	Cercaspora hayi Calpouzos	ATCC 12234 (Banana, Cuba)
CK35	Mycosphaerella state unknown	Cercospora kikuchii (T. Matsumoto & Tomoy.) M.W. Gardner	Soybean, Illinois, U.S.A.
CK39	Mycosphaerella state unknown	Cercospora kikuchii (T. Matsumotoa & Tomoy.) M.W. Gardner	Soybean, Illinois, U.S.A.
CBS 131.32	Mycosphaerella state unknown	Cercospora nicotiana Ellis & Everh.	Nicotiana, location unknown
CS13	Mycosphaerella state unknown	Cercospora sojina Hara	ATCC 44531 (Soybean, Illinois, U.S.A.)
CBS 108.85	M. tassiana (De Not.) Johanson	Cladosporium herbarum (Pers. : Fr.) Link	Diospyros. Morocco
CBS 223.31	M. tassiana (De Not.) Johanson	Cladosporium herbarum (Pers. : Fr.) Link	Allium, Switzerland
STE-U 784	M. molleriana (Thüm.) Lindau	Colletogloeum mollerianum Crous & M.J. Wingf.	Eucalyptus, California, U.S.A.
STE-U 1214	M. molleriana (Thüm.) Lindau	Colletogloeum mollerianum Crous & M.J. Wingf.	Eucalyptus, Portugal
STE-U 1534	Unknown	Dissoconium aciculare de Hoog, van Oorschot & Hijwegen	CBS 342.82 (ex Erysiphe sp. on Medicago, Germany)
STE-U 1233	M lateralis Crous & M.J. Wingf.	Dissoconium dekkeri de Hoog & Hijwegen	Eucalyptus, Zambia
STE-U 1232	M. lateralis Crous & M.J. Wingf.	Dissoconium dekkeri de Hoog & Hijwegen	Eucalyptus, Zambia
STE-U 825	M. lateralis Crous & M.J. Wingf.	Dissoconium dekkeri de Hoog & Hijwegen	Eucalyptus, South Africa
STE-U 1535	M. lateralis Crous & M.J. Wingf.	Dissoconium dekkeri de Hoog & Hijwegen	CBS 567.89 (Juniperus, The Netherlands)
CBS 150.49	Mycosphaerella state unknown	Mycovellosiella bellynckii (Westend.) Constant.	Cynanchum, Switzerland
STE-U 1457	Mycosphaerella state unknown	Mycovellosiella eucalypti Crous & A.C. Alfenas	Eucalyptus, Brazil
CBS 140.34	Mycosphaerella state unknown	Mycovellosiella vaginae (W. Krüger) Deighton	Saccharum, Taiwan
PF7	M. fijiensis M. Morelet	Paracercospora fijiensis (M. Morelet) Deighton	ATCC 22116 (Banana, Philippines)
PF8	M. fijiensis M. Morelet	Paracercospora fijiensis (Morelet) Deighton	ATCC 22117 (Banana, Hawaii)
PFD9	M. fijiensis var. difformis J.L. Mulder & R.H. Stover	Paracercospora fijiensis var. difformis (J.L. Mulder & R.H. Stover) Deighton	ATCC 36054 (Musa, Honduras)
PP16	Mycosphaerella arachidis Deighton	Passalora arachidicola (Hori) U. Braun	MPPD L2122 (Peanut, Oklahoma, U.S.A.)
PP15	M. berkeleyi W.A. Jenkins	Passalora personata (Berk. & M.A. Curt.) S.H. Khan & M. Kamal	MPPD L2121 (Peanut, Oklahoma, U.S.A.)
CBS 277.39	Mycosphaerella state unknown	Passalora robinae (Shear) S. Hughes (?)	Robinia, U.S.A.
CBS 194.47	Mycosphaerella state unknown	Phaeoisariopsis griseola (Sacc.) Ferr.	Phaseolus, Portugal
STE-U 1346	M. suttoniae Crous & M.J. Wingf. F.A. Ferreira & B. Sutton	Phaeophleospora epicoccoides (Cooke & Massee) Crous,	Eucalyptus, Indonesia
CBS 219.77	Mycosphaerella state unknown	Phaeoramularia dissiliens (Duby) Deighton	Vitis, Iraq
STE-U 2221	Mycosphaerella state unknown	Phaeoramularia saururi (Ellis & Everh.), Crous & El-Gholl	Saururus, U.S.A.
STE-U 1266	Mycosphaerella state unknown	Pseudocercospora basiramifera Crous	Eucalyptus, Thailand
STE-U 1106	M. colombiensis Crous & M.J. Wingf.	Pseudocercospora colombiensis Crous & M.J. Wingf.	Eucalyptus, Colombia
PCR18	M. cruenta Latham	Pseudocercospora cruenta (Sacc.) Deighton	ATCC 262271 (Cowpea, Puerto Rico)
STE-U 801	M. crystallina Crous & M.J. Wingf.	Pseudocercospora crystallina Crous & M.J. Wingf.	Eucalyptus, South Africa
CBS 130.32	Mycosphaerella state unknown	Pseudocercospora eriodendri (Raciborski) U. Braun	Eriodendron, Somalia
STE-U 760	M. heimii Crous	Pseudocercospora heimii Crous	Eucalyptus, Madagascar
STE-U 1312	M. heimioides Crous & M.J. Wingf.	Pseudocercospora heimioides Crous & M.J. Wingf.	Eucalyptus, Indonesia
STE-U 1360	M. irregulariramosa Crous & M.J. Wingf.	Pseudocercospora irregulariramosa Crous & M.J. Wingf.	Eucalyptus, South Africa
PM10	M. musicola J.L. Mulder	Pseudocercospora musae (Zimm.) Deighton	ATCC 22115 (Banana, Philippines)
PM11	M. musicola J.L. Mulder	Pseudocercospora musae (Zimm.) Deighton	ATCC 36143 (Musa, Honduras)

STE-U 1264	Mycosphaerella state unknown	Pseudocercospora natalensis Crous & T. Coutinho	Eucalyptus, South Africa
STE-U 1458	Mycosphaerella state unknown	Pseudocercospora paraguayensis (Kobayashi) Crous	Eucalyptus, Brazil
CBS 282.66	Mycosphaerella state unknown	Pseudocercospora rhapisicola (Tominaga) Goh & W.H. Hsieh	Rhapis, Japan
CBS 149.53	Mycosphaerella state unknown	Pseudophaeoramularia angolensis (T. Carvalho & O. Mendes) U. Braun	Citrus, Angola
STE-U 2428	Mycosphaerella state unknown	Ramularia collo-cygni B. Sutton & J.M. Waller	Barley, Germany
STE-U 656	M. fragariae (Tul.) Lindau	Ramularia grevilleana (Tul. & C. Tul.) Jørst.	Strawberry, South Africa
CBS 942.97	M. punctiformis (Pers. : Fr.) Starbäck	Ramularia sp.	Quercus, Belgium
RAE22	Tapesia state unknown	Ramulispora aestiva (Nirenberg) E.L. Stewart & Crous	Wheat, Washington, U.S.A.
RAN45	Tapesia state unknown	Ramulispora anguioides (Nirenberg) Crous	ATCC 60971 (Wheat, Germany)
STE-U 353	M. parkii Crous & M.J. Wingf.	Stenella parkii Crous & Alfenas	Eucalyptus, Brazil
CBS 336.33	Mycosphaerella state unknown	Stigmina platani (Fuckel) Sacc.	Platanus, Location unknown
STE-U 794	M. africana Crous & M.J. Wingf.	Unknown	Eucalyptus, South Africa
CBS 266.53	M. asteroma (Fr.) Lindau (?)	Unknown	Polygonatum, Switzerland
STE-U 1084	M. keniensis Crous & T. Coutinho	Unknown	Eucalyptus, Kenya
STE-U 935	M. marksii Carnegie & Keane	Unknown	Eucalyptus, Australia
STE-U 1224	M. ellipsoidea Crous & M.J. Wingf.	Uwebraunia ellipsoidea Crous & M.J. Wingf.	Eucalyptus, South Africa
STE-U 1225	M. ellipsoidea Crous & M.J. Wingf.	Uwebraunia ellipsoidea Crous & M.J. Wingf.	Eucalyptus, South Africa
STE-U 1004	M. juvenis Crous & M.J. Wingf.	Uwebraunia juvenis Crous & M.J. Wingf.	Eucalyptus, South Africa
STE-U 1005	M. juvenis Crous & M.J. Wingf.	Uwebraunia juvenis Crous & M.J. Wingf.	Eucalyptus, South Africa

STE-U = Culture collection of the Department of Plant Pathology, University of Stellenbosch, South Africa