INTRODUCTION

Sooty blotch and flyspeck (SBFS) is a fungal disease complex that causes a well known problem in apple fruit production (Schweinitz 1832, Colby 1920, Williamson & Sutton 2000). The number of fungal species causing sooty blotch and flyspeck of apples was previously underestimated (Johnson et al. 1997) and more than 60 taxa are currently known (Díaz Arias et al. 2010), of which the majority are either undescribed or taxonomically unresolved (Batzer et al. 2008, Sun et al. 2008, Schoch et al. 2009, Yang et al. 2010).

The general morphology type termed ‘sooty blotch’ describes species that form colonies on apples that are characterised by superficially spreading, more or less densely branched, dark olive-green, greyish, or brownish black hyphae or mycelial strands with or without sclerotium-like structures or fruiting bodies. Fungal growth of similar or differently shaded or branched colonies of the same or different fungal species can coalesce, resulting into a black or sooty appearance of parts of or the entire apple surface. Sooty blotch fungi are epiphytes and do not cause pre-harvest losses or fruit decay (Colby 1920). In some cases they can cause desiccation of apple fruits during post harvest and storage (R. Godec, Agricultural Institute of Slovenia, pers. comm.). However, sooty blotch reduces the market value of apples and has limited the growth in organic apple production (Williamson & Sutton 2000, Batzer et al. 2005, Yue et al. 2007).

Colby (1920) provided a thorough review on sooty blotch of pomaceous fruits, and accepted that a single species, namely Gloeodes (Dothidea) pomigena (= Phyllachora pomigena), was the responsible casual agent. According to the arrangement of spreading and branching mycelia, sooty blotch was classified into different types such as ramose, fuliginous, punctate, rimate and ridged honeycomb (Groves 1933, Batzer et al. 2005). Furthermore, various mostly dothidealean ascomycetes were identified and classified on the basis of overall morphological characters as species of Colletogloeum, Discoconium, Peltaster, Pseudocercospora, Pseudocercosporella and Xenostigmata (Batzer et al. 2005, Crous et al. 2009b).

During inventories of SBFS fungi in Germany and Slovenia, a number of isolates were retrieved from apples showing sooty blotch that corresponded either to the ramose (RS) or a dense fuliginous (FG) phenotype. Molecular sequencing of phylogenetic marker genes identified these strains as sooty blotch isolates currently placed in genera such as Devriesia, Pseudocercospora or Pseudocercosporella. However, none of these identifications has until now been supported by sequences of reference strains of these genera, and therefore the aim of the present study was to compare these sooty blotch isolates with authentic reference strains, and to resolve their taxonomy.
pellet characterised by a homogeneous pattern of sooty blotch were cut off from selected apples with a sterile scalpel and gently surface sterilised with cotton and 70 % ethanol. Pieces of sooty blotch mycelium were removed with a sterile scalpel and placed onto 2 % potato-dextrose agar (PDA; Crous et al. 2009d). Three to four isolations were made from the same sooty blotch colony. The growth habit on the natural substratum was photographically documented (×10–63 magnification). The remainder of the colony and subtending apple peel was dried down, pressed between layers of filter paper, and retained as voucher specimen. Petri dishes were incubated on the laboratory bench for 10–20 d, until colonies started to develop. Colonies were hyphal tipped, and transferred to clean PDA and synthetic nutrient-poor agar (SNA; Crous et al. 2009d) slants for preservation at 4 °C. The obtained strains were compared with a set of sooty blotch strains isolated from apples collected in Germany (Feldmann 2005).

DNA isolation, amplification and analyses

Genomic DNA was isolated from fungal mycelium grown on MEA, using the UltraCleanTM Microbial DNA Isolation Kit (MoBio Laboratories, Inc., Solana Beach, CA, USA) according to the manufacturer’s protocols. The primers V9G (de Hoog & Gerrits van den Ende 1998) and LR5 (Vilgalys & Hester 1990) were used to amplify part of the nuclear rDNA operon spanning the 3’ end of the 18S rRNA gene (SSU), the internal transcribed spacer 1, the 5.8S rRNA gene, the internal transcribed spacer 2 (ITS) and the first 900 bases at the 5’ end of the 28S rRNA gene (LSU). The primers ITS4 (White et al. 1990) and LR0R (Rehner & Samuels 1994) were used as internal sequence primers to ensure good quality sequences over the entire length of the amplicon. Partial gene sequences were determined for translation elongation factor 1-α (TEF) as described by Crous et al. (2006c). The PCR conditions, sequence alignment and subsequent phylogenetic analysis followed the methods of Crous et al. (2006a, 2009a). Sequences were compared with the sequences available in NCBI’s GenBank nucleotide (nr) database using a MegaBLAST search and results are discussed in the relevant species notes where applicable. Alignment gaps were treated as new character states. Sequence data were deposited in GenBank (Table 1) and alignments in TreeBASE (www.treebase.org).

Morphology

Isolates were established on 2 % malt extract agar (MEA), PDA, SNA and oatmeal agar (OA; Crous et al. 2009d), and subsequently incubated at 25 °C under near-ultraviolet light to promote sporulation. Preparations from cultured fungal colonies were mounted on glass slides with clear lactic acid for microscopic examination after 7 d of incubation. Thirty measurements per relevant microscopic structure were gathered with a compound microscope under ×100 magnification. Colony colours on MEA and OA (surface and reverse) were determined using the colour charts of Rayner (1970) after 1 mo at 25 °C in the dark. Reference strains are maintained in the culture collection of the Centraalbureau voor Schimmelcultures (CBS-KNAW), Utrecht, the Netherlands, the working collection (CPC) of P.W. Crous, and at the Agricultural Institute of Slovenia (Table 1). Nomenclatural novelties and descriptions were deposited in MycoBank (www.MycoBank.org; Crous et al. 2004).

RESULTS

Phylogenetic analysis

Approximately 1 700 bases, spanning the ITS and LSU regions, were obtained for isolates listed in Table 1. The LSU region was used in the phylogenetic analysis for the generic placement

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<th>Species</th>
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1 CBS: CBS-KNAW Fungal Biodiversity Centre, Utrecht, The Netherlands; CPC: Culture collection of P.W. Crous, housed at CBS; SK: S. Kern isolation number (INRIS, Univ. Bonn); JF: J. Frank isolation number (Agricultural Institute of Slovenia).

2 ITS: Internal transcribed spacers 1 and 2 together with 5.8S nrDNA; LSU: 28S nrDNA; SSU: 18S nrDNA (not used in phylogenetic analyses due to limited resolution); TEF: Translation elongation factor 1-α.
The manually adjusted LSU alignment contained 76 taxa (including the *Phaeobotryosphaeria visci* outgroup sequence) and, of the 731 characters used in the phylogenetic analysis, 171 were parsimony-informative, 95 were variable and parsimony-uninformative and 465 were constant. Only the first 1 000 equally most parsimonious trees were retained from the heuristic search, the first of which is shown in Fig. 1 (TL = 758, CI = 0.493, RI = 0.844, RC = 0.416). The phylogenetic tree of the LSU region (Fig. 1) showed the isolates obtained in this study to cluster in three lineages, namely *Devriesia* s.l. and two novel genera described below, the first phylogenetically related to *Penidiella* and the second to a clade containing *Mycosphaerella intermedia*, *Mycosphaerella marksi* and *Mycosphaerella madeirensis*.

Only ITS sequences obtained from the GenBank nucleotide database with an identity of 95 % and higher to the four novel species were included in the ITS analysis. The 1 000 equally most parsimonious trees obtained from the GenBank nucleotide database with an identity of 95 % and higher to the four novel species were included in the ITS analysis. The scale bar shows 10 changes, and, of the 1 000 trees, 54 % were within 10 changes from the strict consensus tree.
Fig. 2 The first of 1 000 equally most parsimonious trees obtained from a heuristic search with 100 random taxon additions of the ITS sequence alignment. The scale bar shows 10 changes, and bootstrap support values from 1 000 replicates are shown at the nodes. The four novel species described in this study are indicated by coloured boxes and sequences from non-apple hosts are red text. Strain numbers of epitype cultures are shown in bold and branches present in the strict consensus tree are thickened. The tree was rooted to a sequence of *Cladosporium bruhnei* (GenBank accession EF679337).
species described in this study were added to the alignment. The manually adjusted ITS alignment contained 64 taxa (including the *Cladosporium bruhnei* outgroup sequence) and, of the 489 characters used in the phylogenetic analysis, 165 were parsimony-informative, 71 were variable and parsimony-uninformative and 253 were constant. Only the first 1 000 equally most parsimonious trees were retained from the heuristic search, the first of which is shown in Fig. 2 (TL = 450, CI = 0.747, RI = 0.949, RC = 0.709). The phylogenetic tree of the ITS region (Fig. 1) showed the novel species obtained in this study to cluster with other sequences in GenBank lodged as 'Pseudocercospora sp.' and 'Pseudocercosporella sp.' and these could represent identical or closely related cryptic species (see species notes below).

**Taxonomy**

Several taxonomic novelties were found to be associated with SBFS blemishes on apple surfaces (Fig. 3) that do not match any species presently described, though some could be linked to taxa deposited under preliminary names in GenBank. These genera and species are described as new below.

**Devriesia pseudoamericana** Jana Frank, B. Oertel, Schroers & Crous, sp. nov. — MycoBank MB516839, Fig. 4

*Teleomorph. Unknown.*

*Devriesiae americanae* morphologic similes, sed conidis longioribus, (7–)10–20(–30) × 2–3 µm.

*Etymology.* Named after its morphological similarity, and close phylogenetic relationship to *D. americana*.

*Colonies* on OA. *Mycelium* consisting of branched, septate, brown, finely verruculose, 2–3 µm wide hyphae; chlamydospores intercalary, globose, 5–7 µm diam, brown, smooth. *Conidiophores* terminal and lateral on hyphae, highly variable in length, at times macroconidial, but also micronematous, reduced to conidiogenous cells; cylindrical, straight to curved, brown, smooth, 10–50 × 2–3 µm, 0–6-septate. *Conidiogenous* cells terminal and lateral on conidiophores, 5–10 × 2–3 µm, proliferating sympodially with terminal and lateral polyblastic loci; scars 1–1.5 µm wide, flattened, somewhat darkened and thickened, not refractive. *Conidia* brown, finely verruculose, subcylindrical, at times somewhat swollen, appearing narrowly ellipsoid, straight to curved or once geniculate, irregular, apex obtuse, base truncate, 0–10–septate, (7–)10–20(–30) × 2–3 µm, occurring in irregular branched chains; hila somewhat
thickened and darkened, 1–1.5 µm diam. Conidia on SNA poorly developed, much narrower, with less septa and more fusoid-ellipsoidal.

Culture characteristics — Colonies after 1 mo at 25 °C in the dark on MEA spreading, with sparse to moderate aerial mycelium; surface folded, irregular, crumpled, at times breaking the agar surface, olivaceous-grey, with thin, iron-grey margin; reverse iron-grey; colonies reaching up to 25 mm diam after 1 mo. On OA spreading, flattened, with sparse aerial mycelium in middle of colony; aerial mycelium pale olivaceous-grey, outer region olivaceous-grey; colonies reaching 25 mm diam after 1 mo. On SNA appearing woolly, erumpent, not spreading, smoke-grey, with sparse aerial mycelium, reaching 8 mm diam after 1 mo. Cultures of *D. pseudoamericana* do not grow at 37 °C.


Notes — Members of *Devriesia* s.l. share morphological similarities in their conidiophores, branched conidial chains, somewhat darkened conidial hila and scars, as well as their terminal and intercalary chlamydospores formed in culture. According to LSU rDNA based phylogenetic analyses (Fig. 1), the genus is, however, paraphyletic. *Devriesia* comprises at least four lineages, of which three are distantly related to *D. staurophora*, the type species of the genus. The others are represented by i) *D. pseudoamericana* and *D. americana* (CBS 117726); ii) *D. strelitziae*; and iii) *D. hiliiana*, *D. lagerstroemiae* (Crous et al. 2009b), *D. strelitziae* (Arzanlou et al. 2008) and *Teratosphaeria knoxdavesii* (Crous et al. 2008), which lacks a known anamorph in culture. The ITS sequences of *D. pseudoamericana* and *D. americana* (GenBank AY251068) have an 88 % (441/496 nucleotides) identity. An ITS sequence lodged in GenBank (FN549915) as *Devriesia* sp. matches *D. pseudoamericana* with 99 % (463/467 nucleotides, 4/467 gaps) and appears to be the same species. The strain (CBS 529.82) from which that sequence derived is lodged in the CBS-KNAW database as *Septonema ochraceum* isolated from needles of *Picea abies* in the Netherlands. More collections are needed to determine whether this implies a wider host range for *D. pseudoamericana*. No type material is available for *Septonema ochraceum* and Koukol (2010) failed to find strains that could be regarded as representative of the species using fresh collections and cultures lodged in public collections under that name. Koukol (2010) included CBS 529.82 in his study and also observed the relationship of this strain to *D. americana* but failed to find morphological differences between these strains.

Taxa presently accommodated in different lineages of *Devriesia* s.l. share a different ecology to members of *Devriesia* s.str. (Seifert et al. 2004), which usually occur in soil, and are thermotolerant (able to grow at high temperatures, namely to survive exposure to 75 °C for 30 min; Samson et al. 2000). In contrast, members of *Devriesia* s.l. are usually associated with leaf spots, or occur on dead plant debris as saprobes, and are not able to grow above 37 °C (Crous et al. 2007b, 2009b, Koukol 2010). In spite of *Devriesia* being paraphyletic, we refrain from describing

Fig. 4 *Devriesia pseudoamericana* (CPC 16174). a. Colony on MEA; b. colony on SNA; c, d, f, g. conidiophores; e, h. conidia in chains; i. chlamydospores. — Scale bars = 10 µm.
novel genera formally because more taxa and strains have to be added first for resolving possible synapomorphies supporting their phylogenetic concept morphologically.

**Microcyclospora** Jana Frank, Schroers & Crous, *gen. nov.*
— MycoBank MB516842

Hyphomycetes. Mycelium ex hyphis ramosis, septatis, pallide brunneis, levi-bus, 2–3 µm latis compositum. Conidiophora in cellulis conidiogenis reducta, in hyphis lateralt integrata, mono- vel polystylistica, subdenticulata, pallide brunnea, levia. Conidia scolecospora, cylindracea, recta vel diverse curvata, flexuosa, apice obtuso, basi truncata, uni- ad multisepitata, ad septa lentier constricta, levia, pallide brunnea, guttulata, aggregata in massa mucosa; hila inconspicua, neque incrassata neque fuscata; in cultura cum formatione microcyclica conidiorum.

Type species. *Microcyclospora pomicola* Jana Frank, Schroers & Crous, *sp. nov.*

Etymology. Named after its resemblance to *Pseudocercospora*, and prominent microcyclic conidiation.

Hyphomycetous. Mycelium consisting of branched, septate, pale brown, smooth, 2–3 µm wide hyphae. Conidiophores reduced to conidiogenous cells, integrated, mono- to polystylistica, lateral on hyphae, subdenticulata, 1 µm wide, 1–2 µm tall, pale brown, smooth. Conidia scolecosporous, cylindrically, straight to variously curved, flexuous, apex obtuse, base truncate, 1–multi-septate, somewhat constricted at septa, smooth, pale brown, guttulate, aggregated in mucoid masses; hila not thickened or darkened; microcyclic conidiation observed in culture.

Notes — Members of the genus *Microcyclospora* have in the past been accommodated in *Pseudocercospora* based on their pigmented conidiophores and pigmented, scolecosporous, transversely septate conidia (Batzer et al. 2005). *Microcyclospora* can be distinguished from *Pseudocercospora* s.str. (Crous et al. 2006b, 2007a) in that conidiophores are mostly reduced to solitary conidiogenous cells on hyphae which appear to be mono- to polystylistica (non fasciculate), conidia that occur in mucoid masses, and commonly undergo microcyclic conidiation. The closest phylogenetic neighbours of *Microcyclospora* are species of *Penidiella* (Fig. 1). Based on comparisons of our ITS sequences to those available on GenBank, more species than the three described here can be predicted in this genus (data not shown).

**Microcyclospora malicola** Jana Frank, Schroers & Crous, *sp. nov.*
— MycoBank MB516843; Fig. 5

Teleomorph. Unknown.

Mycelium ex hyphis ramosis, septatis, pallide brunneis, levi-bus, 2–3 µm latis compositum. Conidiophora in cellulis conidiogenis reducta, in hyphis lateralt integrata, mono- vel polystylistica, subdenticulata, 1 µm lata, 1–2 µm alta, pallide brunnea, levia. Conidia scolecospora, cylindracea, recta vel diverse curvata, flexuosa, apice obtuso, basi truncata, (1–)5–7(–13)-septata, levia, pallide brunnea, guttulata, (30–)45–75(–120) × (2–)2.5(–3) µm; hila inconspicua.

Etymology. Named after its host, *Malus*.

Colony on SNA. Mycelium consisting of branched, septate, pale brown, smooth, 2–3 µm wide hyphae. Conidiophores reduced to conidiogenous cells, integrated, lateral on hyphae, mono- to polystylistica, subdenticulata, 1 µm wide, 1–2 µm tall, pale brown, smooth. Conidia scolecosporous, cylindrically, straight to variously curved, flexuous, apex obtuse, base truncate, (1–)5–7(–13)-septate, somewhat constricted at septa, smooth, pale brown, guttulate, (30–)45–75(–120) × (2–)2.5(–3) µm, after 7 d, and similar after 1 mo; hila not thickened nor darkened; microcyclic conidiation observed in culture.

Culture characteristics — Colonies after 1 mo at 25 °C in the dark on MEA flat, spreading, with moderate, smoke-grey aerial mycelium that collapses with age, becoming iron-grey; margins...
smooth, regular, lobate; reverse iron-grey; colonies reaching 5–6 mm diam after 7 d, and up to 40 mm diam after 1 mo. On OA similar, aerial mycelium sparse, collapsing, becoming iron-grey; colonies reaching 5–6 mm diam after 7 d, and up to 50 mm diam after 1 mo. Microcyclic conidiation commonly observed on all media in culture.


Notes — Considerable morphological variation was observed within *M. malicola*, and it was initially expected that isolate CPC 16186 could represent a distinct species to isolate CPC 16172 based on differences in conidial dimensions and growth rate in culture. However, based on sequence data of the ITS region they were identical, thus we chose to treat them as representative of a single taxon until more strains have been collected. A megablast search of NCBI's GenBank nucleotide database revealed six additional accessions with high identity to *M. malicola* (Fig. 2), currently filed at GenBank under 'Pseudo­cercospora'. The strains from which GenBank FJ808758 and FJ808755 were generated probably represent *M. malicola* originating from Serbia, while sequence DQ363418 represents another strain from Germany. The three other accessions (GenBank FJ438380, AY598858 and AY598857 from the USA) probably represent currently undescribed *Microcyclospora* species because they accumulate up to 6 additional substitutions in their ITS sequences when compared with *M. malicola*. The TEF sequence of the type strain of *M. malicola* is 90 % identical (351/387 bases and 13 gaps) to the second strain of the species sequenced in this study, and 87 % identical (119/136 bases and 2 gaps) and 84 % identical (245/289 bases and 13 gaps) to the sequences of the ex-type strains of *P. pomicola* and *M. tardicrescens*, respectively. It is quite possible that *M. malicola* represents a species complex, but this can only be resolved once more strains of the species are collected.

**Microcyclospora pomicola** Jana Frank, B. Oertel, Schroers & Crous, sp. nov. — MycoBank MB516844; Fig. 6

Teleomorph. Unknown.

*Microcyclosporae malicola*eae morphologicae similis, sed disimilidinitibus in sequentibus nucleotidium (ITS) in sequentibus diversis sequentibus culturae typicae distinguuntur: 136 (T/C), 142 (C/T), 181 (A/G), 187 (T/C), 256 (T/A), 265 (C/T), 271 (A/G), 278 (A/G), 279 (A/T), 556 (A/C), 562 (G/A), 569 (C/T), 570 (T/C), 578 (G/C), extra A inter 581 et 582.

**Etymology.** Named after its occurrence on pome fruit (apples).

Colonies on SNA. Mycelium consisting of branched, septate, pale brown, smooth, 2–3 µm wide hyphae. Conidiophores reduced to conidiogenous cells, integrated, lateral on hyphae, mono- to polyblastic, subdenticulate, 1 µm wide, 1–2 µm tall, pale brown, smooth. Conidia scocolesporous, cylindrical, straight to variously curved, flexuous, apex obtuse, base truncate, somewhat constricted at septa, smooth, pale brown, gutulate, (15–)35–55(–65) × (2–)2.5–3 µm, (3–)4–7-septate after 7 d, (15–)50–75(–120) × (2–)2.5–3 µm, 1–13-septate after 1 mo; hila not thickened nor darkened; microcyclic conidiation observed in culture.

Culture characteristics — Colonies after 1 mo at 25 °C in the dark on MEA spreading, with sparse to moderate aerial mycelium; surface smooth due to collapsing, wet aerial mycelium; margin regular, lobate; surface olivaceous-grey, becoming iron-grey due to collapsing aerial mycelium; reverse iron-grey; colonies reaching 6–8 mm diam after 7 d, and up to 40 mm diam after 1 mo. On OA flattened, submerged, spreading with sparse aerial mycelium and even, smooth margins; colony growth rate similar to that observed on MEA.

**Specimens examined.** GERMANY, Baden-Württemberg, Friedrichshafen, on *Malus domestica* fruit surface, 5 Nov. 1996, S. Kern, CBS H-20412 holotype, culture ex-type 43.1a = CPC 16175 = CBS 126141; Baden-Württemberg, Friedrichshafen, on *M. domestica* fruit surface, 5 Nov. 1996, S. Kern, 51.2a = CPC 16173 = CBS 126140.

Notes — The TEF sequence of the type strain of *M. pomicola* is 90 % identical (351/387 bases and 136 gaps) to the second strain of the species sequenced in this study, and 87 % identical (119/136 bases and 2 gaps) and 84 % identical (150/177 bases and 11 gaps) to the sequences of the ex-type strains of *M. malicola* and *M. tardicrescens*, respectively. It is quite possible that *M. pomicola* represents a species complex, but this can only be resolved once more strains of the species are collected.

**Microcyclospora tardicrescens** Jana Frank, Schroers & Crous, sp. nov. — MycoBank MB516845; Fig. 7

Teleomorph. Unknown.

*Microcyclosporae malicola*eae morphologicae similis, sed conidii minoribus, (15–)35–55(–60) × (1.5–)2(–2.5) µm, in cultura tarde crescent et tamen signis nucleotidium singularium affinis in ITS sequentibus culturae typicae in positionibus 190 (C), 213 (quod additur C), 557 (C), 559 (T) distinguuntur.

**Etymology.** Named after its slow growth rate in culture.

Colonies on SNA. Mycelium consisting of branched, septate, pale brown, smooth, 2–3 µm wide hyphae. Conidiophores reduced to conidiogenous cells, integrated, lateral on hyphae,
Mono- to polyblastic, subdenticulate, 1 µm wide, 1–2 µm tall, pale brown, smooth. Conidia scolecosporous, cylindrical, straight to variously curved, flexuous, apex obtuse, base truncate, 1–6-septate, somewhat constricted at septa, smooth, pale brown, guttulate, (15–)35–55(–60) × (1.5–)2(–2.5) µm; hila not thickened nor darkened; microcyclic conidiation observed in culture; older conidia develop intercalary chlamydospores that are medium brown, up to 5 µm wide.

Culture characteristics — Colonies after 1 mo at 25 °C in the dark on MEA erumpent, with sparse smoke-grey aerial mycelium; colonies reaching up to 2 mm after 7 d, and after 7 d on SNA. Conidia of $M. \text{pomicola}$ and $M. \text{malicola}$ are similar in morphology (1.5–)2(–2.5) µm; hila of $M. \text{tardicrescens}$ is prominent microcyclic conidiation.

Although $M. \text{malicola}$ and $M. \text{pomicola}$ are similar in morphology and culture characteristics after 30 d, they can be distinguished after 7 d on SNA. Conidia of $M. \text{malicola}$ are longer, (30–45–75(–120) µm, 1–13-septate, than those of $M. \text{pomicola}$, (15–)35–55(–65) µm, 3–7-septate. Furthermore, after 7 d on MEA, colonies of $M. \text{malicola}$ are 5–6 mm diam, while those of $M. \text{pomicola}$ grow somewhat faster, reaching 6–8 mm diam. The TEF sequence of the ex-type strain of $M. \text{tardicrescens}$ is 84 % identical (245/289 bases and 13 gaps) and 84 % identical (150/177 bases and 11 gaps) to the sequences of the ex-type strains of $M. \text{pomicola}$ and $M. \text{malicola}$, respectively.

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Fig. 7 Microcyclospora tardicrescens (CPC 16187). a. Colony on MEA; b. colony on SNA; c, d. conidiogenous cells giving rise to conidia; e, f. conidia. — Scale bars = 10 µm.

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Microcyclosporella Jana Frank, Schroers & Crous, gen. nov. — MycoBank MB516840

Hyphomycetes. Mycelium ex hyphis pallide brunneis, levibus vel subtiliter verruculosis, ramosis, septatis, 2–3.5 µm latis compositum, interdum strato mucoso, locis conidiogenis ex transverse integratis, truncatis. Conidiophora saepi in cellulis conidiogenis reducta. Cellulae conidiogenae integratae, intercalares in hyphis, raro terminales, cylindraceae vel doliformes, pallide brunneae, sed hyalinae in partibus sentinoidibus coloniae, leves, mono- vel polyblasticae, sympodiales, locis conidiogenis inconspicuis, truncatis, non incrassatis, non refractivis, pallide brunneis vel hyalinis. Conidia hyalina, levia, subcylindraceae, anguste oblata vel fusiformia, apice acute rotundato, basi obconice truncata, guttulata, 0–6 transverse septata, vulgo cum formatione microcyclica conidiorum.

Type species. Microcyclosporella mali Jana Frank, Schroers & Crous, sp. nov.

Etymology. Named after its resemblance to Pseudocercosporella, and its prominent microcyclic conidiation.

Hyphomycetous. Mycelium consisting of pale brown, smooth to finely verruculose, branched, septate, 2–3.5 µm wide hyphae, at times covered in a mucoid layer, with integrated, lateral, truncate conidiogenous loci. Conidiophores mostly reduced to conidiogenous cells. Conidiogenous cells integrated, intercalary on hyphae, rarely terminal, cylindrical to doliform, pale brown, but hyaline if occurring in yeast-like sectors of colonies, smooth, mono- or polyblastic, proliferating sympodially; loci inconspicuous, truncate, unthickened, not darkened, pale brown to hyaline. Conidia hyaline, smooth, subcylindrical to narrowly oblative or narrowly fusoid with acutely rounded apex and obconically truncate base, guttulate, 0–6 transversely septate; microcyclic conidiation common.

Microcyclosporella mali Jana Frank, Schroers & Crous, sp. nov. — MycoBank MB516841; Fig. 8

Teleomorph. Unknown.

Mycelium ex hyphis pallide brunneis, levibus vel subtiliter verruculosis, ramosis, septatis, 2–3.5 µm latis compositum. Conidiophora saepi in cellulis conidiogenis reducta. Cellulae conidiogenae integratae, intercalares in hyphis, raro terminales, cylindraceae vel doliformes, hyalinae vel pallide

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Notes — After 30 d on SNA, $M. \text{malicola}$ and $M. \text{pomicola}$ have larger conidia (15–120 × 2–3 µm), than $M. \text{tardicrescens}$ (15–60 × 1.5–2.5 µm). Furthermore, $M. \text{tardicrescens}$ also forms intercalary chlamydospores, which are lacking in the other two species, and has a slower growth rate in culture.
brunneae, leves, mono- vel polyblasticae, sympodiales. Conidia hyalina, levia, subcylindracea, anguste obclavata vel fusiformia, apice acute rotundato, basi obconice truncata, guttulata, aseptata usque ad multiseptata, hils inconspicuius.

Etymology. Named after its host, Malus.

Colonies on SNA. Mycelium consisting of pale brown, smooth to finely verruculose, branched, septate, 2–3.5 µm wide hyphae, at times covered in a mucoid layer, with integrated, lateral, truncate conidiogenous loci. Conidiophores mostly reduced to conidiogenous cells. Conidiogenous cells integrated, intercalary on hyphae, 5–7 × 1–2 µm, rarely terminal, cylindrical to doliiform, hyaline to pale brown, smooth, mono- or polyblastic, proliferating sympodially; loci inconspicuous, truncate, unthickened, not darkened, 1–2 µm diam; conidiogenous cells pale brown if integrated on mycelium, but hyaline when occurring in slime, yeast-like parts of colonies, which give rise to conidia via microcyclic conidiation as well as via hyphal loci. Conidia hyaline, smooth, subcylindrical to narrowly obclavate or narrowly fusoid with acutely rounded apex and obconically truncate base, guttulate, (0–)3–6(–10)-septate, (15–)17–25(–40) × 2.5(–3) µm; conidial hila unthickened, not darkened, 1–2 µm diam; microcyclic conidiation common, sporulating profusely on SNA.

Culture characteristics — Colonies after 1 mo at 25 °C in the dark on MEA with sparse aerial mycelium and even, lobate to somewhat feathery margins; surface folded, crumpled, olivaceous-grey with patches of iron-grey and pale olivaceous-grey; reverse olivaceous-black; colonies reaching up to 30 mm diam after 1 mo. On OA similar, except surface not folded and crumpled, but more flattened and spreading; colonies reaching 40 mm diam after 1 mo. On SNA immersed, with sparse grey-olivaceous mycelium and feathery margins, reaching 30 mm diam after 1 mo; colonies developing patches that appear yeast-like, white and slimy. Isolate CBS 126136 exhibited the ability to form microconidia or spermatia in culture, which were hyaline, bacilliform, aseptate, 2–3 × 1–1.5 µm (Fig. 8). This was rarely observed, however, and the formation and role of these structures remain unresolved.


Notes — Members of the genus Microcyclosiorella have thus far been referred to as representative of Pseudocercosporella due to their hyaline conidiophores, and transversely septate, hyaline scolecosporous conidia with inconspicuous hila (Batzer et al. 2005). Although the genus Pseudocercosporella has been shown to be polyphyletic within the Mycosphaerellaceae (Crous 2009, Crous et al. 2003, 2009b, c), it has thus far not been possible to resolve the correct placement of the SBFS isolates, as the type species of Pseudocercosporella, P. pomoeae (= P. bakeri, see Braun 1995), has not been known from culture. In the present study this matter has finally been resolved, as a fresh collection from its centre of origin has allowed us to designate an epitype for P. bakeri (see below). The closest phylogenetic sisters of Microcyclosiorella are Mycosphaerella intermedia, Mycosphaerella markiis and Mycosphaerella madeirae (Fig. 1). A megablast search of NCBI’s GenBank nucleotide database revealed several accessions with high identity to

Fig. 8  Microcyclosiorella mali (CPC 16184). a. Colony on OA; b. colony on MEA; c. spermatia; d. spermatogenous cells; e. conidiophore with conidia; f–j. conidia with microcyclic conidiation. — Scale bars = 10 µm.
**Type species: Pseudocercosporella ipomoeae Deighton.**

Colonies in vivo. Mycelium consisting of primary internal and secondary external hyphe, hyaline to pale brown, septate, branched, smooth; stromata lacking or weakly to well-developed, substomatal to intraepidermal. Conidiophores solitary to fasciculate, emerging through stoma or erumpent through the cuticle, arising from inner hyphae or from stromata, sometimes formed as lateral branches of superficial hyphae, or forming crustose to subglobose sporodochia; conidiophores rarely branched, straight and subcylindric to geniculate-sinuous, hyaline, occasionally faintly pigmented, reduced to conidiogenous cells, or septate. Conidiogenous cells integrated, terminal, mono- to polyblastic, sympodial; conidiogenous loci inconspicuous, unthickened, hyaline. Conidia formed singly, rarely in simple or branched chains, subcylindrical, filiform, somewhat obclavate, eusepaetate, 1–multi-sepate, hyaline, thin-walled, apex obtuse to subacute, base subtruncate, hilum unthickened, not darkened, nor refractive. Adapted from Braun (1995).

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**Fig. 9 Pseudocercosporella bakeri** (CPC 17570). a. Leaf spot on Ipomoea sp. with visible sporulation; b. colony on OA; c. conidiophores in vivo; d. conidiophores in vitro (arrows denote loci); e. conidia in vitro. — Scale bars = 10 µm.
additional collections of *P. bakeri* and *P. ipomoeae*, led Braun (1995) to the conclusion that they represented a single taxon. As shown in the present study, conidial dimensions vary considerably from host material to culture, and hence we support the conclusion of Braun (1995), and treat this as a single species, *P. bakeri*, for which an epitype is designated. This species clusters as a close sister to the ‘Dothistroma clade’ (Clade 7 in Crous et al. 2009c).

**DISCUSSION**

The present study treats three genera of fungi associated with SBFS of apple in Germany and Slovenia, namely *Devriesia*, *Microcyclospora* and *Microcyclosporella*. However, based on sequence similarity to related taxa associated with SBFS in public databases (Batzer et al. 2005, Díaz Arias et al. 2010), it is clear that these fungi have a much wider distribution with apples (see sequences from other hosts currently in GenBank; Fig. 2), and that many other species await description in the new genera introduced in this paper.

One genus newly linked to the SBFS complex on apples is *Devriesia*. As discussed previously, however, the genus *Devriesia* is paraplythic (Crous et al. 2007b, Koukol 2010), and further taxa need to be collected to provide more robust clades, and help delineate the morphological features needed to separate the non-thermotolerant genera from *Devriesia* s.str., which seems to be primarily adapted to burnt soil environments although its type species, *D. staurophora*, has been also isolated from dead leaves of *Pinus sylvestris* (Seifert et al. 2004). *Devriesia* s.str., however, is not commonly associated with leaf spots and blemishes on fruit surfaces as is the case in other lineages in *Devriesia* s.l.

The newly introduced genus *Microcyclosporella* shows clear similarities with genera such as *Pseudocercosporella* and *Ramulispora*. Deighton (1973) established the genus *Pseudocercosporella* for anamorphs of the *Mycosphaerella* complex that were *Cercospora*-like, but had unthickened and inconspicuous conidial scars. The four cercosporoid species known to be associated with eyespot disease of cereals (Nirenberg 1981, Robbertse et al. 1995, Lucas et al. 2000) were subsequently included in *Pseudocercosporella*, even though von Arx (1983) preferred to place them in *Ramulispora*. The genus *Ramulispora* is based on *R. sorghi*, which causes sooty stripe of sorghum, due to the abundant production of microscerotia on the leaf surface (Braun 1995). In a subsequent study, Crous et al. (2003) showed the eyespot fungi of wheat to represent a separate genus, *Helgardia*, which has apothecial teleomorphs in *Oculimacula* (Helotiales, Dermateaceae), while *Ramulispora* represents a genus in the *Mycosphaerellaceae* (Crous et al. 2009b, c), distinct from *Pseudocercosporella*. Interestingly enough, both *Ramulispora* and *Helgardia* exhibit microcyclic conidiation (Robbertse et al. 1995), as does the newly introduced *Microcyclosporella*. It appears that this character is of less taxonomic value at the generic level, and probably more ecologically relevant for pathogens that sporulate on superficial plant surfaces (wheat stems, sorghum leaves and apples), facilitating outward splash dispersal.

The genus *Pseudocercosporella* has recently been shown to include taxa that vary greatly in their conidial morphology, ranging from solitary conidiogenous loci, synnemata, sporodochia or fascicles. Furthermore, conidia were shown to include taxa that are transversely euseptate, but also with some muri or fascicles. Furthermore, conidia were shown to include taxa that vary greatly in their conidiomatal morphology, and their transversely septate Scolecosporous conidia with unthickened hila. Because these taxa have been shown to cluster apart from *Pseudocercosporella* s.str. (based on *P. vitis* in the present study, a new genus, *Microcyclospora*, has been introduced to accommodate the SBFS species. Morphologically, *Microcyclospora* can be distinguished from *Pseudocercosporella* s.str. in that conidiophores are never fasciculate, but are reduced to solitary conidigenous loci on hyphae, and that conidia occur in mucoid masses, which prominently undergo microcyclic conidiation.

Little is presently known about the ecology, epidemiology and host ranges of the SBFS fungi, and more sampling on other substrates or crops growing in the vicinity of apple orchards needs to occur to enable us to resolve these aspects. For *Schizothyrium pomi* for instance, up to 78 different plant hosts have been reported, of which many occurred in close vicinity of apple orchards in temperate North American climates (Baker et al. 1977). However, no modern approaches have been applied until now to test this hypothesis, while molecular analyses indicated that a collection of North American *Schizothyrium pomi* strains comprised more than 10 genotypes, of which four were elevated to species level (Batzer et al. 2008). These results clearly suggest that all published geographic and host distribution records of SBFS fungi will have to be treated with caution until they have been re-evaluated based on the new molecular approach currently employed to resolve species and generic boundaries.

Acknowledgements Parts of this study were supported by the Dutch Research Agency (ARRS) in the form of the Young Researcher grant 1000-06-31056 to J.F. We are grateful to A. van Iperen, M. Vermaas, M. Starink (CBS, Utrecht) and J. Wolter-Sadlers (INRES, Bonn) for providing technical assistance.

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