

A new *Ophiostoma* species from loblolly pine roots in the southeastern United States

James W. Zanzot¹, Z. Wilhelm de Beer², Lori G. Eckhardt¹ and Michael J. Wingfield²

(1) Forest Health Dynamics Laboratory, School of Forestry and Wildlife Sciences, Auburn University, Alabama, 36849, USA

(2) Department of Microbiology and Plant Pathology, Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, Pretoria, South Africa

Z. Wilhelm de Beer

Email: wilhelm.debeer@fabi.up.ac.za

Abstract

During the course of a survey of fungi in loblolly pine (*Pinus taeda*) roots in Georgia, USA, a species of *Ophiostoma* morphologically similar to *O. pluriannulatum*, was isolated. Morphological characteristics and DNA sequence comparisons were used to identify the fungus. The isolates produced perithecia with unusually long necks similar to those of *O. pluriannulatum* but they had few or no annuli. DNA sequences for the ribosomal internal transcribed spacer regions 1 and 2 were identical to those of *O. pluriannulatum*. Sequence data of the β -tubulin gene region revealed the absence of intron 4 and presence of intron 5, distinguishing the isolates from *O. pluriannulatum*, which has intron 4 but not intron 5. Phylogenetic analyses of the β -tubulin sequences showed that the isolates from loblolly pine roots grouped together in a lineage distinct from *O. multiannulatum* and *O. subannulatum*, both of which lack intron 4 and have intron 5. The fungus is consequently described as *O. sparsiannulatum* sp. nov., a novel taxon in the *O. pluriannulatum* complex.

Introduction

Species accommodated in the Ophiostomatales (Ascomycetes) are generally known as blue-stain fungi. This is due to the discoloration that most of these species cause on lumber (Seifert 1993) and sapwood of trees infested by scolytine bark beetles (Coleoptera: Scolytinae) or molytine weevils (Coleoptera: Molytinae), which act as their vectors (Malloch and Blackwell 1993). Some of these fungi are also pathogens of trees, e.g., *Ophiostoma ulmi* and *O. novo-ulmi* causing Dutch elm disease (Brasier 1991), while others are not or only mildly pathogenic to trees (Zhou *et al.* 2002).

Three teleomorph genera are recognized in the Ophiostomatales: *Grosmannia* that have *Leptographium* anamorphs, *Ceratocystiopsis* with *Hyalorhinocladiella* anamorphs, and *Ophiostoma* in which both *Sporothrix* and/or *Pesotum* anamorphs are found (Zipfel *et al.* 2006). Of these, *Ophiostoma* is the largest and most variable genus, and it includes some monophyletic lineages of species that share similar morphological characters. One such lineage has been referred to as the *O. pluriannulatum* complex (Zipfel *et al.* 2006).

The *O. pluriannulatum* complex is one of the lesser-known and studied groups in the Ophiostomatales, but DNA sequences from some recent studies have revealed several previously unknown species relationships in the complex. In a phylogeny based on the internal transcribed spacer (ITS) region sequences of the ribosomal RNA operon, Villarreal *et al.* (2005) showed that *O.*

multiannulatum, *O. subannulatum*, and *O. conicola* grouped close to *O. pluriannulatum*, the oldest species in the complex. Based on mating compatibility and ITS sequences, Thwaites *et al.* (2005) suggested that at least four more species exist in the complex. These included *O. perfectum* and an undescribed species referred to as *Ophiostoma* sp. E. They suggested that the other isolates from their study possibly represent *O. californicum* and *O. populinum* (Thwaites *et al.* 2005). Zipfel *et al.* (2006) added *O. carpenteri* to the complex based on ribosomal large subunit (LSU) and partial β -tubulin sequences, while Kamgan *et al.* (2008) described a new species, *O. longiconidiatum*, from native hardwood trees in South Africa. Most of these species are characterized by ascomata with long, fine necks that typically bear annuli (Davidson 1935; DeVay *et al.* 1968; Hedgcock 1906; Kamgan *et al.* 2008; Livingston and Davidson 1987).

During the course of investigations considering pine decline in the southeastern United States, isolates of an *Ophiostoma* species were collected from loblolly pine (*Pinus taeda* L.) roots in Fort Benning, Georgia. The morphology of these isolates superficially resembled *O. pluriannulatum* and related species. The aim of this study was to identify the new isolates to the species level using morphological criteria and DNA sequence comparisons.

Materials and Methods

Isolation of fungi

Loblolly pine roots were excavated at Fort Benning Military Reservation, Georgia, USA, using a two-lateral root sampling approach described by Otrosina *et al.* (1999) and modified by Eckhardt *et al.* (2007). Roots were surface disinfested in a solution of ethanol, commercial bleach, and distilled water (80:10:10, v:v:v), rinsed in tap water, and then plated on malt extract agar (MEA, 20 g malt extract, 15 g agar/L) and MEA amended with cycloheximide and streptomycin (CSMA) (Zhou *et al.* 2004a). Ophiostomatoid fungi were recognized by their characteristic perithecia or conidiophores and transferred to fresh CSMA or MEA. All isolates are maintained in the culture collection (CMW) of FABI, University of Pretoria, South Africa. Type specimens were deposited in National Collection of Fungi (PREM), Pretoria, South Africa, and ex-type cultures in the Centraalbureau voor Schimmelcultures (CBS), Utrecht, The Netherlands.

Morphology

To assess characters of the mycelium, ascomata, and conidia, isolates were grown on oatmeal agar (OA; 30 g Jungle brand oats/L with 15 g/L agar), 2% MEA or pine twig agar (PTA; water agar 15 g/L with twice-autoclaved Mexican weeping pine [*Pinus patula* Schltdl. & Cham.] twigs embedded into the agar). Fungal characteristics were measured using a Zeiss Axioskop 2 microscope with phase contrast and differential interference contrast (DIC) microscopy, and data were collected using the software package AxioVision Release 4.4.1.0. (Carl Zeiss Microimaging, München, Germany). Fifty measurements were made of each structure, and maxima, minima, and means determined.

Agar plugs (5 mm diameter) taken from actively growing cultures were transferred to fresh MEA plates, incubated at temperatures ranging from 5 to 35°C at five-degree intervals using three replicate plates for each temperature. Radial growth was measured every 2 days until the colonies reached the edges of the plates.

Single ascospore isolates were obtained by streaking spore droplets on MEA plates. Germinating ascospores were transferred to new MEA plates. These were crossed in all possible combinations (Zhou *et al.* 2004b) to test whether the fungus was homo- or heterothallic.

DNA extraction, PCR and sequencing

Genomic DNA was prepared for PCR from fresh mycelium taken from cultures on MEA using PrepMan Ultra Reagent (Applied Biosystems, Foster City, California, USA), following the protocol described by Aghayeva *et al.* (2004). The ITS regions were amplified using the primers ITS1F (Gardes and Bruns 1993) and ITS4 (White *et al.* 1990) and the β -tubulin gene region using the primers T10 (O'Donnell and Cigelnik 1997) together with Bt2b (Glass and Donaldson 1995).

Amplifications were performed in an Eppendorf MasterCycler Personal thermal cycler (Eppendorf, Hamburg, Germany) under the following conditions: an initial denaturation at 95°C for 2 min, followed by 40 cycles of 95°C for 30 sec, 50°C annealing for 30 sec, and 72°C extension for 60 sec, with a final extension period of 8 min at 72°C. Reaction mixtures, 50 μ L total volume, consisted of 5 μ L 10 \times PCR reaction buffer (JMR Holdings, London, UK), 2.5 mM MgCl₂, 10 mM dNTPs, 10 μ M of each primer, 2.5 U Taq polymerase (SuperTherm; JMR Holdings) and 4 μ L of genomic DNA. Amplification success was assessed on a 1% (w/v) agarose gel stained with ethidium bromide and visualized under UV light. PCR products were purified with the High Pure PCR Purification Kit (Roche Pharmaceuticals, Basel, Switzerland) according to the manufacturer's protocol. Amplicons were then cycle sequenced using the Big Dye™ Terminator v. 3.0 cycle sequencing premix kit (Applied Biosystems) with the primers noted above, and analyzed on an ABI PRISM™ 377 or ABI PRISM™ 3100 Genetic Analyzer (Applied Biosystems).

Electropherograms of the resolved sequences were visualized, and contigs of forward and reverse sequences compiled using MEGA 4.0.1 (Tamura *et al.* 2007). Sequences were subjected to BLAST searches in NCBI GenBank to determine similarity to published sequences. Appropriate sequences were downloaded and datasets compiled in MEGA 4.0.1. Alignments were done using the E-INS-I strategy in the online version of MAFFT 6 (Kato and Toh 2008) (<http://align.bmr.kyushu-u.ac.jp/mafft/online/server/>). For the β -tubulin gene region, manual adjustments of the alignment were performed in MEGA 4.0.1 using the translation option to ensure appropriate alignment of exons based on amino acid sequences. All sequences generated in this study were deposited in NCBI GenBank.

Phylogenetic analyses

Aligned sequences of the ITS and β -tubulin gene regions were subjected to phylogenetic modeling using maximum parsimony (MP), maximum likelihood (ML), and neighbor-joining (NJ) in PAUP 4.0 b10 (Sinauer, Sunderland, Massachusetts, USA), and Bayesian analyses using MrBayes 3.1.2 (Ronquist and Huelsenbeck 2003). The most appropriate substitution models for both gene regions were selected using the Akaike information criterion (AIC) in ModelTest 3.7 (Posada and Crandall 1998) for ML and NJ analyses, and MrModelTest 2 (Nylander 2004) for Bayesian analyses. For MP, ML, and NJ, 1,000 bootstrap replicates were performed to test branch stability. Bayesian analyses consisted of four independent runs of 10 million generations each using duplicate Monte Carlo Markov chain searches with four chains. Trees were saved every 100 generations. Burnin was determined for each dataset using Tracer 1.4 (<http://evolve.zoo.ox.ac.uk/software.html>).

Results

Isolation of fungi

Three isolates of an unidentified *Ophiostoma* sp. were recovered from loblolly pine roots. These isolates were cycloheximide-tolerant, and were found in association with other cycloheximide-tolerant taxa including *Leptographium* spp. and *Penicillium* spp.

Morphology

Isolates produced abundant aerial hyphae bearing dry, micronematous conidia typical of the anamorph genus *Sporothrix*. Long-necked, darkly pigmented ascomata formed abundantly in culture within 8 days, but only in crosses between single conidial isolates. The ascomata were variable in size, with large and small ascomata producing droplets with viable ascospores. Colonies tended to darken with age. Morphological characteristics were consistent with those of the *O. pluriannulatum* complex (Table 1), although annuli on the ascomatal necks were few in number or lacking. The ascospores were slightly smaller than those reported for the other species most similar to the isolates from loblolly pine roots (Table 1).

Table 1. Comparison of characters of *Ophiostoma sparsiannulatum* with related and morphologically similar species.

Character	Subcharacter	<i>O. sparsiannulatum</i> sp. nov.	<i>O. pluriannulatum</i>	<i>O. multiannulatum</i>	<i>O. californicum</i>	<i>O. subannulatum</i>
Perithecia	Neck length (mm)	0.8–1.4(–8)	(0.9–)1.5(–2.0)	4.0–8.5	0.6–1.1	0.2–1.2(up to 2.2)
	Annuli	0–2(–4)	3–8	6–9	0 to several	1–2(–4)
	Color of base	Dark brown to black	Black	Black	Hyaline	Black
	Diameter of base (µm)	(54)104–166(201)	(90)120(200)	170–275	130–175	75–200
	Ornamentations of base	Brownish hyphae up to 200µm long	Unornamented or with a few septate, pale brown hyphae	Surface roughened with globose cells	Smooth	Unornamented or with a few septate, brownish hyphal hairs
Ascospores	Shape	Allantoid	Reniform	Reniform	Curved	Allantoid to broadly lunate
	Length (µm)	(2.2–)3.0(–4.0)	(4.0–)4.5(–5.0)	2.8–3.8	3.0–4.5	3.5–5.0(–6.0)
	Width (µm)	(0.7–)1.0(–1.6)	(1.5–)1.5(–1.7)	1.2–1.4	1.0–1.3	1.0–2.5
Anamorph		<i>Sporothrix</i>	<i>Sporothrix</i>	<i>Sporothrix</i>	<i>Sporothrix</i>	<i>Sporothrix</i> (with sticky spores)
Conidia	Length (µm)	2.7–9.2	4.6–18.0	6.0–25.0	3.0–18.0	2.0–6.0
	Width (µm)	0.8–2.3	1.5–3.0	2.0–3.2	1.2–1.6	1.0–4.0
Host	Secondary conidia	Present	Present	Not present	Present	Present
Host		<i>Pinus taeda</i>	<i>Quercus</i> spp., <i>Pinus</i> spp.	<i>Pinus</i> spp.	<i>Prunus domestica</i>	<i>Abies</i> spp.
Distribution		Georgia, USA	Southeastern USA, New Zealand	Southeastern USA	California, USA	Idaho, USA
β-tubulin introns ^a		–/5	4/–	–/5	No β-tubulin sequences	–/5
Morphological descriptions		This study	Hedgecock 1906; Upadhyay 1981	Davidson 1935	Devay et al. 1968; Upadhyay 1981	Livingston and Davidson 1987

PCR and sequencing

Amplicons of the ITS regions yielded 618 bp of readable sequence data with NCBI GenBank accession numbers FJ906816–FJ906818. These sequences were identical to those for *O. pluriannulatum* published in GenBank, and similar to several other *Ophiostoma* spp. Amplicons of the β-tubulin gene yielded 575 bp of readable sequence data (FJ907176–FJ907178), which shared 93% homology with *O. subannulatum*, the closest match in GenBank. The partial β-tubulin sequences of these isolates included 14 bp of exon 3, no intron 3, exon 4 (27 bp), no intron 4, exon 5 (42 bp), intron 5 (49 bp), and 443 bp of the 5' end of exon 6.

Phylogenetic analyses

For the ML and NJ analyses of the ITS dataset, the TrN+G substitution model was selected. For the Bayesian analysis, GTR+I+G was selected. The aligned ITS dataset consisted of 742 characters and confirmed the placement of the isolates in the genus *Ophiostoma* and in the *O. pluriannulatum* complex (Fig. 1). ITS sequences for the isolates from *P. taeda* roots were identical to those of *O. pluriannulatum sensu stricto*, but differed in nine bases from *O. multiannulatum* and six bases from *O. subannulatum*. The *O. californicum* and *Ophiostoma* sp. E isolates of Thwaites *et al.* (2005) formed well-supported clades within the larger *O. pluriannulatum* clade, distinct from the isolates from loblolly pine roots.

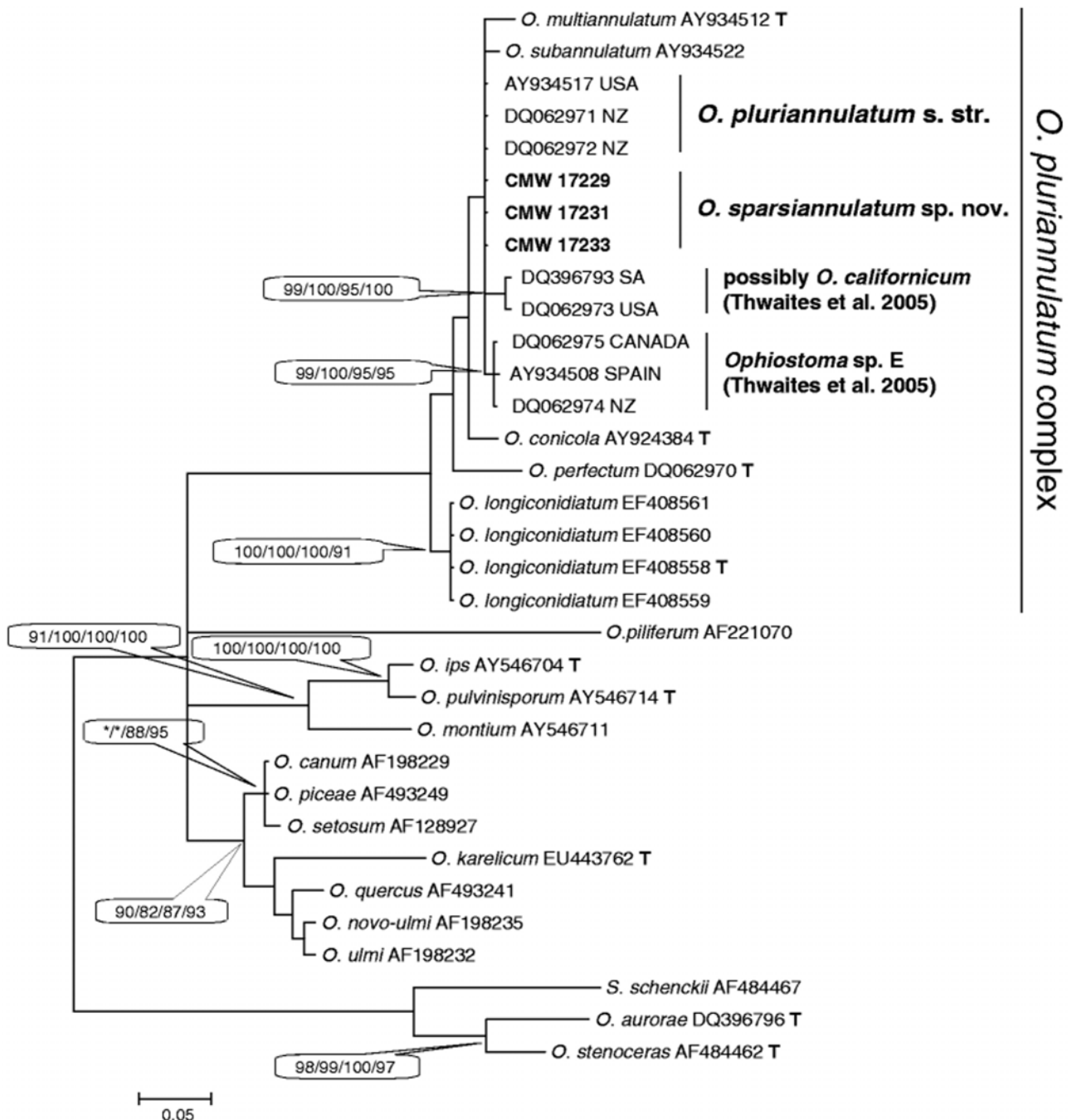


Fig. 1. Bayesian tree of ITS sequences of selected *Ophiostoma* and *Sporothrix* isolates, representing all the major lineages in *Ophiostoma*. Sequences obtained from the present study are printed in bold type. Accession numbers are presented for all sequences obtained from GenBank. Branch support is indicated by bootstrap analyses (1,000 replicates) for MP, ML, and NJ analyses, and posterior probabilities for Bayesian analysis, respectively. *Bootstrap support <70%, T ex-type isolates.

The aligned β -tubulin data used for phylogenetic analyses consisted of 354 characters, including exons 4, 5 and 6, interspersed by introns 4 and 5. In the loblolly pine root isolates, intron 4 was lacking and intron 5 was present, and this was the same for *O. multiannulatum* and *O. subannulatum*. *Ophiostoma pluriannulatum*, *O. carpenteri*, and *O. piliferum* had intron 4 and lacked intron 5, thus separate phylogenetic analyses were conducted for datasets including (Fig. 2a) and excluding (Fig. 2b) species with intron arrangements different from the isolates from *P. taeda* roots. For ML and NJ analyses, the TrN+I model was selected for the β -tubulin dataset including *O. pluriannulatum*, *O. carpenteri*, and *O. piliferum*, and for the β -tubulin dataset excluding these three taxa, the HKY+I model was selected. For the Bayesian analysis, GTR+I+G was selected for the inclusive dataset, and HKY+I for the exclusive dataset.

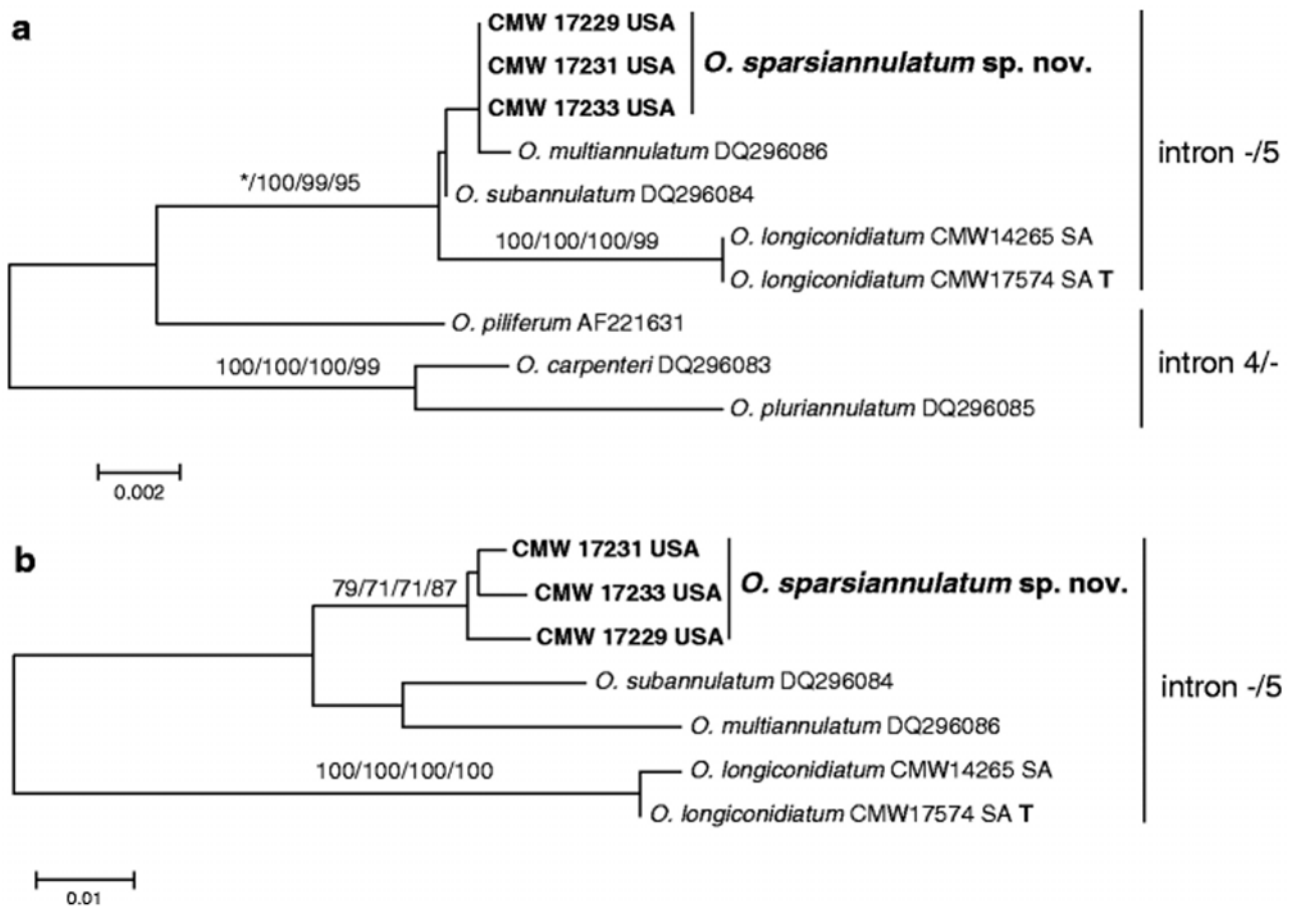


Fig. 2. Bayesian trees of β -tubulin sequences of species from the *O. pluriannulatum* complex, including exons 4, 5, and 6, as well as the introns as specified. Accession numbers are presented for all sequences obtained from GenBank. Branch support is indicated by bootstrap analyses (1,000 replicates) for MP, ML, and NJ analyses, and posterior probabilities for Bayesian analysis, respectively. *Bootstrap support <70%, T ex-type isolates. **a** Dataset including species that have intron 4 but not intron 5, as well as those lacking intron 4, but with intron 5. **b** Dataset including only species without intron 4 and with intron 5

Phylogenetic trees generated from the β -tubulin sequence data showed that the isolates under investigation were distinct from *O. pluriannulatum sensu stricto* (Fig. 2a). The pattern of exons and introns was similar to that of *O. multiannulatum* and *O. subannulatum* (Fig. 2a), but the sequences were distinct from the latter two species, forming a lineage with strong statistical support in the analyses that included only species with intron 4 (Fig. 2b).

Taxonomy

The *Ophiostoma* species from loblolly pine roots is morphologically similar to *O. pluriannulatum* and related species (Table 1). However, based on DNA and morphological comparisons, the new isolates represent a novel species, described here as follows:

Ophiostoma sparsiannulatum Zanzot, Z.W. de Beer and M.J. Wingf., sp. nov. MycoBank MB 515290. Figs. 3 and 4.

Etymology: The name refers to the fact that annuli are produced sparsely on the ascomatal necks of this fungus.

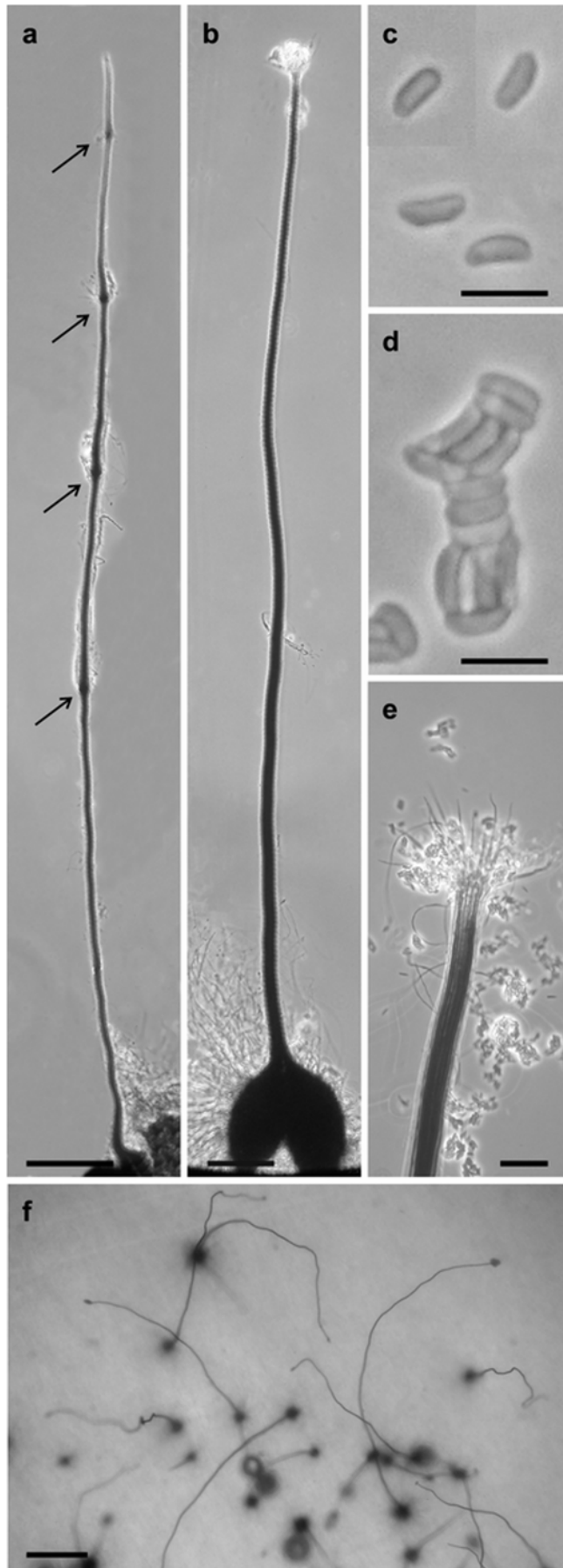


Fig. 3. Teleomorph of *Ophiostoma sparsiannulatum* (ex-holotype, CBS 122815). **a** Neck of mature ascoma with annuli (arrows) present (bar 100 μ m). **b** Mature ascoma (bar 100 μ m). **c, d** Ascospores (bars 5 μ m). **e** Ostiolar hyphae surrounding ostiole at apex of ascomatal neck (bar 20 μ m). **f** Ascomata submerged in pine twig agar (bar 1 mm).

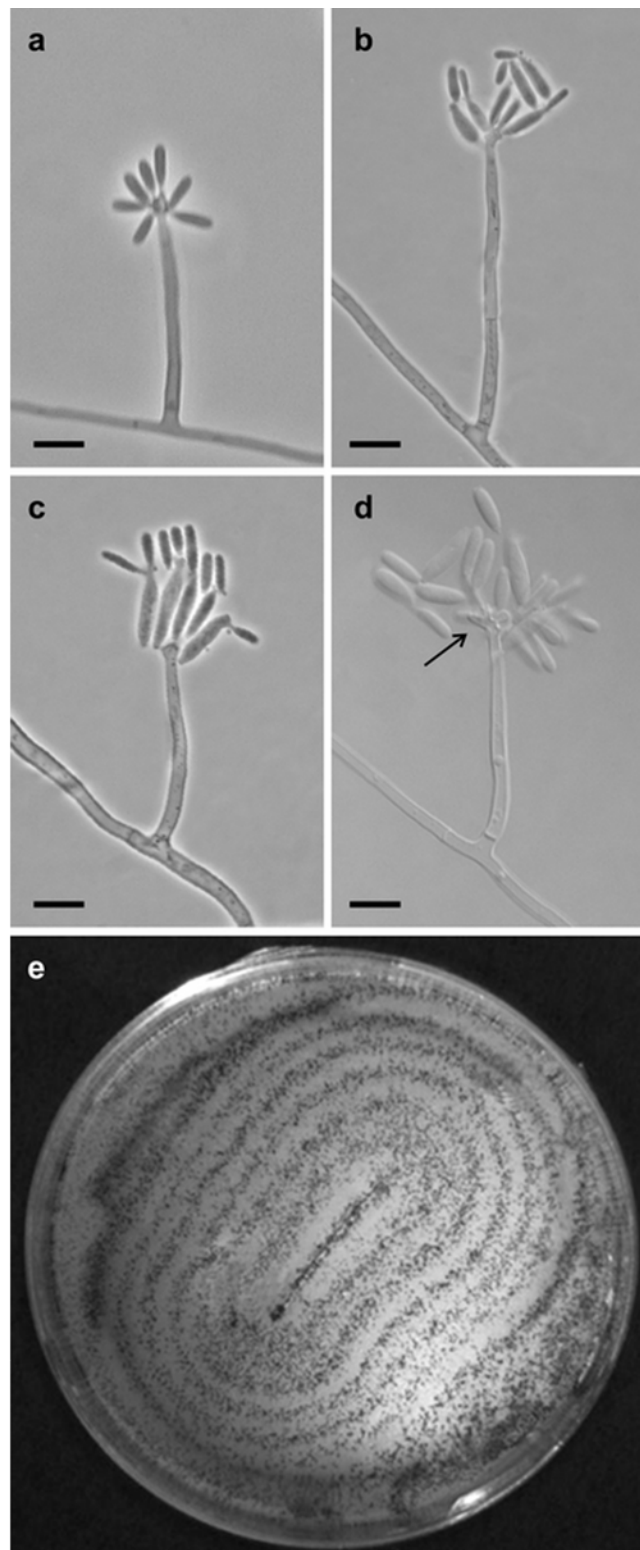


Fig. 4. *Sporothrix* anamorph of *Ophiostoma sparsiannulatum* (ex-holotype, CBS 122815). **a–c** Different stages of conidiophores with conidia, showing the development of secondary conidia (phase contrast, bars 5 μm). **d** Conidiogenous cell with conidia formed on denticles (DIC, bar 5 μm). **e** Culture on oatmeal agar, showing concentric rings of ascomata.

Description: Bases ascomatum atrobrunneae vel nigrae, globosae, (54–)104–166(–201) μm diametro. Colla ascomatum atrobrunnea vel nigra, (500–)800–1,400(–1,850) μm longa, annulis 0–4, apice (7–)10–17(–21) μm , basi (13–)27–48(–65) μm diametro. Ascomata nonnulla in agaro immersa colla valde longiora usque ad 8,600 μm habent. Hyphae ostiolares divergentes, hyalinae (18–)32–62(–80) μm longa. Ascosporae hyalinae, allantoideae, (2.2–)2.8–3.4(–4.0) \times (0.7–)0.9–

1.1(–1.6) μm , in massa pallide flava mucosa in apice cumulantes. Anamorpha *Sporothrix* inter ascomata in massis albis, floccosis, abundanter, crescens. Conidiophorae (9–)13–52(–150) μm longae, cellulae conidiogenae denticulis (0.7–)0.8–1.3(–1.6) μm longis. Conidia plerumque ovoidea vel obovoidea, (2.7–)3.8–6.7(–9.2) \times (0.8–)1–1.8(–2.3) μm , saepe conidia secundaria emittentia.

Colony hyaline at first, darkening with age and with copious white floccose aerial growth of the anamorph, typically forming concentric rings (Fig. 4e). Mycelium aerial and submerged. Optimal temperature for growth 25°C, no growth at 35°C. Hyphae smooth, not constricted at septa. Ascomata typically produced within 8 days, and only in crosses between single spore isolates, suggesting heterothallism. Ascomatal bases dark brown to black, globose, bases (54–)104–166(–201) μm in diameter (Fig. 3b), brownish ornamental hyphae up to 200 μm long. Ascomatal necks dark brown to black, (500–)800–1,400(–1,850) μm long with annuli absent or occasionally 1–4 present (Fig. 3a). Some submerged perithecia observed on PTA (Fig. 3f) have much longer necks, as long as 8,600 μm . Diameter of the necks (7–)10–17(–21) μm at the apex and (13–)27–48–(65) μm at the base. Ostiolar hyphae divergent, hyaline, (18–)32–62(–80) μm long (Fig. 3e). Asci not observed. Ascospores accumulating in a clear white mucilaginous mass at the ascomatal apex, turning light yellow with age, hyaline, allantoid, (2.2–)2.8–3.4(–4.0) \times (0.7–)0.9–1.1(–1.6) μm (Fig. 3c, d).

Sporothrix anamorph: growing abundantly amongst the perithecia, in white fluffy masses. Conidiophores of variable length, (9–)13–52(–150) μm (Fig. 4a–d), conidiogenous cells with denticles (0.7–)0.8–1.3(–1.6) μm long (Fig. 4d). Conidia typically ovoid-obovoid with a basal abscission collar evident, (2.7–)3.8–6.7(–9.2) \times (0.8–)1–1.8(–2.3) μm , often giving rise to secondary conidia (Fig. 4b–d).

Specimens examined: USA, Georgia, Fort Benning, from roots of *Pinus taeda*, July 2004, L.G. Eckhardt. Holotype PREM 59847, ex-holotype culture CBS 122815=CMW 17231, Paratype PREM 59848, ex-paratype culture CBS 122816=CMW 17233. Additional culture CMW 17229.

Notes: Morphological differences between *O. sparsiannulatum* and similar species are listed in Table 1. *Ophiostoma pluriannulatum* differs from *O. sparsiannulatum* in having longer ascospores and longer conidia, while *O. multiannulatum* has longer perithecial necks, greater numbers of annuli, longer conidia, no secondary conidia, and characteristic globose cells on the perithecial bases. *Ophiostoma californicum* has hyaline bases that are not ornamented as opposed to the dark bases with ornamental hyphae in *O. sparsiannulatum*. Of all these species, *O. subannulatum* is most similar to *O. sparsiannulatum*, but its conidia are generally shorter and rounder than those of the latter species. The conidia of *O. subannulatum* are also produced in slimy masses (Livingston and Davidson 1987), while those of *O. sparsiannulatum*, *O. pluriannulatum*, and *O. multiannulatum* are borne in dry, floccose clusters.

Discussion

This investigation revealed a new species of *Ophiostoma* isolated from roots of loblolly pine that is morphologically similar and closely related to *O. pluriannulatum*. Recognition of the new species initially rested strongly on DNA sequence comparisons with *O. pluriannulatum* and related species. Yet having recognized that it represented a novel taxon, it is also possible to distinguish *O. sparsiannulatum* from related species based on morphological characteristics.

DNA sequence comparisons, especially the ITS regions, are often used to address species level questions in fungi, including the Ophiostomatales (De Beer *et al.* 2003; Harrington *et al.* 2001). However, in the present study, ITS sequences did not distinguish satisfactorily between *O. sparsiannulatum* and other species in the complex. This is not unique as there are other cases where

ITS sequences have failed to resolve closely related species within *Ophiostoma* (Chung *et al.* 2006; Harrington *et al.* 2001). Partial β -tubulin sequences, including intron data, are now routinely used to improve resolution when comparing closely related species (Gorton *et al.* 2004; Grobbelaar *et al.* 2009). The presence or absence of introns within the β -tubulin gene region have been shown to be constant within major lineages of the Ophiostomatales, and can be used as a tool to distinguish between these lineages (Zipfel *et al.* 2006). However, results from the present study showed for the first time that two closely related *Ophiostoma* species (*O. pluriannulatum* and *O. subannulatum*) can have identical ITS sequences but a different intron/exon composition in the β -tubulin gene. This emphasizes the fact that extreme care should be taken when aligning β -tubulin sequences, always ensuring that exon data are aligned appropriately based on their amino acid sequences. The risk here lies in the fact that a small exon might not be recognized by the alignment program if it is not designated. This is especially true when an alignment is enforced onto longer introns on either side of the exon. The exon sequence would thus be aligned with intron sequences and resulting phylogenies would not accurately reflect the evolution of the gene region.

An important morphological characteristic in most members of the *O. pluriannulatum* complex is that they have long perithecial necks with annuli. These annuli are likely vestiges of ostiolar hyphae through which the necks have proliferated (DeVay *et al.* 1968; Thwaites *et al.* 2005). Both *O. pluriannulatum* and *O. multiannulatum* have perithecia with several annuli. While *O. sparsiannulatum* has perithecial necks similar in length to these species, it has no or very few annuli. In this regard, it is most similar to *O. subannulatum*.

The importance of annuli as a diagnostic character for *Ophiostoma* spp. has been debated, with their importance discounted (De Hoog 1974) or emphasized (Upadhyay 1981). Phylogenetic inference based on DNA sequences support the *O. pluriannulatum* complex as a monophyletic clade, now accommodating seven *Ophiostoma* spp. with annulate perithecial necks: *O. sparsiannulatum*, *O. pluriannulatum*, *O. multiannulatum*, *O. californicum*, *O. subannulatum*, *O. longiconidiatum* (Thwaites *et al.* 2005; Zipfel *et al.* 2006; Kamgan *et al.* 2008), and an as yet undescribed species, *Ophiostoma* sp. E (Thwaites *et al.* 2005). Species in the complex that do not form long necks with annuli are *O. carpenteri* (Hausner *et al.* 2003), *O. conicola* (Marmolejo and Butin 1990), and *O. perfectum* (Davidson 1958). The position of *O. perfectum* is uncertain, since ITS sequences of the ex-type isolate (CBS 636.66) produced by Thwaites *et al.* (2005) and Villarreal *et al.* (2005) are very different from each other. The sequence produced by Thwaites *et al.* (2005) is included in our analyses (Fig. 1), but those of Villarreal *et al.* (2005) correspond to *O. floccosum*, which groups outside the *O. pluriannulatum* complex and closer to *O. piceae* (Zipfel *et al.* 2006). Furthermore, De Hoog (1974) described a synnematous synanamorph from the ex-type isolate of *O. perfectum*, while Davidson (1958) only mentioned a *Sporothrix* anamorph in his original description. It is thus possible that the ex-type isolate represents a mixed culture. The relatedness of *O. perfectum*, *O. carpenteri*, and *O. conicola* to the *O. pluriannulatum* complex should be re-evaluated with critical study of the type material, and where possible additional isolates of each species. Such a study would reveal whether annuli on elongated perithecial necks should be considered a unifying characteristic of the *O. pluriannulatum* complex.

The first species to be described in the *O. pluriannulatum* complex was *O. pluriannulatum sensu stricto*, which was isolated from stained northern red oak (*Quercus rubra* L.) sapwood (Hedgcock 1906). This species has been recorded by far the most often of all species in the complex, and also from many other hardwood genera and species in North America (Appel *et al.* 1990; Davidson 1935; Griffin 1968), Europe (Gregor 1932) and Japan (Aoshima 1965). The fungus has, furthermore, been described from conifers such as pine, spruce, and fir from North and Central America (Davidson 1935; Zhou *et al.* 2004b), Europe (Mathiesen-Käärik 1953; Romón *et al.* 2007), Japan (Aoshima 1965), New Zealand (Thwaites *et al.* 2005), and South Africa (Zhou *et al.* 2006). Identification of *O. pluriannulatum* in most of these reports was based on morphology, and only in

more recent cases on ITS sequences (Thwaites *et al.* 2005; Romón *et al.* 2007; Villarreal *et al.* 2005; Zhou *et al.* 2006). Results of the present study reinforce the fact that additional sequence data such as those for the β -tubulin gene region are required to delineate species in the complex. Clearly, overlapping morphological features in many of these species have resulted in isolates reported in the older literature as *O. pluriannulatum*, while they could represent other species in the complex. For example, the fungus reported by Verrall (1939, 1941) from loblolly pine in Georgia, and for which cultures are no longer available, could have been the same as *O. sparsiannulatum*, given that it was isolated from the same host species in the same region.

Ophiostoma pluriannulatum has been isolated from several wood- and bark-infesting insects and their galleries (Appel *et al.* 1990; Mathiesen-Käärik 1953; Romón *et al.* 2007; Verrall 1941; Zhou *et al.* 2004b, 2006). Again, the identifications of the fungus were based on morphology and only in the recent cases on ITS sequence. It is, therefore, impossible to draw meaningful conclusions relating to the insect–fungus interactions in this complex, apart from the fact that most species appear to be associated with bark beetles and nitidulids. Any conclusions regarding the specificity of these insect–fungus–host interactions will only be possible when identifications of the fungal species have been resolved based on sufficiently variable gene regions.

Ophiostoma sparsiannulatum is known only from the roots of loblolly pine damaged by various root-infesting insects. Given that these insects specialize on *Pinus* spp., it seems likely that the fungus will share this habit. While *O. sparsiannulatum* has been recovered from the living roots of declining loblolly pine trees, its recovery was infrequent. This suggests that it is not an important contributing factor in southern pine decline (Eckhardt *et al.* 2007; Menard 2007). Virtually no pathogenicity data are available for the members of the *O. pluriannulatum* complex, but these fungi are not considered to be pathogens (Davidson 1935; Hedgcock 1906). Nonetheless, further studies should include root inoculations with *O. sparsiannulatum* to evaluate its capacity to infect roots, as well as consideration of how it is transmitted and whether it may infect other hosts.

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