#### **ORIGINAL RESEARCH ARTICLE**



# *Botryosphaeriaceae* species associated with branch dieback and decline of macadamia trees in South Africa

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#### Abstract

*Botryosphaeriaceae* species are important latent pathogens causing diseases on trees utilized in forestry and agriculture. In recent years, there has been an increase in the incidence and severity of branch dieback and decline on macadamia trees in South Africa, and species of *Botryosphaeriaceae* have been considered as a possible cause. Although botryosphaeria dieback has been well-studied in Australia, there is little information regarding these fungi on Macadamia in South Africa. The aims of this study were consequently to (i) identify species of *Botryosphaeriaceae* from Macadamia branches from main production regions in South Africa, (ii) compare the diversity of species between symptomatic and asymptomatic branches, as well as between different growing regions, (iii) and to consider their relative importance in causing dieback. Eight species and three putative hybrids of the *Botryosphaeriaceae* were identified based on a phylogenetic comparison of sequence data from the ITS rDNA, *tub2*, *tef-1α* and *rpb2* loci. These included an unidentified *Diplodia* sp., and *Lasiodiplodia* sp., as well as *L. gilanensis, L. theobromae, L. pseudotheobromae, Neofusicoccum kwambonambiense, N. luteum, N. parvum* and three hybrid species. The unidentified species of *Diplodia., Lasiodiplodia* sp., *L. gilanensis,* and *N. kwambonambiense* are reported for the first time on Macadamia in South Africa. All species showed a potential to cause branch dieback symptoms, with species of *Neofusicoccum* identified as the most aggressive species. This study revealed a high level of diversity of *Botryosphaeriaceae* species and illustrates their potential as causal agents of dieback on Macadamia in South Africa.

Keywords Fungal endophyte · Species diversity · Pathogen · Aggressiveness

# Introduction

*Macadamia* is a nut-bearing tree belonging to the family *Proteaceae*. It is indigenous to southeastern Australia, but is planted commercially in several tropical and subtropical countries of Africa, Asia, as well as North and South America in addition to Australia (Cann 1965; McConachie 1980; McHargue 1996). The Macadamia industry has expanded rapidly in recent years and today large areas in various countries are under cultivation. South Africa is currently the leading producer of Macadamia nuts, followed

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<sup>2</sup> Plant Microbiology Division, ARC-Plant Health and Protection, P/Bag X134, Queenswood, Pretoria 0121, South Africa by Australia (SAMAC 2022). The increase in Macadamia production in the last few years is linked to the fact that it is the highest priced processed nut on the world market. Furthermore, Macadamia nuts are also highly sought after for their rich flavor, texture, and health benefits (Macfarlane and Harris 1981).

Macadamia trees, both in their native and introduced ranges, are commonly affected by various fungal pathogens that have resulted in significant yield losses affecting the productivity of the Macadamia industry (Akinsanmi et al. 2017; Drenth et al. 2009; Hunter and Kunimoto 1973; Ko and Kunimoto 1994; Wrona et al. 2020). Notable diseases include stem canker caused by *Phytophthora* species (Jeff-Ego et al. 2020), husk spot caused by *Pseudocercospora macadamiae* (Beilharz et al. 2003), dry flower disease caused by *Pestalotiopsis macadamiae* and *Neopestalotiopsis macadamiae* (Akinsanmi et al. 2017), phomopsis husk rot caused by *Diaporthe* species and botryosphaeria dieback caused by *Botryosphaeriaceae* species (Jeff-Ego and Akinsanmi 2019; Wrona et al. 2020). Branch dieback is one of the most commonly reported symptoms on woody crop plants (Jurskis 2005; Landsberg and Wylie 1988). It can be caused by multiple biotic factors including pests and diseases, as well as environmental stressors such as drought and salt damage (Jurskis 2005; Landsberg and Wylie 1988). *Phytophthora* and *Botryosphaeriaceae* species are known to cause branch dieback on Macadamia (Jeff-Ego and Akinsanmi 2019; Jeff-Ego et al. 2020). Branch dieback symptoms on Macadamia caused by *Phytophthora* include leaf chlorosis and loss of foliage (Jeff-Ego et al. 2020), while botryosphaeria branch dieback is characterized by browning of leaves that remain attached to trees accompanied by a wedge-shaped discoloration in cross-sections of the affected wood (Jeff-Ego and Akinsanmi 2019).

The Botryosphaeriaceae is a diverse family of fungi with a cosmopolitan distribution. Species are considered to be class III endophytes implying that they have the potential to become latent or opportunistic pathogens (Sakalidis et al. 2011; Mehl et al. 2013). As pathogens they often cause serious dieback and decline on a wide range of hosts (Slippers and Wingfield 2007; Mehl et al. 2013; Jami et al. 2014; Moral et al. 2019). Another notable aspect of the ecology of this group of fungi is their lack of host specificity, thus enabling them to colonize and cause disease on diverse native and introduced plant hosts in a particular region (Slippers and Wingfield 2007; Pérez et al. 2008; Mehl et al. 2013; Jami et al. 2014). Due to the endophytic nature of these fungi, they are easily introduced into new areas, unnoticed either on seeds or cuttings, later infecting other trees (Slippers and Wingfield 2007; Mehl et al. 2013; Marsberg et al. 2017).

The incidence and severity of Botryosphaeria dieback disease on Macadamia has increased significantly in Australia over the last 15 years (Jeff-ego and Akinsanmi 2019; Mohankumar et al. 2022). In total, thirteen *Botryosphaeriaceae* species have been associated with the branch dieback disease in Australia namely, *Botryosphaeria dothidea, Lasiodiplodia jatrophicola, L. iraniensis, L. pseudotheobromae, L. theobromae, Neofusicoccum australe, N. luteum, N. mangroviorum, N. parvum,* and four undescribed *Lasiodiplodia* species (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). While these fungi are not as virulent as some primary pathogens, the dieback and canker diseases caused by *Botryosphaeriaceae* on Macadamia are amongst the most common and, under some conditions, the most serious problems affecting these trees.

In recent years, there have also been increasing reports of branch dieback and decline of Macadamia trees in South Africa. While considerable work has been done to identify species of *Botryosphaeriaceae* and their association with branch dieback in Australia, these fungi have hardly been studied on this host in South Africa. Only *Botryosphaeria ribis* has been reported causing disease on Macadamia in South Africa (Herbert and Grech 1985). The aim of this study was therefore to characterize species of *Botryosphaeriaceae* from Macadamia in South Africa and to consider their pathogenicity on this host.

# Materials and methods

# Disease symptoms, sampling, and fungal isolation

Samples were collected from the KwaZulu-Natal, Limpopo and Mpumalanga growing regions of South Africa in 2018. Samples were collected on two farms per region and at two sites per farm to capture cultivar diversity. At each farm, twenty asymptomatic branches were collected from 20 individual trees using random sampling. A total of 40 asymptomatic samples were therefore collected per growing region, with the number of diseased samples collected being dependent on availability. Symptoms observed in the field included general branch and shoot dieback, internal wood discoloration, and tree decline. The collected samples were placed in brown paper bags and processed the same day. To increase the number of samples obtained from diseased material, additional branch dieback samples received via the Forestry and Agricultural Biotechnology Institute (FABI) disease diagnostic clinic, University of Pretoria, were also included.

Branches were surface disinfested in 10% hydrogen peroxide for 1 min, in 70% alcohol for 1 min, and then rinsed in sterile water and dried on a paper towel. The branches were cut in half to expose the pith. For symptomatic samples, branches were inspected for evidence of discoloration in the pith and sections were taken from the edge of discoloured (necrotic) tissue. Four sterilized tissue sections of approximately 3-5 mm<sup>2</sup> from both symptomatic and asymptomatic branches were then placed onto the 2% malt extract agar (Biolab, MEA; 20 g malt extract, 20 g agar/ 1000 ml distilled water). The isolation plates were incubated at 25 °C for seven days. Single hyphal tips of isolates displaying characteristics of Botryosphaeriaceae (white to grey mycelium with aerial hyphae) were transferred to 2% MEA plates to obtain pure cultures. The pure cultures were subsequently deposited and are maintained in the culture collection (CMW) of the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, Pretoria.

# DNA extraction, PCR amplification and sequencing

DNA was extracted from fungal mycelium of 7-day-old cultures using the PrepMan ® Ultra kit (Thermo fisher Scientific) extraction method. Mycelium was scraped from the edges of colonies for each isolate using a sterile needle tip and transferred into a sterile 1.5-ml tube containing

50  $\mu$ l of the PrepMan Ultra preparation reagent. Mycelial suspensions were shaken in a vortex mixer for 30 s and incubated at 95 °C in a heating block for 10 min. The tubes were centrifuged for 2 min and 50  $\mu$ l of the supernatant was transferred to a new sterile 1.5-ml tube and stored at -20 °C until used for Polymerase Chain Reaction (PCR) reactions. DNA concentrations were determined using a ND-1000 spectrophotometer V3.7.1 (Thermo Fisher Scientific, USA) and diluted with Sabax water to 50 ng/ $\mu$ l.

The Internal Transcribed Spacer (ITS) gene region of the rRNA operon was amplified using the primers ITS1/ ITS4 (White et al. 1990) for initial identification and placement of isolates in the genera of the Botryosphaeriaceae. Thereafter, additional gene regions were amplified for a subset of samples chosen to represent the various genera collected at different geographical origins, farm sites and cultivars (Online Resource Table 1). The Translation Elongation Factor (*tef-1* $\alpha$ ) gene region was amplified using the primers EF1-728F/EF1-986R (Carbone and Kohn 1999), the β-tubulin (tub2) gene region using BT-2a/BT-2b (Glass and Donaldson 1995), and the RNA polymerase II subunit (rpb2) gene was amplified using the primers rpb2-LasF/ rpb2-LasR for Lasiodiplodia (Cruywagen et al. 2017) and rpb2 bot6F/ RPB-2bot7R for Neofusicoccum (Pavlic et al. 2009; Sakalidis et al. 2011).

The total volume of the PCR mixture was 25 µL which consisted of 100-200 ng of genomic DNA, 5 µl of 5 mM MyTaq<sup>™</sup> reaction buffer containing MgCl<sub>2</sub> and dNTPs (Bioline South Africa), 0.5 µl of MyTaq<sup>™</sup> DNA polymerase, 0.5 µl of 10 mM primer for each gene region used, and 17.5 µl of ddH<sub>2</sub>O (Adcock Ingram, Bryanston, South Africa). PCR was conducted in a thermal cycler (Bio-Rad, BioRad Laboratories Inc., Hercules, CA, USA) with the following cycles: initial denaturation at 95 °C for 4 min followed by 35 cycles of denaturation at 95 °C for 35 s; annealing at 55 °C (ITS, *tef1-\alpha*, *tub2*), 56 °C (*rpb2*), for 1 min and extension at 72 °C for 90 s; followed by final extension for 10 min at 72 °C. The PCR amplification products were separated by electrophoresis in 1% agarose gels and run at 100 V and 400 mA for 40 min in 1.0 × Trisborate-EDTA (TBE) buffer. PCR product sizes were estimated using a 100 bp marker and visualized using Bio-Rad Molecular Imager® Gel Doc™ XR System. This was followed by a PCR product clean up reaction using ExoSAP-IT<sup>TM</sup> (Applied Biosystems, Foster city, CA) according to the manufacturer's instructions.

BigDye Terminator v3.1 Cycle Sequencing Kit was used to sequence PCR products in both directions. The thermal cycler was programmed as follows: initial denaturation at 96 °C for 2 min, followed by 25 cycles at 96 °C for 10 s, 50 °C for 5 s and 60 °C for 4 min. The samples were sequenced using an ABI 3100 Automated Capillary DNA Sequencer (Applied Biosystems, USA) at the Bioinformatics Sequencing facility, University of Pretoria.

# **Phylogenetic analyses**

Forward and reverse sequences were checked for accuracy and manually edited using CLC BioWorkbench v.5. Consensus sequences were generated by aligning the forward and reverse sequences. Sequences were submitted to GenBank (Online Resource Table 1). The generated ITS sequences were then subjected to Basic Local Alignment Search Tool (BLAST) searches against the National Center for Biotechnology Information (NCBI) (www.ncbi.nlm.nih.gov) to group the isolates into different genera. Thereafter, ITS, *tub2*, *tef-1* $\alpha$  and *rpb2* sequences of each genus were compiled into a dataset that included known species. The reference sequences and type strains were obtained from papers published by Cruywagen et al. (2017), Pavlic et al. (2009), Phillips et al. (2013) and Zhang et al. (2021). Additional four novel species recently described by Mohankumar et al. (2022) were also included. The sequence dataset was then aligned with Multiple Alignment using Fast Fourier Transform (MAFFT) version 7 (https://mafft.cbrc.jp/alignment/ software).

Phylogenetic analyses of single genes and the concatenated sequence data based on maximum likelihood (ML) were performed by using Randomized Axelerated Maximum Likelihood (RaxML) version 8 (Stamatakis 2014) and viewed in Mega 7.0. Individual trees were generated, and the best substitution models were determined for each dataset with Modeltest v.2.1.3 using the Akaike Information Criterion (AIC). The best-fit evolution model for ITS and *tef1-\alpha* was GTR + G (Stamatakis and Alachiotis 2010) whereas HKY + G was the best model for *tub2* and *rpb2* (Hasegawa et al. 1985). For ML analyses, a bootstrap value of 1000 was set to determine the robustness of the trees. Bayesian inference, based on a Markov Chain Monte Carlo (MCMC) approach, was also performed in MrBayes v.3.1.2 (Ronquist and Huelsenbeck 2003). Datasets for each of the gene regions sequenced were analysed separately and combined to investigate the congruency of the phylogenetic clades obtained. All the phylogenetic trees generated in this study are rooted to Botryosphaeria dothidea. Based on the results of ML, species recognition was determined according to the genealogical species concordance.

#### Pathogenicity trials

Species of *Botryosphaeriaceae* that were isolated more than once were used in pathogenicity tests. A single isolate representing each of these species was randomly selected for the pathogenicity trials under glasshouse conditions (Online Resource Table 1). The isolates were grown on 2% MEA at

 Table 1
 Botryosphaeriaceae isolates used in the phylogenetic analysis and pathogenicity trials in this study

CMW	Species Tissue <sup>3</sup>		ssue <sup>3</sup>	Province Farm Cultivar Date				GenBank Accession number			
								ITS	tub2	tef-1α	rpb2
57,110	Diplodia sp.	Н	Limpopo		1	Beaumont	2018	OR671504	OR770372	OR761223	-
57,111	Diplodia sp.	Н	Limpopo		1	Beaumont	2018	OR671505	OR761373	OR761224	-
57,172	L. gilanensis	D	KwaZulu-Natal		2	Beaumont	2018	OR659658	OR770462	OR772784	OR772795
57,188	L. gilanensis	D	KwaZulu-Natal		2	Beaumont	2018	OR659659	OR770463	OR772783	OR772796
57,182	L. gilanensis	D	KwaZulu-Natal		2	Beaumont	2018	OR659660	OR770464	OR772782	OR772797
57,180	L. gilanensis	D	KwaZulu-Natal		2	Beaumont	2018	OR659661	OR770465	OR772781	OR772798
57,181	L. gilanensis	Н	Limpopo		3	Beaumont	2018	OR659662	OR770466	OR772785	OR772799
57,163	L. gilanensis	Η	Limpopo		3	Beaumont	2018	OR659663	OR770467	OR772786	OR772800
57,164	L. gilanensis	Η	Limpopo		3	Beaumont	2018	OR659664	OR770468	OR772787	OR772801
57,166	L. gilanensis	Н	Limpopo		3	Beaumont	2018	OR659665	OR770469	OR772788	OR772802
57,167	L. gilanensis	Н	Limpopo		1	Beaumont	2018	OR659666	OR770470	OR772789	OR772803
57,184	Lasiodiplodia sp.	D	Mpumalanga		1	Mixed int	2018	OR659667	OR770481	OR772900	OR773001
57,185	Lasiodiplodia sp.	D	Mpumalanga		1	Mixed int	2018	OR659677	OR770482	OR773670	OR773002
57,195	Lasiodiplodia sp.	Н	Mpumalanga		4	Nelmak D	2018	OR659678	OR770483	OR773671	OR773003
57,196	Lasiodiplodia sp.	Н	Mpumalanga		5	816	2018	OR659679	OR770484	OR773672	OR773004
57,187	Lasiodiplodia sp.	Н	KwaZulu-Natal		2	788	2018	OR659680	OR770486	OR773673	OR773005
57,191	Lasiodiplodia sp.	Н	KwaZulu-Natal		2	788	2018	OR659681	OR770487	OR773674	OR773006
57,192	Lasiodiplodia sp.	Н	Mpumalanga		4	Nelmak D	2018	OR659683	OR770488	OR773675	OR773007
57,201	Lasiodiplodia sp.	Η	Mpumalanga		4	344	2018	OR659682	OR770489	OR773676	OR773008
57,202	Lasiodiplodia sp.	Н	Mpumalanga		5	816	2018	OR659676	OR770470	OR773677	OR773009
57,206	Lasiodiplodia sp.	Н	KwaZulu-Natal		2	Beaumont	2018	OR671508	OR770362	OR770678	OR770299
57,200	Lasiodiplodia sp.	Н	Mpumalanga		1	788	2018	OR671509	OR770364	OR770679	OR770297
57,203	Lasiodiplodia sp.	Н	KwaZulu-Natal		6	Nelmak D	2018	OR772782	OR770355	OR671549	OR671674
57,199	Lasiodiplodia sp.	Η	KwaZulu-Natal		6	Nelmak D	2018	OR772783	OR770354	OR671547	OR671673
57,197	Lasiodiplodia sp.	Н	KwaZulu-Natal		2	Beaumont	2018	OR772784	OR770353	OR671548	OR671672
57,204	Lasiodiplodia sp.	Н	KwaZulu-Natal		2	788	2018	OR772786	OR770352	OR671546	OR671671
57,207	Lasiodiplodia sp.	Н	Mpumalanga		4	Nelmak D	2018	OR772787	OR770351	OR671543	OR671670
57,209	Lasiodiplodia sp.	Н	Mpumalanga		4	344	2018	OR772788	OR770350	OR671542	OR671669
57,309	L. pseudotheobromae	D	KwaZulu-Natal		2	Beaumont	2018	OR772796	OR680840	OR772794	OR772801
57,310	L. pseudotheobromae	D	KwaZulu-Natal		2	Beaumont	2019	OR772792	OR680841	OR770468	OR772802
57,302	L. pseudotheobromae	D	Limpopo		1	Mixed int	2018	OR772793	OR680842	OR649170	OR772803
57,321	L. pseudotheobromae	D	Limpopo		7	814	2018	OR772790	OR680844	OR649171	OR772804
57,319	L. pseudotheobromae	D	KwaZulu-Natal		2	Beaumont	2021	OR772794	OR680845	OR649172	OR772805
57,316	L. pseudotheobromae	D	KwaZulu-Natal		2	Beaumont	2020	OR772797	OR680846	OR649173	OR772806
57,327	L. pseudotheobromae	D	KwaZulu-Natal		2	Beaumont	2022	OR772798	OR680847	OR649174	OR772807
57,140	L. pseudotheobromae	D	Limpopo		7	814	2018	OR772799	OR680848	OR649182	OR772808
57,160	L. pseudotheobromae	D	Mpumalanga		8	Mixed int	2020	OR772800	OR680849	OR770468	OR772809
57,161	L. pseudotheobromae	D	Mpumalanga		8	Mixed int	2020	OR772801	OR680850	OR770441	OR772900
57,162	L. pseudotheobromae	D	Mpumalanga		9	Mixed int	2020	OR772802	OR680861	OR770442	OR680832
57,153	L. pseudotheobromae	D	KwaZulu-Natal		10	Mixed int	2019	OR772803	OR680862	OR770443	OR680833
57,154	L. pseudotheobromae	D	KwaZulu-Natal		10	Mixed int	2019	OR//2804	OR680863	OR//0444	OR680834
57,155	L. pseudotheobromae	D	KwaZulu-Natal		10	Mixed int	2019	OR//2805	OR680864	OR//0446	OR680832
57,156	L. pseudotheobromae	D	Kwa∠ulu-Natal		10	Mixed int	2019	OK/72806	UK680865	OK/70447	OK680833
57,157	L. pseudotheobromae	D	KwaZulu-Natal		11	Beaunont	2019	OR7/2807	OR680866	OR//0448	OR680815
57,158	L. pseudotheobromae	D	KwaZulu-Natal		11	Beaumont	2019	OR772808	OR680843	OR / /0449	OR680816
57,159	L. pseudotheobromae	D	KwaZulu-Natal		11	Beaumont	2019	OK//2809	OR680842	OK/72/93	OK680817
57,170	L. pseudotheobromae	ע יי	Kwa∠ulu-Natal		11	Beaumont	2019	OR772900	OR680841	OR772792	OR680818
57,311	L. pseudotheobromae	Н	Limpopo		12	Mixed int	2018	OR//2901	OK680840	OR/72/91	OK680819
37,314	L. pseudotheobromae	н	ситроро		12	wiixed int	2018	OK//2902	OK680839	UK//0490	0K680820

# Table 1 (continued)

57,306       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR770371       OR770339       OR80821         57,304       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR859669       OR770370       OR770339       OR860822         57,307       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR859670       OR770369       OR770334       OR860824         57,307       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR859670       OR770360       OR770334       OR860825         57,301       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR85975       OR77036       OR770331       OR860825         57,301       L pseudotheobromae       H Limpopo       12       Mixed int       2018       OR85975       OR77036       OR770331       OR860825         57,305       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770361       OR770330       OR860825         57,315       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770460       OR770470<
57,304       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659669       OR770370       OR770330       OR680823         57,307       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659670       OR770360       OR770330       OR680823         57,307       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659670       OR770360       OR770330       OR680823         57,301       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659670       OR770360       OR770330       OR680829         57,313       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770360       OR770330       OR680829         57,131       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770460       OR770340       OR680829         57,132       L pseudotheobromae       H       Kupunalanga       4       Nelmak       2018       OR659675       OR770460       OR77048       OR680833         57,132       L pseudotheobromae       H       Kupunalanga <t< td=""></t<>
57,308       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659669       OR770369       OR770330       OR680824         57,307       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659671       OR770360       OR770334       OR680824         57,301       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659671       OR770360       OR770330       OR680825         57,301       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770360       OR770330       OR680829         57,313       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770460       OR770480       OR680830       OR770449       OR680831         57,152       L pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659676       OR770460       OR770440       OR680831         57,152       L pseudotheobromae       H       Kimpopo       12       Mixed int       2018       OR659676       OR770460       OR770440       OR680831         57,134       L theobromae <td< td=""></td<>
57,307       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659671       OR770368       OR770334       OR680824         57,303       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659671       OR770360       OR770334       OR680825         57,300       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770360       OR770331       OR680828         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770360       OR770330       OR680828         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR7040       OR770440       OR680830         57,154       L pseudotheobromae       H       Kamzulu-Natal       13       788       2018       OR659675       OR770460       OR770440       OR680835         57,134       L theobromae       D       Limpopo       1       Mixed int       2018       OR659651       OR770460       OR770480       OR770480       OR70483       OR680835         57,135       L theobromae       H
57,303       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659671       OR70367       OR70332       OR680825         57,301       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659674       OR70366       OR70332       OR680825         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR70360       OR70332       OR680825         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR70360       OR70480       OR70480       OR680829         57,152       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR70460       OR70440       OR680825         57,152       L. pseudotheobromae       H       KwaZulu-Natal       13       788       2018       OR659651       OR70460       OR70470       OR680832         57,130       L. theobromae       D       Mupumalanga       4       344       2018       OR659651       OR70460       OR70480       OR680830         57,132       L. theobromae       H       Mpumalanga
57,301       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659674       OR770366       OR770333       OR680826         57,300       L. pseudotheobromae       H       Mpumalanga       4       Nelmak D       2018       OR659675       OR770364       OR770332       OR680827         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770364       OR770330       OR680829         57,150       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770360       OR770448       OR680830         57,152       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659657       OR770469       OR770449       OR680831         57,130       L. theobromae       D       Limpopo       1       Mixed int       2018       OR659651       OR770467       OR770481       OR680833         57,132       L. theobromae       D       Mpumalanga       4       344       2018       OR659653       OR770464       OR770483       OR680836         57,132       L. theobromae       H       Mpumalanga       4
57,300       L. pseudotheobromae       H       Mpumalanga       4       Nelmak D       2018       OR659675       OR770364       OR770332       OR680827         57,312       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659672       OR770363       OR770330       OR680828         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770360       OR770330       OR680829         57,150       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR7049       OR70449       OR680831         57,150       L. pseudotheobromae       H       KwaZulu-Natal       13       788       2018       OR659651       OR7046       OR70449       OR680835         57,130       L. theobromae       D       Mpumalanga       4       344       2018       OR659651       OR70460       OR70480       OR680836         57,132       L. theobromae       H       Mpumalanga       4       Nelmak D       2018       OR659651       OR70460       OR70480       OR680836         57,132       L. theobromae       H       Mpumalanga       S       816
57,312       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659672       OR770363       OR770313       OR680828         57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770362       OR770330       OR680829         57,159       L. pseudotheobromae       H       Mumalanga       4       Nelmak D       2018       OR659673       OR770360       OR770449       OR680831         57,150       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR770469       OR770449       OR680835         57,130       L. theobromae       D       Limpopo       1       Mixed int       2018       OR659651       OR77046       OR770481       OR680833         57,132       L. theobromae       D       Limpopo       7       Mixed int       2018       OR659653       OR770460       OR770482       OR680836         57,131       L. theobromae       H       Mpumalanga       4       Nelmak D       218       OR659654       OR71046       OR77048       OR680835         57,135       L. theobromae       H       Mpumalanga       5       81
57,313       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659673       OR770362       OR770300       OR680829         57,169       L. pseudotheobromae       H       Mpumalanga       4       Nelmak D       2018       OR659675       OR70469       OR770448       OR680830         57,152       L. pseudotheobromae       H       Limpopo       12       Mixed int       2018       OR659675       OR70469       OR770449       OR680833         57,130       L. theobromae       D       Limpopo       1       Mixed int       2018       OR659651       OR770460       OR770481       OR680833         57,130       L. theobromae       D       Limpopo       1       Mixed int       2018       OR659653       OR770460       OR770481       OR680833         57,130       L. theobromae       D       Limpopo       7       Mixed int       2018       OR659653       OR770460       OR770482       OR680834         57,131       L. theobromae       H       Mpumalanga       4       Nelmak D       2018       OR59656       OR70463       OR770468       OR70483       OR680840         57,132       L. theobromae       H       Mpumalanga       4<
57,169         L. pseudotheobromae         H         Mpumalanga         4         Nelmak D         2018         OR659676         OR680843         OR770448         OR680830           57,152         L. pseudotheobromae         H         Limpopo         12         Mixed int         2018         OR659657         OR770469         OR770449         OR680831           57,130         L. theobromae         D         Limpopo         1         Mixed int         2018         OR659651         OR770460         OR770481         OR680833           57,130         L. theobromae         D         Mpumalanga         4         344         2018         OR659651         OR770460         OR770481         OR680833           57,130         L. theobromae         D         Mpumalanga         4         344         2018         OR659653         OR70460         OR770481         OR680830           57,131         L. theobromae         H         Mpumalanga         5         816         2018         OR659656         OR770460         OR770480         OR680830           57,135         L. theobromae         H         Mpumalanga         4         Nelmak D         2018         OR659656         OR770460         OR770480         OR680831 <tr< td=""></tr<>
57,152L pseudotheobromaeHLimpopo12Mixed int2018OR65967OR770469OR770449OR68083157,138L pseudotheobromaeHKwaZulu-Natal137882018OR65967OR770460OR770461OR68083557,130L theobromaeDLimpopo1Mixed int2018OR659651OR770460OR770481OR68083257,134L theobromaeDMpumalanga43442018OR659653OR770460OR770483OR68083457,132L theobromaeHMpumalanga4Nelmak D2018OR659654OR70460OR770480OR68084057,131L theobromaeHMpumalanga58162018OR70468OR770460OR770480OR68083057,134L theobromaeHMpumalanga58162018OR659656OR70461OR70480OR68083057,135L theobromaeHKwaZulu-Natal2Beaumont2018OR659669OR70460OR70480OR68083157,135L theobromaeHMpumalanga4Nelmak D2018OR659669OR70460OR70480OR68083157,135L theobromaeHMpumalanga4Nelmak D2018OR649170OR680821OR70480OR68083157,135L theobromaeHKwaZulu-Natal2Beaumont2018OR649171OR680823OR70460OR70480OR68081557,116N.kwambona
57,138L pseudotheobromaeHKwaZulu-Natal137882018OR65967OR770468OR770470OR68083557,130L theobromaeDLimopopo1Mixed int2018OR659651OR770467OR770481OR68083257,134L theobromaeDMpumalanga43442018OR659653OR770460OR770483OR68083357,129L theobromaeDLimopopo7Mixed int2018OR659653OR770460OR770484OR68083457,131L theobromaeHMpumalanga4Nelmak D2018OR659654OR770460OR770480OR68084057,131L theobromaeHMpumalanga58162018OR659656OR770460OR770480OR68083057,132L theobromaeHKwaZulu-Natal2Beaumon2018OR659656OR770460OR770480OR68083057,133L theobromaeHMpumalanga4Nelmak D2018OR659659OR770460OR770480OR68083057,134L theobromaeHMpumalanga4Nelmak D2018OR659659OR770460OR770480OR68083057,135L theobromaeHMpumalanga2Beaumon2018OR649170OR68021OR770340OR68081557,114N.kwambonambienseDLimopo2Beaumon2018OR649171OR68023OR770340OR68081657,116N.kwambonambiense
57,130L. theobromaeDLimpopo1Mixed int2018OR659651OR770467OR770481OR68083257,134L. theobromaeDMpumalanga43442018OR659652OR770466OR770482OR68083357,129L. theobromaeDLimpopo7Mixed int2018OR659653OR770466OR770484OR68083457,132L. theobromaeHMpumalanga4Nelmak D2018OR659654OR770464OR770484OR68084057,131L. theobromaeHMpumalanga58162018OR659656OR770463OR770483OR68083057,132L. theobromaeHLimpopo1Mixed int2018OR659666OR770460OR770484OR68083057,133L. theobromaeHKwaZulu-Natal2Beaumont2018OR659669OR770460OR770489OR68083257,133L. theobromaeHMpumalanga4Nelmak D2018OR659669OR770460OR770489OR68083257,135L. theobromaeHMpumalanga4Nelmak D2018OR659669OR770460OR770489OR68083257,135L. theobromaeHMpumalanga2Beaumont2018OR649170OR680821OR770489OR68082157,135L. theobromaeDLimpopo2Beaumont2018OR649170OR680821OR770340OR6808157,116N. kwambonambienes<
57,134L. theobromaeDMurualanga43442018OR659652OR770466OR770482OR68083357,129L. theobromaeDLimpopo7Mixed int2018OR659653OR770465OR770483OR68083457,132L. theobromaeHMpumalanga4Nelmak D2018OR659654OR770464OR770484OR68083657,131L. theobromaeHMpumalanga58162018OR659656OR770463OR770480OR68083057,132L. theobromaeHLimpopo1Mixed int2018OR659666OR770461OR770480OR68083057,133L. theobromaeHKwaZulu-Natal2Beaumont2018OR659669OR770460OR770480OR68083157,133L. theobromaeHMpumalanga4Nelmak D2018OR659669OR770460OR770480OR68083257,135L. theobromaeHKwaZulu-Natal2Beaumont2018OR649170OR680821OR770480OR68083257,112N. kwambonambienseDLimpopo2Beaumont2018OR649171OR680821OR770440OR68081657,114N. kwambonambienseDLimpopo1Beaumont2018OR649173OR680821OR770440OR68081757,115N. kwambonambienseDLimpopo2Beaumont2018OR649173OR680821OR770440OR68081957,113N. k
57,129L. theobromaeDLimpopo7Mixed int2018OR659653OR70465OR770463OR68083457,132L. theobromaeHMpumalanga4Nelmak D2018OR659654OR70463OR770463OR70486OR68084057,131L. theobromaeHMpumalanga58162018OR659656OR770463OR770463OR70486OR68084057,132L. theobromaeHLimpopo1Mixed int2018OR659656OR770460OR770480OR68083057,133L. theobromaeHKwaZulu-Natal2Beaumont2018OR659669OR770460OR770480OR68083257,135L. theobromaeHKwaZulu-Natal2Beaumont2018OR659669OR770460OR770480OR68083257,135L. theobromaeHKwaZulu-Natal2Beaumont2018OR69170OR680821OR770468OR68083257,135L. theobromaeDLimpopo2Beaumont2018OR649170OR680821OR770468OR68081557,114N. kwambonambienseDLimpopo2Beaumont2018OR649171OR680821OR770446OR68081657,116N. kwambonambienseDLimpopo14Beaumont2018OR649171OR680821OR770446OR68081757,115N. kwambonambienseDLimpopo1Beaumont2018OR649171OR680822OR770440OR680816 </td
57,132       L theobromae       H       Mumalanga       4       Nelmak D       2018       OR659654       OR770464       OR770484       OR680836         57,131       L theobromae       H       Mpumalanga       5       816       2018       OR70468       OR770463       OR770486       OR680836         57,132       L theobromae       H       Limpopo       1       Mixed int       2018       OR659656       OR770461       OR770488       OR680830         57,133       L theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR659669       OR770460       OR770489       OR680830         57,133       L theobromae       H       Mpumalanga       4       Nelmak D       2018       OR659669       OR770460       OR770489       OR680835         57,133       L theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR649170       OR680821       OR770468       OR680815         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR770446       OR680817         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont
57,131       L. theobromae       H       Mpumalanga       5       816       2018       OR770468       OR770463       OR770486       OR680840         57,128       L. theobromae       H       Limpopo       1       Mixed int       2018       OR659656       OR770461       OR770488       OR680830         57,136       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR659668       OR770460       OR770489       OR680830         57,133       L. theobromae       H       Mpumalanga       4       Nelmak D       2018       OR659669       OR770460       OR770488       OR680832         57,135       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR649170       OR680821       OR770468       OR870348       OR680832         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR770468       OR680815         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649172       OR680821       OR770346       OR680817         57,113       N. kwambonambiense       D       Limpopo       14
57,128       L. theobromae       H       Limpopo       1       Mixed int       2018       OR659656       OR770462       OR770487       OR680830         57,136       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR659668       OR770461       OR770487       OR680830         57,133       L. theobromae       H       Mpumalanga       4       Nelmak D       2018       OR659669       OR770460       OR770488       OR680832         57,135       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR649170       OR680821       OR770468       OR770488       OR680832         57,112       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR7047048       OR680815         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649171       OR680821       OR770450       OR680816         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649173       OR680821       OR770345       OR680819         57,113       N. kwambonambiense       D       Limpopo
57,136       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR659668       OR770461       OR770488       OR680831         57,133       L. theobromae       H       Mpumalanga       4       Nelmak D       2018       OR659669       OR770460       OR770489       OR680835         57,135       L. theobromae       H       KwaZulu-Natal       2       Beaumont       2018       OR649170       OR680821       OR770468       OR680832         57,112       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR770348       OR680815         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680822       OR770346       OR680816         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649172       OR680823       OR770346       OR680817         57,118       N. kwambonambiense       D       Limpopo       14       Beaumont       2018       OR649173       OR680821       OR770344       OR680827         57,113       N. kwambonambiense       D       Limpopo       1       <
57,133L. theobromaeHMpumalanga4Nelmak D2018OR659669OR770460OR770489OR68083557,135L. theobromaeHKwaZulu-Natal2Beaumont2018OR770468OR770459OR770468OR68083257,112N. kwambonambienseDLimpopo2Beaumont2018OR649170OR680821OR770348OR68081557,114N. kwambonambienseDLimpopo2Beaumont2018OR649171OR680822OR770347OR68081657,116N. kwambonambienseDLimpopo2Beaumont2018OR649172OR680823OR770346OR68081757,118N. kwambonambienseDLimpopo14Beaumont2018OR649173OR680824OR770345OR68081957,113N. kwambonambienseDLimpopo2Beaumont2018OR649174OR680821OR770344OR68081957,113N. kwambonambienseDLimpopo1Beaumont2018OR649174OR680822OR770343OR68082757,123N. kwambonambienseDLimpopo15Beaumont2018OR649175OR770330OR770342OR68082057,120N. kwambonambienseHLimpopo14Beaumont2018OR649175OR77031OR770340OR68082057,117N. kwambonambienseHLimpopo14Beaumont2018OR649175OR770331OR770340OR680820<
57,135L. theobromaeHKwaZulu-Natal2Beaumont2018OR770468OR770459OR770468OR68083257,112N. kwambonambienseDLimpopo2Beaumont2018OR649170OR680821OR770348OR68081557,114N. kwambonambienseDLimpopo2Beaumont2018OR649171OR680822OR770347OR68081657,116N. kwambonambienseDLimpopo2Beaumont2018OR649172OR680823OR770346OR68081757,118N. kwambonambienseDLimpopo14Beaumont2018OR649173OR680824OR770345OR68081857,113N. kwambonambienseDLimpopo2Beaumont2018OR649174OR680821OR770344OR68081957,115N. kwambonambienseDLimpopo1Beaumont2017OR649182OR770330OR770342OR68082757,123N. kwambonambienseDLimpopo15Beaumont2018OR649175OR770331OR770342OR68082057,120N. kwambonambienseHLimpopo14Beaumont2018OR649175OR770331OR770340OR68082057,117N. kwambonambienseHLimpopo14Beaumont2018OR649175OR770331OR770341OR68082057,117N. kwambonambienseHLimpopo14Beaumont2018OR649175OR770331OR770340OR680820 </td
57,112       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR770348       OR680815         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649170       OR680821       OR770348       OR680816         57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649171       OR680822       OR770346       OR680816         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649172       OR680823       OR770346       OR680817         57,118       N. kwambonambiense       D       Limpopo       14       Beaumont       2018       OR649173       OR680824       OR770345       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649174       OR680821       OR770344       OR680819         57,113       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR770340       OR680827         57,123       N. kwambonambiense       D       Limpopo       15       Beaumont
57,114       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649171       OR680822       OR770347       OR680816         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649171       OR680823       OR770347       OR680816         57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649172       OR680823       OR770346       OR680817         57,118       N. kwambonambiense       D       Limpopo       14       Beaumont       2018       OR649173       OR680824       OR770345       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649174       OR680821       OR770344       OR680819         57,113       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR770343       OR680827         57,115       N. kwambonambiense       D       Limpopo       15       Beaumont       2010       OR649181       OR770330       OR770342       OR680826         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont
57,116       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649172       OR680823       OR770346       OR680817         57,118       N. kwambonambiense       D       Limpopo       14       Beaumont       2018       OR649173       OR680824       OR770346       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649173       OR680824       OR770345       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649174       OR680821       OR770344       OR680819         57,115       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR680822       OR770343       OR680827         57,123       N. kwambonambiense       D       Limpopo       15       Beaumont       2020       OR649181       OR770330       OR770342       OR680820         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14
57,118       N. kwambonambiense       D       Limpopo       14       Beaumont       2018       OR649173       OR680824       OR770345       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649173       OR680824       OR770345       OR680818         57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649174       OR680821       OR770344       OR680819         57,115       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR680822       OR770343       OR680827         57,123       N. kwambonambiense       D       Limpopo       15       Beaumont       2020       OR649181       OR770330       OR770342       OR680820         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770332       OR770340       OR680820         57,117       N. kwambonambiense       H       Limpopo       14
57,113       N. kwambonambiense       D       Limpopo       2       Beaumont       2018       OR649174       OR680821       OR770344       OR680819         57,115       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649174       OR680821       OR770344       OR680827         57,115       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR680822       OR770342       OR680826         57,123       N. kwambonambiense       D       Limpopo       15       Beaumont       2020       OR649181       OR770330       OR770342       OR680826         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649176       OR770332       OR770340       OR680821
57,115       N. kwambonambiense       D       Limpopo       1       Beaumont       2017       OR649182       OR680822       OR770343       OR680827         57,123       N. kwambonambiense       D       Limpopo       15       Beaumont       2020       OR649181       OR770330       OR770342       OR680826         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649176       OR770332       OR770340       OR680820
57,123       N. kwambonambiense       D       Limpopo       15       Beaumont       2020       OR649181       OR770330       OR770342       OR680826         57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649176       OR770332       OR770340       OR680820
57,120       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649175       OR770331       OR770341       OR680820         57,117       N. kwambonambiense       H       Limpopo       14       Beaumont       2018       OR649176       OR770332       OR770340       OR680821
57,117         N. kwambonambiense         H         Limpopo         14         Beaumont         2018         OR649176         OR770332         OR770340         OR680821
I I
57.121 N. kwambonambiense H KwaZulu-Natal 13 Beaumont 2018 OR649177 OR770334 OR770339 OR680822
57,119 <i>N. kwambonambiense</i> H Limpopo 14 Beaumont 2018 OR649178 OR770335 OR770338 OR680823
57.124 N. kwambonambiense H Limpopo 2 Mixed int 2018 OR649179 OR770328 OR770337 OR680824
57.122 N. kwambonambiense H Limpopo 2 Mixed int 2018 OR649180 OR770327 OR770336 OR680825
57.125 N. luteum D KwaZulu-Natal 16 Mixed int 2020 OR649183 OR770326 OR770468 OR680843
57.126 <i>N. luteum</i> D KwaZulu-Natal 16 Mixed int 2020 OR649184 OR770325 OR770469
57.127 N. luteum D Mpumalanga 9 Mixed int 2020 OR649185 OR770324 OR770470 OR680844
<b>57,140</b> <i>N. parvum</i> D Limpopo 18 Mixed int 2020 OR649168 OR770323 OR770481 OR680829
57.145 <i>N. parvum</i> D KwaZulu-Natal 16 Mixed int 2020 OR649167 OR770322 OR770482 OR680830
57.146 <i>N. parvum</i> D. Limpopo 17 Beaumont 2019 OR649166 OR770321 OR770483 OR680831
57.149 <i>N. parvum</i> D Limpopo 17 Beaumont 2019 OR649162 OR770320 OR770484 OR680835
57.143 <i>N. parvum</i> D. Limpopo 17 Beaumont 2019 OR649165 OR770319 OR770486 OR680832
57.144 <i>N. parvum</i> D. Limpopo 17 Beaumont 2019 OR649164 OR770318 OR770487 OR680833
57.142 <i>N. parvum</i> D Limpopo 17 Beaumont 2019 OR649163 OR772801 OR770488 OR680834
57,150 <i>N. parvum</i> H Limpopo 2 Beaumont 2018 OR649161 OR772802 OR770489 OR680836
57.155 <i>N. parvum</i> H Limpopo 2 Mixed int 2018 OR649157 OR772803 OR770468 OR680840
57.152 <i>N. parvum</i> H Limpopo 2 Beaumont 2018 OR649160 OR772804 OR770469 OR680837
57.154 <i>N. parvum</i> H Limpopo 2 Mixed int 2018 OR649158 OR772805 OR770470 OR680839
57,158 <i>N. parvum</i> H Limpopo 2 Beaumont 2018 OR649159 OR772806 OR770481 OR680838
57,156 <i>N. parvum</i> H Limpopo 2 Beaumont 2018 OR649156 OR772807 OR770482 OR680841

Table 1 (continued)

CMW 57,151	Species	Tissue <sup>3</sup>	Province	Farm	Cultivar Beaumont	Date 2018	GenBank Accession number			
	N. parvum	H Limpopo					OR649155	OR772808	OR770483	OR680842
57,141	N. parvum	H Limpopo		2	Mixed int	2018	OR649169	OR772809	OR770484	OR680828

<sup>1</sup>ID numbers for isolates used in pathogenicity trials in this study are given in bold

<sup>2</sup>Isolates shown to hybridize are given in italics

<sup>3</sup>Isolates obtained from asymptomatic tissues = H, and from symptomatic tissues = D

<sup>4</sup>Mixed Int=Mixed integrifolia where the correct identification of the cultivar was not clear or unknown

room temperature for seven days prior to inoculation. Oneyear-old trees of the Beaumont cultivar, planted in black plastic bags, were left to acclimatize in the glasshouse for 3 weeks prior to inoculation. Glasshouse conditions were set with daily temperature fluctuations (approximately 25 °C Day/15 °C night), and a controlled photoperiod of 16/8 h (day/night).

Ten inoculations per isolate were performed at the base of the stem of each tree by firstly removing the bark tissue with a 5 mm cork borer to expose the cambium. Thereafter a 5 mm diameter mycelial plug, cut from the actively growing margin of seven-day-old culture, were placed mycelium surface down into the wounded area with control inoculations that were made using a clean MEA plug. The wounds and mycelial plugs were then sealed with Parafilm (Bemis®, USA) to prevent desiccation and contamination. The inoculated trees were randomly arranged in the glasshouse. Following inoculation, trees were watered every two days for the first 9 weeks. After the ninth week, trees were watered every fourth day. Measurements of lesions were taken when the first dieback symptoms were observed on the trees 12 weeks after inoculation. Data visualization and statistical analyses for all data were carried out in R software (RStudio Team, 2022). The Kruskal-Wallis test was used to identify the significant difference (P < 0.05) between species used in the pathogenicity trials in this study.

Re-isolations were made from two trees inoculated with each of the *Botryosphaeriaceae* isolates and the controls. This was done by cutting out a small piece of wood at the edge of the lesion and placing this on 2% MEA. DNA sequences of the *tef-1* $\alpha$  gene region were used to confirm the identity of the fungal isolates obtained.

# Results

#### **Fungal Isolates**

A total number of 138 *Botryosphaeriaceae* isolates were obtained from symptomatic and asymptomatic branches. We therefore isolated 92 *Botryosphaeriaceae* isolates from 120 asymptomatic branches. Of the 46 isolates obtained from

symptomatic tissues, 27 were obtained from disease clinic samples and 15 were from samples collected from the survey of the three regions. With regards to geographical location, 58 isolates were obtained from Limpopo, 33 isolates from Mpumalanga, and 47 isolates from KwaZulu-Natal. BLASTn results revealed that the isolates obtained belonged to three genera, namely *Diplodia* (two isolates) *Lasiodiplo-dia* (88 isolates) and *Neofusicoccum* (48 isolates).

## **Identity of isolates**

In the phylogenetic analysis the two Diplodia isolates grouped in a clade with D. alatafructa, D. pseudoseriata and D. insularis (Fig. 1). The two isolates were identical to the type strains of D. pseudoseriata (EU080927) and D. alatafructa (FJ888460) in the ITS dataset (Online Resource 1a). However, the two isolates formed a sub-clade within a larger clade in both the *tub2* (Online Resource 1b) the *tef-1* $\alpha$  (Online Resource 1c) phylogenies with a strong bootstrap support. In the combined phylogeny, the two isolates grouped in a clade with D. alatafructa, D. pseidoseriata and D. insularis, in agreement to the grouping observed in the ITS phylogeny (Fig. 1). The two isolates in the present study differed from D. pseudoseriata with three and two nucleotides on *tef-1* $\alpha$  and *tub2*, respectively, and consequently could not be assigned to a known species and are treated as Diplodia sp.

The phylogenetic analyses of ITS, tub2,  $tef-1\alpha$ , rpb2 and combined sequence datasets of the 31 representative isolates of *Neofusicoccum* resulted in trees with similar topologies (Fig. 2, Online resource 2). The phylogenies grouped the isolates into three clades representing *N. parvum* (n=15), *N. kwambonambiense* (n=13), and a clade containing *N. luteum* and *N. mangroviorum* (n=3) (Fig. 2). Based on our phylogenetic analysis, the latter two species could not be confidently separated from one another, which agrees with Zhang et al. (2021).

A total of 60 *Lasiodiplodia* representative isolates were selected for further analysis, however only a few representatives are shown in the individual and combined phylogenies. Phylogenetic analyses of the representative isolates based on both the ITS and *tub2* datasets lacked **Fig. 1** Maximum Likelihood (ML) tree of the genus *Diplodia* produced with from a combined ITS, *tub2*, and *tef-1a* sequences of *Diplodia*. Bootstrap values above 70% are given at the nodes. Isolates sequenced in this study appear in bold and type species are in blue. The tree was rooted to *Botryosphaeria dothidea* (CBS115476 and CBS110302)



phylogenetic resolution to differentiate between all the known *Lasiodiplodia* species (Online Resources 3a, b). Eight of the isolates grouped closely but separate from *L. gonubiensis*, while the remaining *Lasiodiplodia* isolates grouped within a larger clade representing the *L. theobromae* species complex (Online Resources 3a, b). The *tef-la* and *rbp2* phylogenies (Online Resources 3c, d) had similar topologies and could distinguish between

most *Lasiodiplodia* species in this study. In both phylogenies, nine isolates grouped with *L. gilanensis* and *L. missouriana* (syn. *L. gilanensis*; Zhang et al. 2021) and another eight isolates grouped with *L. pseudotheobromae*. The combined dataset of the four gene regions (Fig. 3) confirmed the grouping of isolates from this study in *L. pseudotheobromae*, *L. theobromae*, *L. gilanensis* and *Lasiodiplodia* sp.

**Fig. 2** Maximum Likelihood (ML) tree of *Neofusicoccum* produced with a combined sequence data of ITS, *tub2*, *rpb2* and *tef-1a*. Bootstrap values above 70% are given at the nodes. Isolates sequenced in this study appear in bold and type species are in blue. The tree was rooted to *Botryosphaeria dothidea* (CBS115476 and CBS110302)



Three of the isolates obtained in this study (CMW57135, CMW57129 and CMW57133) displayed incongruency between different gene regions and might represent hybrids. In ITS and *tub2* (Online Resources 3a, b), all three isolates grouped within the *L. theobromae* complex. However, for

*tef-1a* (Online Resource 3c), CMW57133 grouped with *L. pseudotheobromae* and with *L. theobromae* in the *rpb2* dataset. Isolate CMW57129 grouped separately, but close to *L. theobromae* in *tef-1a*, but grouped with *L. mahajangana* (syn. *L. exigua*) in the *rpb2* analysis (Online Resource 3d)

**Fig. 3** Maximum Likelihood (ML) tree of *Lasiodiplodia* produced with a combined sequence data of *ITS, tub2, rpb2* and tef-1 $\alpha$ . Bootstrap values above 70% are given at the nodes. Isolates sequenced in this study appear in bold and type species are in blue. Potential hybrid isolates are given in red and isolates from Australia are in green. The tree was rooted to *Botryosphaeria dothidea* (CBS115476 and CBS110302)





and in the combined dataset (Fig. 3). Isolate CMW57135 grouped with *L. theobromae* in all individual datasets other than for *tef-1a* where it grouped close to both *L. pseudotheobromae* and *L. lignicola*. Based on the decision tree provided by Cruywagen et al. (2017), all three isolates were thus treated as hybrids because they move between nonsister species in phylogenetic analyses.

# Species distribution per region

Botryosphaeriaceae species were recovered from all three major Macadamia growing regions in South Africa, however the species diversity and distribution differed between regions. Seven species was found in KwaZulu-Natal, with six species present in Limpopo and four in Mpumalanga (Fig. 4). One putative hybrid species was also detected in each growing region. Lasiodiplodia pseudotheobromae was the most frequently isolated species (n = 33), followed by Lasiodiplodia theobromae (n = 14) from all three Macadamia production regions. Lasiodiplodia sp. (n = 29) was isolated in relative high numbers from Mpumalanga and KwaZulu-Natal while N. parvum (n = 24) was dominant in Limpopo with only one isolate obtained from KwaZulu-Natal. Neofusicoccum kwambonambiense (n = 21), N. *luteum* (n=3) and *L. gilanesis* (n=9) were obtained from both KwaZulu-Natal and Limpopo. The *Diplodia* sp. were

**Fig. 4** *Botryosphaeriaceae* species diversity and distribution on Macadamia across three the major growing regions in South Africa

present in very low numbers and were only isolated from a single orchard in Limpopo (Fig. 4).

# Isolates from asymptomatic versus symptomatic trees

A total of 92 isolates isolates were obtained from 120 asymptomatic tissues, while all 46 symptomatic tissues yielded 46 isolates. Six of the eight species (N. kwambonambiense N. parvum, Lasiodiplodia sp. L. pseudotheobromae, L. gilanensis, and L. theobromae) identified in this study were present on both tissue types (Fig. 5). Lasiodiplodia pseudotheobromae was the most common and dominant species in both tissue types. However, it was isolated in higher numbers from diseased tissues (n = 19) compared to healthy tissues (n = 14). Diplodia sp. was obtained from only asymptomatic tissues but was isolated in only low numbers (n=2) and from a single orchard. *Neofusicoccum luteum* was also represented in low numbers (n = 3)and was isolated only from diseased samples submitted to the diagnostic clinic. Of the three *Lasiodiplodia* hybrids, two were isolated from asymptomatic tissues (CMW57133 and CMW57135) while CMW 57129 was isolated from a symptomatic tissue.



**Fig. 5** Venn diagram showing species overlap and diversity of species obtained from both asymptomatic and symptomatic tissues



# **Pathogenicity trials**

For the first nine weeks, the inoculated plants showed no visible leaf symptoms and no stem discoloration (necrotic lesion) (Fig. 6a). Lesions and branch dieback symptoms were however observed 12 weeks post inoculation (Fig. 6b, c, d), three weeks after watering of plants was reduced. All the *Botryosphaeriaceae* isolates tested in this study produced lesions after wounding and are therefore considered pathogenic to Macadamia. Significant (P < 0.001) differences in aggressiveness were observed among the species. *Neofusicoccum luteum*, *N. kwambonambiense* and *Lasiodiplodia* sp. produced the largest lesions and their lesions differed significantly from those of the controls (Fig. 7). *Lasiodiploidia gilanensis, L. pseudotheobromae* and *L. theobromae* produced the smallest lesions of all tested isolates.

# Discussion

This study represents the first comprehensive investigation of the diversity, distribution, and the pathogenicity of *Botryosphaeriaceae* species associated with macadamia branches in South Africa. Isolates obtained from different growing regions were analysed, resulting in the identification of eight species and three putative hybrids residing in *Diplodia*, *Neofusicoccum* and *Lasiodiplodia*. These included a *Diplodia* sp., three *Neofusicoccum* species and four *Lasiodiplodia* species. In addition, three putative *Lasiodiplodia* hybrids were also identified. The pathogenicity trials demonstrated that all *Botryosphaeriaceae* species linked to Macadamia in South Africa can induce vascular discoloration and/or dieback in Macadamia seedlings, although symptoms only appeared after a significant reduction in irrigation.

This study revealed a high diversity of Botryosphaeriaceae species on Macadamia in South Africa. This in agreement with previous studies from other woody hosts (McDonald and Eskalen 2011; Jami et al. 2013; Carlucci et al. 2015; Scarlett et al. 2019; Hilário et al. 2020). A total of thirteen species were also obtained from surveys conducted on Macadamia in Australia (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). The results obtained from this study are therefore in agreement with the surveys from Australia that revealed a high diversity of species present on this host in both its native and non-native range. However, in terms of species overlap, only L. pseudotheobromae, L. theobromae, N. parvum and N. luteum (syn. N. mangroviorum) were isolated from Macadamia in both countries. These shared species are known to have a cosmopolitan distribution associated with various hosts including wood, fruit and nuts crops (Slippers and Wingfield 2007; Trouillas et al. 2010; Hui-Fang et al. 2012; Chen et al. 2014; Pavlic-Zupanc et al. 2017; Burgess et al. 2019; Moral et al. 2019). It is, therefore, not surprising that they were isolated from Macadamia in both countries.

The largest group of isolates belonged to *Lasiodiplodia*. Species in the genus are found on a wide range of hosts including nut and fruits trees globally (Coutinho et al. 2017; Moral et al. 2019). *Lasiodiplodia pseudotheobromae* accounted for the highest percentage of isolates obtained in this study. This species was found in all the growing regions sampled and was also the dominant species isolated from Macadamia in Australia (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). *Lasiodiplodia gilanensis* is reported on Macadamia for the first time but was present in low numbers. This species was first described from Iran from twigs of an unknown woody plant (Abdollahzadeh et al. 2010). However, it has since been reported in China **Fig. 6** Macadamia seedlings pathogenicity trials. (A) Seedlings nine weeks post inoculation forming a callus around the point of inoculation; (B) Dieback symptoms observed on leaves just below the point of inoculation. (C) Seedlings twelve weeks post inoculation showing typical branch dieback symptoms. (D) The difference in lesion length between the eight species used plus the control



(Li et al. 2018), Mexico (Rangel-Montoya et al. 2021) and the United States (Úrbez-Torres et al. 2017). In South Africa, *L. gilanensis* was first detected in KwaZulu-Natal on native *Syzygium cordatum* (Vivas et al. 2021). It is, therefore, possible that the species has moved between native *S. cordatum* and non-native Macadamia since they are often planted in close proximity to natural environments.

Hybridisation has been reported in a variety of important fungal pathogens. The present work reports three hybrids in *Lasiodiplodia*, which is currently the only genus in the *Botryosphaeriaceae* where hybridization has been described (Sakalidis et al. 2011; Cruywagen et al. 2017; Rodríguez-Gálvez et al. 2017). It is possible that some hybrids in other genera have been overlooked due to the reliance on ITS and *tef-1* $\alpha$  sequence data, as the identification of hybrids requires consideration of incongruence between multiple loci (Cruywagen et al. 2017). Hybridization events leads to the generation of unique genetic diversity that could result in speciation events, host range expansions or other ecological changes (Brasier and Kirk 2001). The presence and frequency of such hybrids should therefore be studied in more detail in future.

*Neofusicoccum* was the second most dominant genus present in this study. Out of the three species found in this



**Fig. 7** Necrotic lesion length (mm) caused by isolates of *Botryosphaeriaceae* on one-year-old Beaumont plants. Bars with same letter are not significantly different from each other

study, *N. parvum* was the most common across the different regions in this study and in Australia (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). In contrast, *N. luteum* was the least frequently isolated in Australia (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022) and in the current study. The second most prevalent species in this genus was *N. kwambonambiense*, which has been reported on various hosts in countries such as Brazil, South Africa, Uruguay and Australia (Pérez et al. 2008; Pavlic et al. 2009; Lopes et al. 2016; Liddle et al. 2019). However, despite its occurrence on other hosts in Australia, *N. kwambonambiense* has not previously been isolated from Macadamia.

This study is the first report of the genus *Diplodia* on Macadamia. Currently, the genus comprises of 25 species based on morphology and DNA sequence data of the ITS, *tef-1a*, and *tub2* gene regions (Dissanayake et al. 2016; Slippers et al. 2017; Yang et al. 2017; Zhang et al. 2021). The two isolates (CMW46906, CMW48899) obtained in the current study, however, could not be assigned to a specific species due to the lack of bootstrap support to confidently separate *D. insularis*, *D. alatafructa*, and *D. pseudoseriata* from one another. Our results are therefore in agreement to Zhang et al. (2021) and Phillips et al. (2012) which suggested that *D. alatafructa*, and *D. insularis* should be reduced to synonymy with *D. pseudoseriata*.

The comparison between asymptomatic and symptomatic trees revealed that six of the eight species found in this study occurred on both tissues. In addition, species unique to either tissue was also detected in low numbers and was unique to a specific area and/or farm. The results suggest a higher diversity of species present on symptomatic tissues than in asymptomatic tissues, in agreement to what was found in a survey from Macadamia branches in Australia (Mohankumar et al. 2022). Laurent et al. (2020), however, used a metabarcoding strategy to study the health status of *Quercus, Vitis* and *Pinus* and found no significant difference in species diversity (Laurent et al. 2020). This study is in agreement with the results of Cruywagen et al. (2017) where there was no distinction in the species diversity between asymptomatic and symptomatic baobab trees. This is however contrary to the study by Jami et al. (2013) where they found a greater species diversity in asymptomatic branches of *Acacia karoo* in South Africa. It is clear from these studies that a number of species can be involved in disease symptoms on a particular host, but that it does not represent the full diversity of the *Botryosphaeriaceae* on the host.

All eight *Botryosphaeriaceae* species were able to cause lesions on Macadamia seedlings. Therefore, the ability of these species to cause disease was irrespective of whether the species was isolated from asymptomatic or symptomatic tissue. In addition, the results also suggests that the initiation of the pathogenic potential of the species occurred subsequent to the reduction in irrigation. The results presented here therefore aligns with the concept that *Botryosphaeriaceae* species are latent pathogens that can live in plants without causing any symptoms until some external factor, such as water stress, results in the alteration in the plantpathogen interaction and induces a switch to a pathogenic state (Slippers & Wingfield 2007).

The *Botryosphaeriaceae* species differed in aggressiveness. *Neofusicoccum luteum* was the most aggressive species in our study and was also described as one of the most aggressive species on Macadamia in Australia (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). The novel *Lasiodoplodia* sp. was the second most aggressive and should be monitored in future. *Neofusicoccum kwambonambiense* was the third most aggressive species and was also described as one of the most aggressive species on Macadamia leaves in Australia (Liddle et al. 2019). It is however also known that there is variation in species aggressiveness of the *Botryosphaeriaceae* and that isolates of the same species may differ in their aggressiveness (Moral et al. 2019). Future research should therefore include multiple isolates per species.

In this study, the aggressiveness of *Botryosphaeriaceae* species on Macadamia was not necessarily correlated with its isolation frequency. For example, *L. pseudotheobromae* was the most dominant species in both Australia and South Africa, however it was not the most aggressive species on Macadamia in both countries (Jeff-Ego and Akinsanmi 2019; Mohankumar et al. 2022). Similarly, this species was also frequently isolated in a study by Coutinho et al. (2017) from branches of hog plum, tamarind and cashew. However, its aggressiveness was lower compared to other species

when inoculated into mango fruit (Coutinho et al. 2017). To the contrary, *N. luteum* was found in low numbers but is very aggressive. The relative contribution of the different species could thus be influenced by both their frequency, as well as their aggressiveness, and both factors should be considered when considering management options.

In summary, this study expands the knowledge of the occurrence of Botryosphaeriaceae on commercially planted Macadamia trees in South Africa. The study emphasizes the extensive diversity of the Botryosphaeriaceae community that can exist on both asymptomatic and symptomatic Macadamia trees. Furthermore, all species showed a potential to cause branch dieback symptoms, with Neofusicoccum species identified as the most aggressive in this study, while Lasiodiplodia species were the most common. The rapid expansion of Macadamia plantations and the association of Botryosphaeriaceae with extreme weather conditions, raise concerns about the threat of Botryosphaeriaceae-related diseases in South Africa in the future (Desprez-Loustau et al. 2006; Slippers and Wingfield 2007). Future work should therefore focus on understanding the effect of stress on the disease expression of *Botryosphaeriaceae* species on different Macadamia cultivars. This should also include pathogenicity assays that mimic the natural penetration and infection process.

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Writing – review & editing: All authors.

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**Data availability** All sequence data generated for this study have been submitted to GenBank (accession numbers are listed in Online Resource Table 1) and any additional data are available on request.

# Declarations

**Conflict of interest** The authors have no conflict of interest to declare that are relevant to this article. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; and in the writing of the manuscript.

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