

***Pseudocercospora mapelanensis* sp. nov., associated with a fruit and leaf disease of
Barringtonia racemosa in South Africa**

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ABSTRACT

Barringtonia racemosa (Lecythidaceae) is a widely distributed mangrove associate in coastal areas of Africa, Asia and Australia. During routine disease surveys along the east coast of the KwaZulu-Natal Province in South Africa, *B. racemosa* trees were observed showing symptoms of leaf infection and necrotic lesions on fruits. A previously undescribed species of *Pseudocercospora* was commonly associated with these symptoms.

This fungus is described as *Pseudocercospora mapelanensis* sp. nov., based on multi-gene sequence analyses of the ACT, ITS, LSU and TEF genome regions, as well as morphological characteristics.

Key Words: *Capnodiales, Lecythidaceae, Mangroves, Mycosphaerellaceae*

INTRODUCTION

Mangroves, including true mangroves and mangrove associates (Tomlinson 1986; Ellison and Farnsworth 2001; Kathiresan and Bingham 2001; Hogarth 2007; Wang et al. 2010), are adapted to survive in saline-rich water and anoxic soils. They are found along tropical and subtropical coastlines where they provide many environmental and economic benefits (Spalding et al. 2010). Mangroves are, however, frequently threatened by anthropogenic activities and/or environmental factors (Spalding 2010). Despite their importance there have been relatively few studies to consider the impact of microbial diseases on these trees (Osorio et al. 2014). Diseases that have been reported from mangroves include branch cankers, leaf spots and die-back of trees (Barnard and Freeman 1982; Teas et al. 1982; Wier et al. 2000; Tattar and Scott 2004; Osorio et al. 2014).

Barringtonia racemosa (L.) Spreng. (powder-puff tree, fresh water mangrove) is a mangrove associate that belongs to the family Lecythidaceae, found along canals and rivers of coastal areas of Africa, Asia and Australia (Chantaranothai 1995; USDA-ARS 2014). *Barringtonia racemosa* trees are valuable as a source of medicinal products, timber, food and tannin (Lim 2012). Although several fungal species have been reported from *Barringtonia* species globally, nothing is known regarding fungal pathogens of *B. racemosa* in South Africa. During a survey of the health of mangrove species in this country, a fruit, floral and leaf disease was observed on *B. racemosa* trees in the Zululand

region. The aim of this study was to describe the symptoms of the disease and to identify its causal agent.

Materials and Methods

Disease observations and sample collection

In February 2011, a disease affecting the flowers, fruits and leaves of *Barringtonia racemosa* was observed in the Mapelane area of the Kwazulu-Natal Province in South Africa. In order to identify the fungus most closely associated with the disease symptoms, infected fruits, floral sepals and leaves were collected from several sites, placed in paper bags and transported to the laboratory for further study. During 2012 and 2013, additional areas where *B. racemosa* is known to occur were also surveyed for the occurrence of the disease, and additional samples were collected for analyses.

Cultures

A single fungus was found sporulating profusely on infected green leaf and fruit tissue. Fungal structures were transferred directly from plant material, mounted on microscope slides and examined under a Zeiss Axioskop microscope (Carl Zeiss, Germany). Images were obtained by using an Axiocam digital camera connected to the microscope. To isolate this fungus, stromatic tissue was removed from symptomatic plant material and transferred to 500 µl autoclaved Sabax water. Of this, 100 µl were spread onto 2% malt extract agar (MEA, Biolab Malt Extract, 17g Biolab Agar amended with 0.4g of streptomycin to suppress bacterial growth, in 1L distilled water) and incubated overnight. Germinating conidia were transferred to fresh 2% MEA. Pure cultures obtained from single spore isolates were deposited in the culture collection (CMW) of the Tree Protection Co-operative Programme (TPCP) at FABI, University of Pretoria. Duplicate cultures were

deposited in the CBS-KNAW Fungal Biodiversity Centre's culture collection (Centraalbureau voor Schimmelcultures) in Utrecht, The Netherlands.

Pathogen Identification

DNA Extraction, PCR Amplification and Sequencing

To extract genomic DNA, mycelium from colonies of three isolates, no older than four weeks of age, were placed into 1.5mL sterile Eppendorf tubes and freeze dried. Isolates for identification were selected based on plant part and collection site, so as to represent all symptom types and regions, as well as multiple trees. DNA was extracted using the phenol-Chloroform method described by Raeder and Broda (1985). After DNA was obtained, a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Rockland, DE) was used to measure DNA concentrations, and to calculate the volumes to be used in PCR reactions.

Four gene regions were sequenced for all isolates. A portion of the mitochondrial large subunit (LSU) was amplified using primers LR5 (Vilgalys and Hester 1990) and LR0R (Moncalvo et al. 1993). The internally transcribed spacer (ITS) regions (ITS1, ITS2) were amplified using the primers ITS1 and ITS4 (White et al. 1990). A portion of the translation elongation factor 1- α (TEF) was amplified using the primers EF1F and EF2R (Jacobs et al. 2004). A portion of the actin (ACT) gene was amplified using the primers ACT-512F and ACT-783R (Carbone and Kohn 1999).

For all gene regions, each reaction contained 2.5 μ L of PCR buffer, 2 μ L dNTP, 1 μ L of each primer, 0.3 μ L of fast *Taq* polymerase and 3 μ L of DNA (60ng/ μ l). Sterile Sabax water was added to obtain a final volume of 25 μ L for each reaction. Reactions were run using the following thermal cycling conditions: initial denaturation at 94°C for four minutes

followed by a denaturation step of ten cycles consisting of 94°C for 20 seconds, annealing at 55°C (ITS and TEF), 48°C (LSU) and 61°C (ACT) for 48 seconds, and elongation at 72°C for 45 seconds, followed by a further 25 cycles of 94°C for 20 seconds, followed by an annealing step with temperatures as previously indicated for each gene region (55°C, 48°C, 61°C), for 40 seconds with a time increase of five seconds per cycle, and then 45 seconds at 72°C. This was concluded by a final step of 72°C for 10 minutes. An aliquot of five µl of each PCR product was separated by gel electrophoresis at 90V for 20 min in a 2% agarose gel in 5% TAE Buffer (40 mM Tris, 40 mM acetate, 2 mM EDTA, pH 8.0) to evaluate the success of the reactions using GelRed™ nucleic acid gel stain (Biotium, Hayward, CA, USA). PCR products were cleaned using Sephadex G-50 columns, following the instructions provided by the manufacturers (Sigma Aldrich, Sweden).

Purified PCR products were used as template DNA for sequencing reactions, using a Big-Dye terminator cycle sequencing kit (Perkin-Elmer Applied Biosystems, Foster City, California, USA) and the same primers and annealing temperatures that were used in the PCR. The final products were also cleaned in Sephadex G-50 columns. Sequencing was carried out on an ABI PRISM 3100 DNA sequencer (Applied Biosystems).

Morphology

Colony morphology and microscopic features of the isolated fungus were examined directly from plant material and from cultures growing on 2% MEA. Mounts of the fungus were prepared in 85% lactic acid on microscope slides for detailed observations of structures such as stromata, conidiophores and conidia. Where structures were dry, these were mounted in 2% KOH. Characters such as size of conidiophores and conidiogenous cells as well as pigmentation of the conidia were used for description of the species based on morphology. Fifty measurements were made for each relevant morphological

characteristic and the standard deviation (S.D.) was calculated for measurements of the stromata, conidia and conidiophores. Minimum and maximum dimensions are given in parentheses.

Colony colors (surface and reverse) were assessed after 30 days on MEA, using the color chart of Rayner (1970). Growth of cultures was assessed by measuring single conidial cultures after 15, 30 and 60 days maintained at a temperature of 25°C. Culture characteristics were determined from cultures grown under natural day/night conditions.

Phylogenetic analyses

Sequences of the isolates obtained from *B. racemosa* were assembled using CLC Main Workbench 6.7.1 (<http://www.clcbio.com/genomics/>), and compared with published sequences using a Blast search in the GenBank (<http://blast.ncbi.nlm.nih.gov/blast.cgi>) data base. DNA sequences for closely related species, previously published, were retrieved from GenBank and combined into data sets with the *B. racemosa* sequences (Table 1). The data matrices were aligned online using MAFFT (<http://align.bmr.kyushuu.ac.jp/mafft/online/server/>) version 6 (Katoh et al. 2005) and edited manually for alignment errors with MEGA version 6 (Tamura et al. 2013).

Phylogenetic analyses of sequence data for Maximum Parsimony (MP) were performed with the software package PAUP* 4.0b10 (Swofford 2003). Maximum parsimony (MP) genealogies for single genes were constructed with the heuristic search option (1000 random taxa additions, tree bisection and reconstruction or TBR in PAUP). Uninformative characters in each data set were excluded from the analyses, and the consistency index (CI), homoplasy index (HI), rescaled consistency index (RC), retention index (RI) and tree length (TL) were determined for the resulting trees. Partition homogeneity tests (PHT) were conducted to determine the congruence of trees obtained from the different gene

regions with PAUP* 4.0b10 (Swofford 2003) to define whether data from the different gene regions could be combined.

Phylogenetic analyses of sequence data for Maximum Likelihood (ML) were performed with the program PhyML version 3.0 (Guindon and Gascuel 2003). The confidence levels were estimated with 1000 bootstrap replicates. The best fit substitution models for each of the data combinations were determined using jModeltest 0.1.1 (Posada 2008). MrBayes v. 3.2 (Ronquist et al. 2012) was implemented to perform the Bayesian inference (BI) analyses. Trees were sampled at every 100th generation for six million generations and the posterior probability values above 0.95 were accepted.

Results

Disease symptoms and sample collection

The fruit, leaf and floral sepal disease of *B. racemosa* was found in the Mapelane area on the southern border of the Isimangaliso Wetland Park, the city of Richards Bay and in Mlalazi Nature Reserve near the town of Mtunzini. Symptoms were mostly observed during summer, particularly January to April. Isolates of the fungus consistently associated with this disease were obtained from the aerial organs of ten trees from Mapelane, as well as from ten trees each in two areas in Richards Bay. Three representative isolates from fruits, leaves and sepals were selected for identification based on DNA sequence data.

The disease affecting *B. racemosa* was characterized by sooty black spots on the fruits (Fig. 1A, 1B); on the floral sepals (1C, 1D) as well as dark stromata becoming confluent to produce large spots on the leaves (Fig. 1E, 1F). These spots could merge and cover the entire surface of the aerial organs affected. Leaf infection often resulted in leaf



Fig. 1 Signs and symptoms of disease caused by *Pseudocercospora mapelanensis* on *B. racemosa* (A, B). Fruit disease caused by *P. mapelanensis* (C, D). Sooty black spots on floral sepals (E). Leaf spots on the abaxial part of the leaf (F). Close up of leaf spots caused by *P. mapelanensis*.

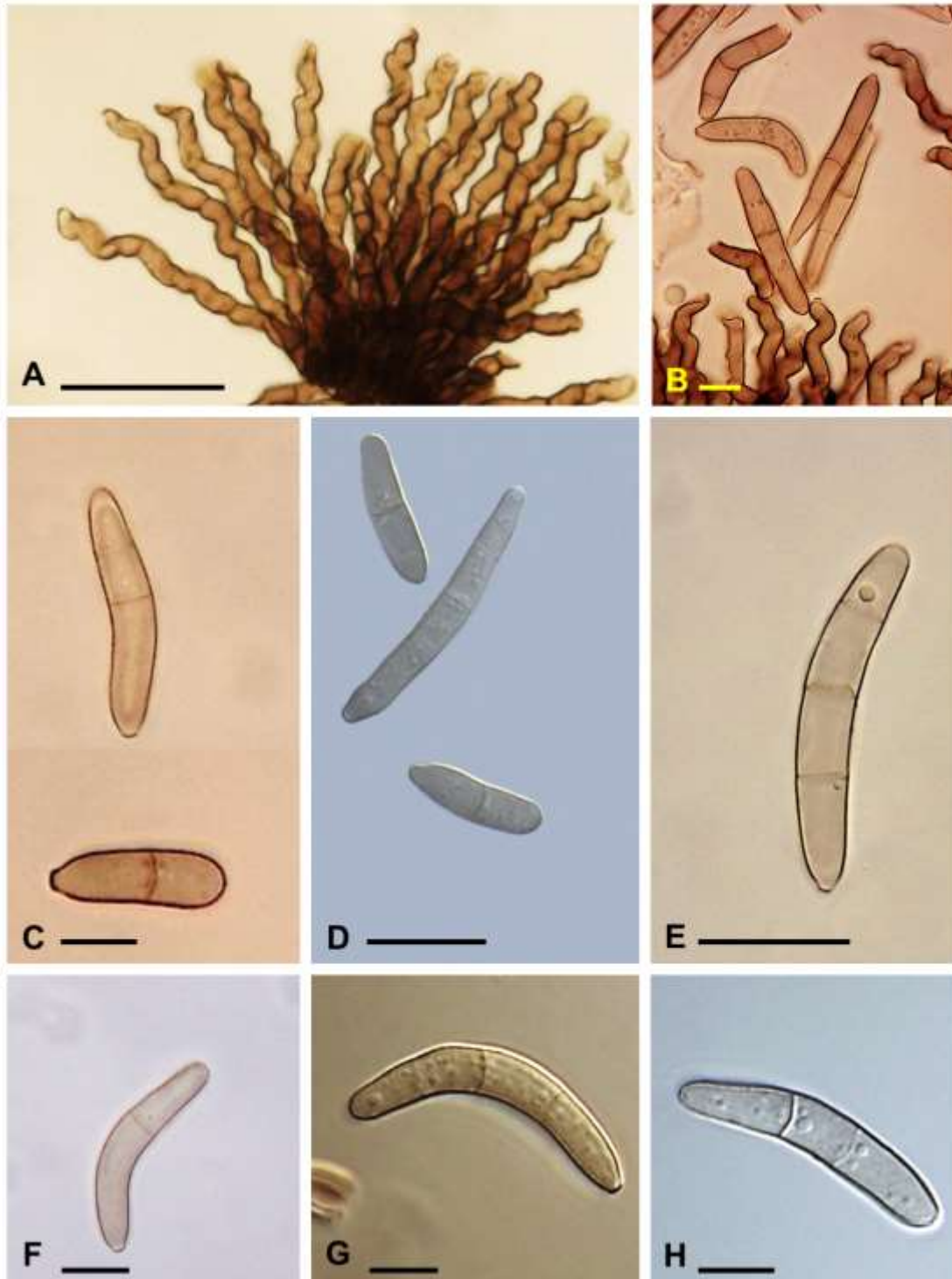


Fig. 2 Conidiophores and conidia of *Pseudocercospora mapelanensis* (A). Geniculate conidiophores, observed with zoom (40x) bar (50 µm). (B) Conidiophores and conidia observed with zoom (40x) (bar 10 µm). (C, D, E, F, G, H) Pale brown conidia with truncate base, different shape, length and septa observed with zoom (100x) (bar C, F, G, H = 10 µm), (bar D, E = 20 µm).

deformation (Fig. 1E). Brownish conidiophores and pale brown conidia were commonly observed under the microscope (Fig. 2). Although the disease did not result in defoliation of trees, flower and fruit abortion was common on affected trees at Mapelane and Richards Bay.

Identification

PCR Amplification and Sequencing

DNA was successfully extracted from the isolates and PCR and sequence products generated for all selected gene regions. Sequence fragments were approximately 367bp in size for the ACT, 485bp for the ITS, 826bp for the LSU and 456bp for the TEF-1 α . All sequences used in this study have been deposited in GenBank (Table 1).

Phylogenetic analyses

Blast searches in Genbank, of sequences generated for the LSU and ITS gene regions, showed that all isolates obtained from diseased *B. racemosa* represented a species of *Pseudocercospora*. Based on these results, data sets were assembled for all gene regions sequenced, including sequence data for previously described, published species of *Pseudocercospora* (Table 1).

Sequence data sets of the ACT, ITS, LSU, and TEF-1 α gene regions were analyzed individually (Table 1). A phylogenetic re-construction was conducted for the aligned LSU data set to determine generic relationships. The LSU data set comprised species from the five closest related sister genera in the Mycosphaerellaceae, based on Blast searches and published literature. These included *Cercospora* Fresen., *Mycosphaerella* Johanson., *Passalora* Fresen., *Pseudocercospora* Speg., *Septoria* Sacc., and *Pallidocercospora* Crous.

Table 1 Isolates included in phylogenetic analyses. Accession numbers of DNA sequences generated in the present study are printed in bold.

Species	Isolate No	Collector	Host	Country	GenBank Accession Numbers			
					LSU	ITS	TEF-1- α	ACT
<i>Cercospora capsici</i>	CBS118712	P. Tyler	Unknown	Fiji	GU214653			
<i>C. sojina</i>	CPC12322	H.D. Shin	<i>Glycine soja</i>	South Korea	GU253861			
<i>Cladosporium cladosporioides</i>	CBS 10921				EU019262			
<i>C. herbarum</i>	AFTOL-ID 1591				DQ678074			
<i>Mycosphaerella africana</i>	CMW4945	P.W. Crous	<i>Eucalyptus viminalis</i>	South Africa	DQ246257			
<i>M. ellipsoidea</i>	CMW4934	Unknown	<i>Eucalyptus</i> sp.	South Africa	DQ246253			
<i>M. madeirae</i>	CBS112895	S. Denman	<i>E. globulus</i>	Portugal	DQ204756			
<i>M. laricina</i>	CBS326.52	E. Müller	<i>Larix decidua</i>	Switzerland	GU253693			
<i>Pallidocercospora acaciigena</i>	CPC3838	M.J. Wingfield	<i>Acacia mangium</i>	Venezuela	GU214661			
<i>Pal. crystallina</i>	CMW3033	M.J. Wingfield	<i>E. bicostata</i>	South Africa	DQ204747			
<i>Pal. heimii</i>	CMW4942	P.W. Crous	<i>Eucalyptus</i> sp.	Madagascar	DQ204751			
<i>Pal. heimioides</i>	CMW3046	M.J. Wingfield	<i>Eucalyptus</i> sp.	Indonesia	DQ204753			
<i>Pal. irregulariramosa</i>	CMW4943	M.J. Wingfield	<i>E. saligna</i>	South Africa	DQ204754			
<i>Pal. konae</i>	CBS120748	W. Himamam	<i>E. camaldulensis</i>	Thailand	GU253852			
<i>Passalora eucalypti</i>	CBS111318	P.W. Crous	<i>E. saligna</i>	Brazil	GU253860	GU269845	GU384558	GU320548
<i>Pseudocercospora abelmoschi</i>	CPC14478	H.D. Shin	<i>Hibiscus syriacus</i>	South Korea	GU253696	GU269647	GU384365	GU320355
<i>P. acericola</i>	CBS122279	R. Kirschner	<i>Acer albopurpurascens</i>	Taiwan	GU253699	GU269650	GU384368	GU320358
<i>P. ampelopsis</i>	CPC11680	H.D. Shin	<i>Ampelopsis brevipedunculata</i>	South Korea	GU253846	GU269830	GU384542	GU320534
<i>P. angolensis</i>	CBS14953	M.C. Pretorius	<i>Citrus</i> sp.	Zimbabwe	JQ324941	JQ324975	GU384548	JQ325010
<i>P. arecacearum</i>	CBS118406	C.F. Hill	<i>Rhopalostylis sapidis</i>	New Zealand	GU253704	GU269655	GU384373	GU320363
<i>P. atromarginalis</i>	CBS114640	C.F. Hill	<i>Solanum</i> sp.	New Zealand	GU253706	GU269658	GU384376	GU320365
<i>P. avicenniae</i>	Mycobank: MB515468	R.G. Shivas & P.W. Crous	<i>Avicennia marina</i>	Australia				
<i>P. chengtuenensis</i>	CPC10696	H.D. Shin	<i>Lycium chinense</i>	South Korea	JQ324942	GU269673	GU384390	GU320379
<i>P. cladosporioides</i>	CBS117482	P.W. Crous	<i>Olea europaea</i>	Tunisia	JQ324944	GU269678	GU384395	GU320383
<i>P. coprosmae</i>	CBS114639	C.F. Hill	<i>Coprosma robusta</i>	New Zealand	JQ324946	GU269680	GU384397	GU320386

Species	Isolate No	Collector	Host	Country	GenBank Accession Numbers			
					LSU	ITS	TEF-1- α	ACT
<i>P. crispans</i>	CPC14883	P.W. Crous	<i>Eucalyptus</i> sp.	South Africa	GU253825	GU269807	GU384518	GU320510
<i>P. cruenta</i>	CPC10846	H. Booker	<i>Vigna</i> sp.	Trinidad	GU269688	GU269688	GU384404	JQ325012
<i>P. cydoniae</i>	CPC10678	H.D. Shin	<i>Chaenomeles speciosa</i>	South Korea	GU253732	GU269691	GU384407	GU320396
<i>P. dodonaeae</i>	CBS114647	C.F. Hill	<i>Dodonaea viscosa</i>	New Zealand	JQ324948	GU269697	GU384413	JQ325013
<i>P. dovyalidis</i>	CPC13771	P.W. Crous	<i>Dovyalis zeyheri</i>	South Africa	GU253818	GU269800	GU384513	GU320503
<i>P. eucalyptorum</i>	CPC10916	P.W. Crous	<i>Eucalyptus</i> sp.	South Africa	GU253788			
<i>P. eustomatis</i>	CBS110822	G. Dal Bello	<i>Eustoma grandiflorum</i>	Argentina	GU253744	GU269705	GU384421	GU320409
<i>P. fori</i>	CMW9095	G.C. Hunter	<i>E. grandis</i>	South Africa	DQ204748	AF468869	DQ211664	DQ147618
<i>P. fraxinites</i>	CPC10743	H.D. Shin	<i>Fomtanesia phillyraeoides</i>	South Korea	GU253720	GU269672	GU384389	GU320378
<i>P. fuligena</i>	CPC12296	Z. Mersha	<i>Lycopersicon</i> sp.	Thailand	JQ324953	GU269711	GU384427	GU320415
<i>P. hakeae</i>	CBS112226	P.W. Crous & B. Summerell	<i>Grevillea</i> sp.	Australia	GU253805	GU269784	GU384495	JQ325017
<i>P. humulicola</i>	CPC10049	H.D. Shin	<i>Humulus scandens</i>	South Korea	JQ324955	GU269724	JQ324996	JQ325018
<i>P. mapelanensis</i>	CMW40579	J.A. Osorio & J. Roux	<i>B. racemosa</i>	South Africa	KM203119	KM203116	KM203122	KM203125
<i>P. mapelanensis</i>	CMW40580	J.A. Osorio & J. Roux	<i>B. racemosa</i>	South Africa	KM203120	KM203117	KM203123	KM203126
<i>P. mapelanensis</i>	CMW40581	J.A. Osorio & J. Roux	<i>B. racemosa</i>	South Africa	KM203121	KM203118	KM203124	KM203127
<i>P. indonesiana</i>	CBS122473	I.W. Buddenhagen	<i>Musa</i> sp.	Sumatra	GU253765	GU269735	GU384448	GU320437
<i>P. kiggelariae</i>	CPC11853	W. Gams	<i>Kiggelaria africana</i>	South Africa	GU253762	GU269730	GU384443	GU320432
<i>P. libertiae</i>	CBS114643	C.F. Hill	<i>Libertia ixioides</i>	New Zealand	JQ324959	GU269733	GU384446	GU320435
<i>P. longispora</i>	CBS122470	D.R. Jones	<i>Musa</i> sp.	Malaysia	GU253764	GU269734	GU384447	GU320436
<i>P. luzardii</i>	CPC2556	A.C. Alfenas	<i>Hancornia speciosa</i>	Brazil	GU214477	GU269738	GU384450	GU320440
<i>P. musae</i>	CBS116634	J. Carlier	<i>Musa</i> sp.	Cuba	GU253775			
<i>P. oenotherae</i>	CPC10290	H.D. Shin	<i>Oenothera odorata</i>	South Korea	JQ324961	GU269856	GU384567	GU320559
<i>P. plectranthi</i>	CPC11462	H.D. Shin	<i>Plectranthus</i> sp.	South Korea	JQ324962	GU269791	GU384501	GU320492
<i>P. proteae</i>	CPC15217	F. Roets	<i>Protea mundii</i>	South Africa	GU253826	GU269808	GU384519	GU320511
<i>P. prunicola</i>	CPC14511	H.D. Shin	<i>Prunus x yedoensis</i>	South Korea	GU253723	GU269676	GU384393	GU320382
<i>P. punctata</i>	CPC14734	P.W. Crous	<i>Syzygium</i> sp.	Madagascar	GU253791	GU269765	GU384477	GU320468
<i>P. ranjita</i>	CPC11141	M.J. Wingfield	<i>Gmelina</i> sp.	Indonesia	GU253810	GU269790	GU384500	GU320491

Species	Isolate No	Collector	Host	Country	GenBank Accession Numbers			
					LSU	ITS	TEF-1- α	ACT
<i>P. ravenalicola</i>	CBS122468	M. Arzanlou & W. Gams	<i>Ravenala</i> <i>madagascariensis</i>	India	GU253828	GU269810	GU384521	GU320513
<i>P. rhoina</i>	CPC11464	H.D. Shin	<i>Rhus chinensis</i>	South Korea	JQ324966			
<i>P. rubi</i>	MUCC875	T. Kobayashi & C. Nakashima	<i>Rubus allegheniensis</i>	Japan	GU253795	GU269773	GU384485	GU320476
<i>P. rumohrae</i>	CBS117747	C.F. Hill	<i>Marattia salicina</i>	New Zealand	GU253796	GU269774	GU384486	GU320477
<i>P. subsessilis</i>	CBS136.94	R.F. Castaneda		Cuba	GU253832	GU269815	GU384527	GU320517
<i>P. subtorulosa</i>	CBS117230	R. Kirschner	<i>Melicope</i> sp.	Taiwan	GU253833			
<i>P. tereticornis</i>	CBS124996	A.J. Carnegie	<i>E. nitens</i>	Australia	GQ852647			
<i>P. timorensis</i>	MUCC 819	C. Nakashima & T. Akashi	<i>Ipomoea indica</i>	Japan	GU253840	GU269823	GU384536	GU320526
<i>P. viticicola</i>	MUCC777	C. Nakashima	<i>Vitex trifolia</i>	Japan	GU253845	GU269828	GU384540	GU320532
<i>P. vitis</i>	CPC11595	H.D. Shin	<i>V. vinifera</i>	South Korea	GU214483	GU269829	GU384541	GU320533
<i>P. xanthocercidis</i>	CPC11665	A.R. Wood	<i>Xanthocercis</i> <i>zambesiaca</i>	South Africa	JQ324971			
<i>P. xanthoxyli</i>	CPC10065	H.D. Shin	<i>Xanthoxylum ailathoides</i>	South Korea	GU253848	GU269832	GU384544	GU320536
<i>Septoria cerastii</i>	CPC12343	H.D. Shin	<i>Cerastium holosteoides</i> var. <i>hallasanense</i>	South Korea	GU253869			
<i>S. chelidonii</i>	CPC12337	H.D. Shin	<i>Chelidoniummajur</i> var. <i>asiaticum</i>	South Korea	GU253870			
<i>S. crepidis</i>	CPC12539	H.D. Shin	<i>Crepis japonica</i>	South Korea	GU253871			
<i>S. dysentericae</i>	CPC12328	H.D. Shin	<i>Inula britannica</i> var. <i>chinensis</i>	South Korea	GU253866			
<i>S. erigerontis</i>	CPC12340	H.D. Shin	<i>Erigeron annuus</i>	South Korea	GU253872			
<i>S. eucalyptorum</i>	CPC11282	W. Gams	<i>Eucalyptus</i> sp.	India	GU253873			
<i>S. justiciae</i>	CPC12509	H.D. Shin	<i>Justicia procumbens</i>	South Korea	GU253874			
<i>S. quercicola</i>	CBS663.94	H.A. Van der Aa	<i>Quercus robur</i>	Netherlands	GU253867			

CBS: Centraalbureau,voor Schimmelcultures, CBS-KNAW Fungal Biodiversity Centre, Utrecht, The Netherlands; CMW: Culture Collection of the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, Pretoria, South Africa; MUCC Murdoch University Algal Culture Collection, Murdoch, Western Australia; CPC Collection of Pedro Crous housed at CBS.

LSU: partial 28S nrRNA gene; ITS: internal transcribed spacer regions 1 & 2 including 5.8S nrRNA gene; EF-1 α : partial translation elongation factor 1-alpha gene; ACT: partial actin gene.

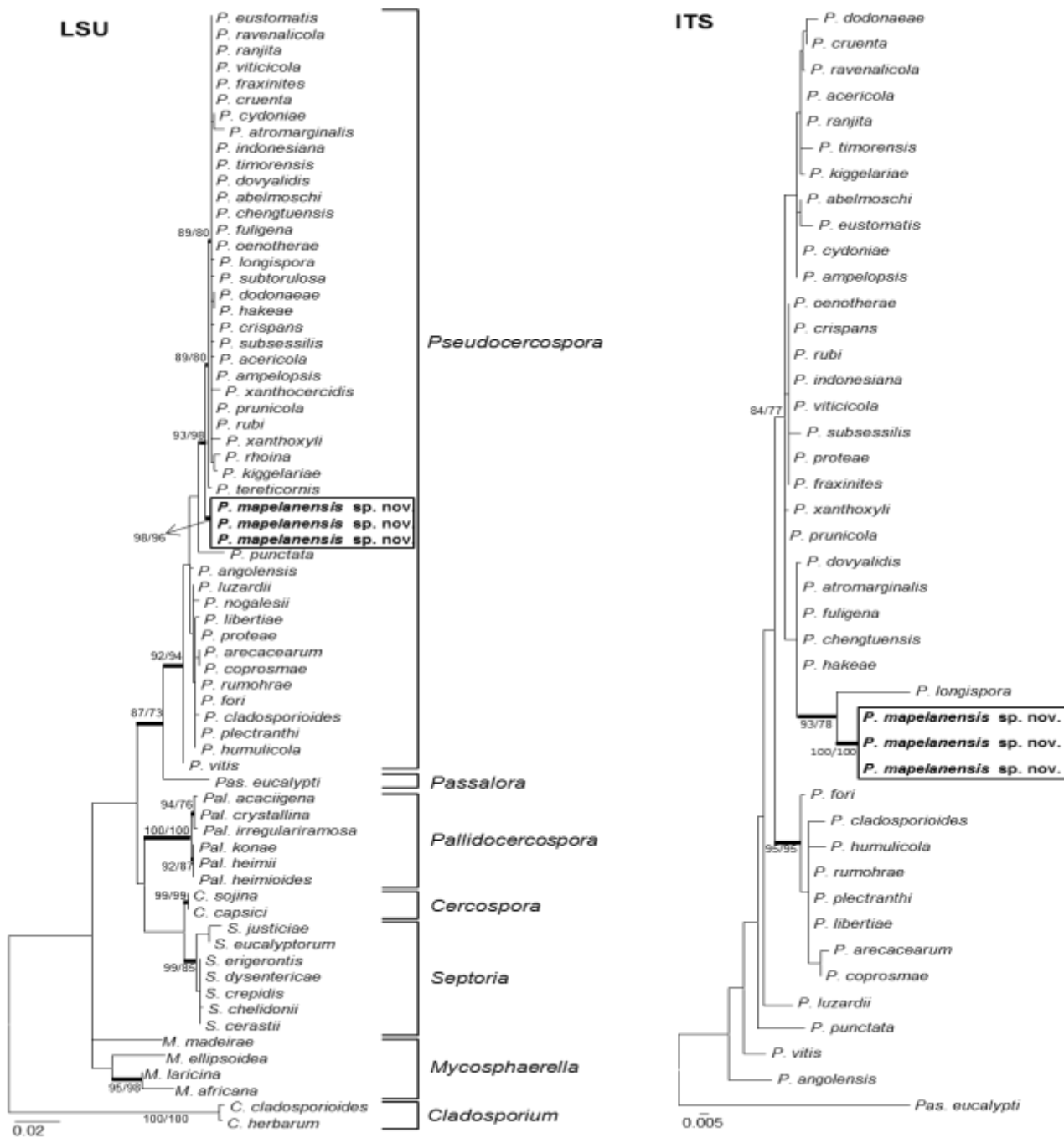


Fig. 3 Phylograms obtained from (BI), (ML) and (MP) analyses of the LSU and ITS data sets. These analyses provide evidence of isolates obtained from *B. racemosa* grouping into the *Pseudocercospora* clade, indicated in the LSU tree. The ITS analyses suggest that isolates from *B. racemosa* represent an undescribed species, with *P. longispora* as the most closely related species to *P. mapelanensis* sp. nov., (BI) posterior probabilities $\geq 95\%$ are represented by thick branches. Bootstrap support values $>70\%$ are indicated near the nodes as MP/ML. * = bootstrap support values $<70\%$.

Cladosporium cladosporioides (Fresen.) G.A. de Vries., and *Cladosporium herbarum* (Pers.) Link, were included as outgroups. The dataset for the LSU phylogenetic analyses comprised a total of 69 taxa. ML analyses resulted in 1000 trees and MP analyses in 7304 trees (Fig. 3).

The isolates obtained from *B. racemosa* grouped strongly within the genus *Pseudocercospora* in all analyses. Subsequently, a species level phylogeny was derived from individual analysis of the ACT, ITS and TEF-1 α alignment of described species of *Pseudocercospora*, with *Passalora eucalypti* (Crous & Alfenas) Crous & U. Braun, included as outgroup. A total of 43 taxa were included in the analyses for each of the three gene regions. ML analyses of the ITS data set resulted in 1000 trees and MP analyses in 93765 trees. These trees all suggested that isolates from *B. racemosa* represent an undescribed species, most closely related to, but distinct from, *P. longispora* (Fig. 3).

Phylogenetic analyses involving sequence data from the ACT and TEF gene regions helped to clarify the placement of the species from *B. racemosa* (Fig. 4). ML analysis of the ACT gene region resulted in 1000 trees and MP analysis in 4274 trees. The ML analysis of the TEF gene region data set resulted in 1000 trees and MP analysis in 156 trees. These trees also suggested that isolates from *B. racemosa* represent an undescribed species of *Pseudocercospora*, showing *P. dodonaeae* as the closest relative in the ACT gene region and *P. xanthoxyli* as the closest relative in the TEF-1 α gene region.

Taxonomy

Based on sequence comparisons for multiple gene regions, the fungus obtained from diseased *B. racemosa* in South Africa represents an undescribed species of *Pseudocercospora*. We provide the following description based on morphological characteristics.

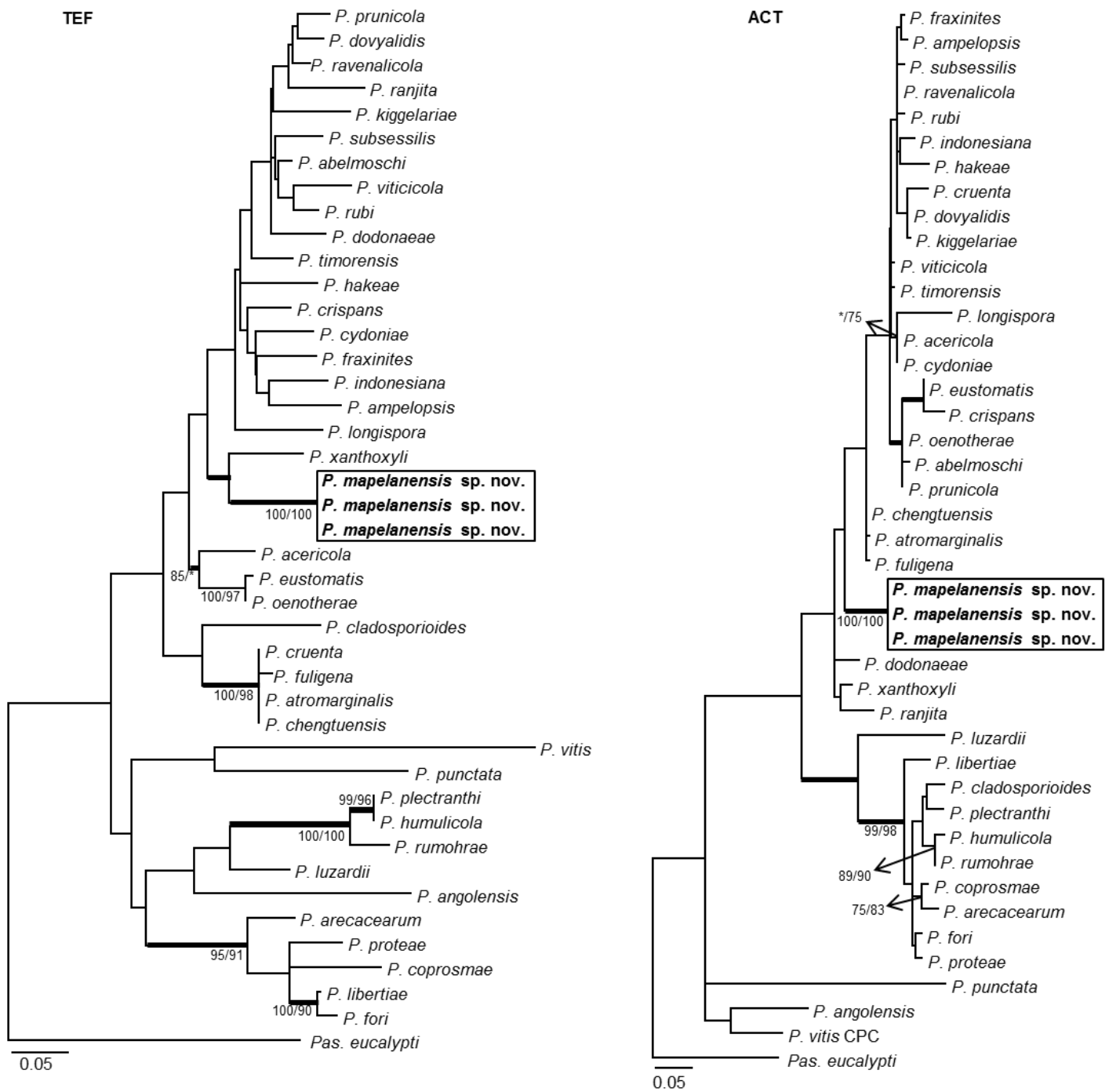


Fig. 4 Phylograms obtained from (BI), (ML) and (MP) analyses of the TEF-1 α and ACT genes. These analyses show *P. xanthoxyli* as the closest relative in TEF and *P. dodonaeae* as the closest species in ACT. These analyses suggest that isolates from *B. racemosa* represent an undescribed species of *Pseudocercospora* (printed in bold and included in the box). (BI) posterior probabilities $\geq 95\%$ are represented by thick branches. Bootstrap support values $>70\%$ are indicated near the nodes as MP/ML. * = bootstrap support values $<70\%$.

Pseudocercospora mapelanensis J.A Osorio & Jol. Roux (Fig. 1, 2)

MycoBank MB 809625.

Diagnosis: Morphologically akin to *Pseudocercospora barringtoniicola*, but conidia shorter and wider, (19-)24-38(-45) x (4-)5-7(-10) μm , only 1-3-septate, conidiogenous loci sometimes subconspicuous, circle-like in front view, and hila of the conidia often somewhat refractive or even darkened-refractive and thus more conspicuous.

No sexual morph was observed on leaves, fruits or flowers. Leaf spots amphigenous, subcircular to angular-irregular, 2-18 mm diam or confluent and larger, margin indefinite or surrounded by a diffuse, narrow to wider, paler halo, pale greenish, yellowish to ochraceous. Caespituli amphigenous, also on fruits, punctiform to pustulate, scattered to gregarious, dark brown. Asexual *stromata* (10-)20-60 μm diam, finally sometimes expanded or confluent and larger, to 100 μm diam or even larger, brown to blackish brown, sub-globose, erumpent on leaves and fruits, composed of swollen hyphal cells, subglobose to slightly angular-irregular in outline, 2-6 μm diam, amphigenous, gregarious on both surfaces, denser on the abaxial surface, circular to irregular in shape, at first smaller and substomatal, later expanding or confluent and also immersed, giving rise to conidiophores. *Conidiophores* (10-)20-116(-136) x (3-)4-7(-8) μm , mostly hypophyllous, in small, loose to often large or even very large and dense fascicles, usually arising from a dark stroma, emerging through stomata or erumpent, septate, growing sympodially, moderately to mostly strongly geniculate-sinuuous, truncate apex, sometimes obtusely rounded, unbranched, at first sub-hyaline to pale olivaceous, later brownish, ranging from pale to medium olivaceous-brown to brown, wall smooth, thin, to about 0.5 μm , occasionally somewhat thicker near the base in older conidiophores, to about 1 μm ; conidiogenous cells integrated, terminal or conidiophores reduced to conidiogenous cells,

conidiogenous loci inconspicuous or subconspicuous by being somewhat refractive around the rim, in from view sometimes visible as minute circle, about 1.5-2 μm diam. *Conidia* solitary, subcylindrical, fusiform to obclavate, straight to somewhat curved, (19-) 24-38(-45) x (4-)5-7(-10) μm , 1-3-septate, subhyaline to pale brown or olivaceous-brown, smooth, thin-walled, apex obtuse and rounded, base short obconically truncate, (1.5-)2(-2.5) μm wide, hila unthickened, usually not darkened, but often somewhat refractive or occasionally even slightly darkened-refractive.

Colonies slow growing (reaching 2 mm diameter after 15 days, and 5-6 mm after 30 days) on MEA (Malt Extract Agar) at 25°C. Colonies with circular to irregular blackish margins, raising centrally, of dense cottony mycelium and hard texture. Young colonies dark, turning to greyish white or green after one month, reverse colony blackish to dark blackish brown. Despite trying different culture media types no sporulation of *P. mapelanensis* was observed in cultures.

Etymology: Epithet refers to one of the areas in South Africa (Mapelane Nature Reserve) where *Barringtonia racemosa* occurs and where the disease was first observed.

Habitat. Symptomatic fruits, floral sepals and leaves of *Barringtonia racemosa*.

Known distribution in South Africa: Kwazulu-Natal Province.

Specimens examined. SOUTH AFRICA, KWAZULU-NATAL PROVINCE: Mapelane Nature Reserve, from symptomatic fruits, flowers and leaves. In Herb. PREM (**Holotype** PREM 61109, Paratype PREM 61107, PREM 61108, col. J.A Osorio & Jol. Roux). Isotype HAL 2680 F. Other cultures: CMW (Ex-type CMW 40581, Ex-Paratypes CMW 40579, CMW 40580). CBS (ex-type CBS 138923, Ex-Paratype CBS 138922).

The following is a dichotomous key to *Pseudocercospora* species on *Barringtonia* species, adapted by Braun from the Cercosporoid hyphomycetes on *Barringtonia* spp. (Braun and Mouchacca 2000).

1. Stromata lacking or very small, only with a few substomatal cells; conidiophores in small, loose fascicles, 2-8, very long, to 185 μm ; on *B. acutangulae*
..... *P. barringtoniae-acutangulae*
- 1* Stromata well-developed, about 10–100 μm diam 2
2. Conidiophores to 200 μm long, pluriseptate; conidia long and narrow, filiform-acicular to narrowly obclavate-cylindrical, 20–90 x 1.5-5 μm , 2-8-septate, subhyaline to pale green-olivaceous; on *B. asiatica*
..... *P. barringtoniigena*
- 2* Conidiophores shorter, usually < 100 μm ; conidia obclavate-subcylindrical, fusiform, shorter and broader, about 20-60 x 4-7(-10) μm 3
3. Conidiogenous loci and conidial hila unthickened, neither darkened nor refractive; conidia 30-60 x 4-6.5 μm , 3-6-septate; on *B. speciosa*
..... *P. barringtoniicola*
- 3* Conidiogenous loci often subconspicuous, visible as minute circle in from view, conidial hila often conspicuous by being somewhat refractive or even darkened-refractive; conidia shorter, (19-)24-38 x (4-)5-7(-10) μm , only 1-3-septate; on *B. racemosa*
..... *P. mapelanensis*

Discussion

This study provides the first description of a disease of the mangrove associate, *B. racemosa* in South Africa. A fungus in the genus *Pseudocercospora* was consistently and intimately associated with early symptoms of the disease. It readily sporulated on green tissue and all indications are that this is the causal agent of the disease. The *Pseudocercospora* sp., was shown to represent a novel taxon for which the name *Pseudocercospora mapelanensis* has been provided.

Three species of *Pseudocercospora* have previously been described causing infections on *Barringtonia* species (Braun et al. 1999, 2000). These include *P. barringtoniae-acutangulae* U. Braun & Mouch., *P. barringtoniigena* U. Braun & Mouch., and *P. barringtoniicola* U. Braun & Mouch. Unfortunately neither cultures, sequences nor herbarium specimens are available for these species and they could thus not be included in this study for comparative purposes. However, based on the published descriptions of the symptoms and micro-morphology, these species are all clearly different from *P. mapelanensis*.

Pseudocercospora barringtoniae-acutangulae was first described from *B. acutangulae* in India causing irregular to sub-circular small leaf spots and later becoming confluent to form large patches on the leaves (Braun and Mouchacca 2000). *Pseudocercospora barringtoniigena* was reported from *B. asiatica* in the South Pacific (Futuna) and it is characterized causing sub-circular and amphigenous leaf spots, and similar to *B. acutangulae* in causing large patches on the leaves when lesions become confluent (Braun and Mouchacca 2000). *Pseudocercospora barringtoniicola* was first described from *B. speciosa* in French Polynesia, Tahiti. This species forms large sub-circular to irregular spots on both sides of the leaves and the lesions become blackish during fungus

Table 2 Comparison between described species of *Pseudocercospora* associated with *Avicennia marina* and *Barringtonia* species

	<i>Pseudocercospora mapelanensis</i>	<i>P. barringtoniae-acutangulae</i>	<i>P. barringtoniigena</i>	<i>P. barringtonicola</i>	<i>P. avicenniae</i>
Host Species	<i>B. racemosa</i>	<i>B. acutangulae</i>	<i>B. asiatica</i>	<i>B. speciosa</i>	<i>Avicennia marina</i> (Acanthaceae)
Conidiophores (µm)	(10-)20-116(-136) x (3-)4-7(-8) pluriseptate	30-185 x (3-)4-7 2-6 septa	20-200 x 3-7 pluriseptate	10-50 x 3-6 0-3 septa	20-90 x 3-5.5
Conidia (µm)	(19-)24-38(-45) x (4-)5-7(-10) (0-)1-3(-4) septate	20-70(-85) x 4-4.5-7 (-8) 1-6(-7) septate	20-90 x 1.5-5 2-8-septate	30-60 x 4-6.5 3-6 septate	30-100 x 3-5 3-12-septate
Region	South Africa (Mapelane and Richards Bay)	India (Uttar Pradesh, Gorakhpur)	South Pacific (Futuna)	French Polynesia, Tahiti, Rurutu	Australia
Reference	Current study	Braun & Mouchacca 2000	Braun & Mouchacca 2000	Braun et al. 1999	Shivas et al. 2009

fructification (Braun et al. 1999). Based on morphological descriptions provided for these fungi they differ from the South African fungus in a number of characteristics (Table 2). Conidiophores and conidia of *P. barringtoniae-acutangulae* and *P. barringtoniigena* are larger than those of *P. mapelanensis*. Conidiophores of *P. barringtonicola* are shorter, and conidia larger, than those of *P. mapelanensis*. Species previously described from *Barringtonia* also differ in the number of conidial septa, with *P. mapelanensis* having between one and three septa while the other species have more than three transverse septa.

Sequence data for the *Pseudocercospora* species known to occur on *Barringtonia* are not available for comparison and the lack of cultures preclude this work. However, multiple phylogenetic analyses showed that *P. mapelanensis* is most closely related to, but distinct from, *P. dodonaeae* Boesewinkel, *P. longispora* Arzanlou & Crous, and *P. xanthoxyli* (Cooke) Y.L. Guo & X.J. Liu. Based on multiple phenotypic characters such as conidial and conidiophores shape, dimensions and number of septa, these species can also be distinguished from *P. mapelanensis*.

It was not possible to conduct pathogenicity tests with *P. mapelanensis* in this study. This was due to the fact that the fungus does not sporulate in culture and the host trees are rare and could not be obtained for intensive study. However, all aspects of infections indicated that this is a highly host specific fungus, sporulating on freshly infected green tissue. This is consistent with many *Pseudocercospora* species and their *Mycosphaerella* sexual states that are highly host-specific and that are commonly treated as pathogens in the absence of pathogenicity tests (Crous and Braun 2003, Crous et al. 2004, Burgess et al. 2007, Crous et al. 2008, Crous 2009). All indications from this study are that *P. mapelanensis* is the causal agent of the disease of *B. racemosa* and it was the only fungus found consistently on infected tissue. The disease of *B. racemosa* was often severe and although it did not

cause defoliation, it appeared to result in significant levels of fruit abortion. It is probable that the pathogen is native on its host but this is a question that deserves further study.

Acknowledgements

This work was financially supported by the Department of Science and Technology (DST) and National Research Foundation (NRF) Center of Excellence in Tree Health Biotechnology (CTHB). We thank Ezemvelo KZN Wildlife and the Isimangaliso Wetland Park for sampling permits and assistance in the field. The material was collected under EKZMW permit no OP 4776. The authors gratefully acknowledge Prof. dr U. Braun (Martin-Luther-University, Institute of Biology, Halle, Germany), for his assistance in revising the fungal material and for his valuable input in the diagnosis of the new *Pseudocercospora* species. We also acknowledge Ariska Van der Nest, Arista Fourie and James Mehl (FABI, University of Pretoria), for technical assistance.

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