

Three *Ganoderma* species, including *Ganoderma dunense* sp. nov., associated with dying *Acacia cyclops* trees in South Africa

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ABSTRACT

Large numbers of *Acacia cyclops* trees are dying along the coastal plains of the Eastern and Western Cape Provinces of South Africa. The cause of the deaths has been attributed to a root and butt rot disease caused by the basidiomycete fungus *Pseudolagarobasidium acaciicola*. However, many signs (e.g. basidiomes) and symptoms reminiscent of *Ganoderma* root-rot disease are commonly associated with dying trees. In this study, isolates collected from basidiomes resembling species of *Ganoderma*, as well as from root and butt samples from

diseased *A. cyclops* trees were subjected to DNA sequencing and morphological studies to facilitate their identification. Multi-locus phylogenetic analyses and morphological characterisation revealed that three species of *Ganoderma* are associated with dying *A. cyclops* trees. These included *G. destructans*, a recently described species causing root-rot on trees elsewhere in South Africa. The remaining two were novel species, one of which is described here as *G. dunense*. The novel species is distinguished by its mucronate basidiome, laccate shiny pileus surface, duplex context and ovoid basidiospores. Only an immature specimen was available for the second species and a name was consequently not provided for it. Interestingly, only a single isolate representing *P. acaciicola* was recovered in this study, suggesting that further investigations are needed to ascertain the role of each of the four basidiomycetous root-rot fungi in the death of *A. cyclops* trees.

Key words: Ganodermataceae, phylogeny, *Pseudolagarobasidium*, taxonomy, rooikrans, root rot

INTRODUCTION

Acacia cyclops A. Cunn. ex G. Don (Fabaceae) is a woody shrub that occurs mainly in arid and coastal areas due to its ability to withstand severe environmental pressures such as drought, soil salinity and sand blasts (Gill 1985). In South Africa, the first trees of this species were introduced from Australia in the early 1830's to contain and stabilise the movement of sand dunes in the coastal areas of the country, including the Eastern and Western Cape Provinces (Avis 1989). Subsequently, the shrub developed rapidly to become an invasive weed and a serious environmental threat because it forms tall, thick, and almost impenetrable stands that

stifle the establishment of native plants (Henderson 1998, 2007). However, despite being detrimental to the diversity of local plants, *A. cyclops* trees also provide an important source of firewood, charcoal and construction materials (Shackleton et al. 2006).

During the course of the last four decades, *A. cyclops* trees have been dying along the coastal planes in the Eastern and Western Cape Provinces of South Africa, particularly between the towns of George and Stilbaai (Taylor 1969; Wood and Ginns 2006; Kotzé et al. 2015). The dying trees suffer from a rapidly developing root and butt-rot disease that results in die-back, wilt, and a white rot of the affected roots and root collars. The disease has been attributed to the basidiomycete root-rotting fungus *Pseudolagarobasidium acaciicola* Ginns (Wood and Ginns 2006; Kotzé et al. 2015). However, basidiomes of another fungus resembling a species of *Ganoderma* are also regularly seen attached to the bases of the dying trees, but no attempt has been made to determine the identity of this fungus (Taylor 1969; Wood and Ginns 2006).

The genus *Ganoderma* P. Karst. (Basidiomycota, Polyporales, Ganodermataceae) has a worldwide distribution and includes both saprophytic and parasitic species. They cause white rot on a wide range of host trees (Flood et al. 2000), but are also of significant medicinal and cultural importance (Bishop et al. 2015). *Ganoderma* species are known to kill a wide variety of trees, including those with high economic value such as rubber (*Hevea brasiliensis* Müll.Arg.), tea [*Camellia sinensis* (L.) Kuntze], oil palm (*Elaeis guineensis* Jacq.) and ornamental and forest trees (Ramasamy 1972; Paterson 2007; Kinge and Mih 2011; Coetzee et al. 2015). They are also the main causal agents of root and butt rot diseases of numerous *Acacia* species in tropical regions of the world (Glen et al. 2009; Coetzee et al. 2011).

Identification and circumscription of species of *Ganoderma* has mainly relied on the description of morphological characteristics of the basidiomes, resulting in considerable taxonomic confusion (Richter et al. 2015). Most recently, the Phylogenetic Species Recognition (PSR) concept has improved species delimitation in the genus (Hong and Jung 2004, Zhou et al. 2015). Analyses of DNA sequences for loci such as the internal transcribed spacer (ITS: ITS1-5.8S-ITS2), the translation elongation factor 1- α (TEF1- α) and β -tubulin, among others, have facilitated the inference of relationships between species of *Ganoderma* (Park et al. 2012; Zhou et al. 2015; Xing et al. 2016). At least 438 specific and infraspecific names are listed for *Ganoderma* in Index Fungorum (<http://www.indexfungorum.org/names/Names.asp>). It has however, been suggested that less than one third of these names are valid (Kirk et al. 2008; Richter et al. 2015).

Studies by Taylor (1969) and Wood and Ginns (2006) revealed the recurring presence of basidiomes reminiscent of *Ganoderma* attached to the bases of dying *A. cyclops* trees in the Western Cape Province of South Africa. However, having demonstrated in pathogenicity trials that *P. acaciicola* was the causal agent of the tree death, the authors did not attempt to determine the identity of the *Ganoderma* species associated with the declining trees (Wood and Ginns 2006). The report of Wood and Ginns (2006) suggests that a single *Ganoderma* species is associated with dying *A. cyclops* trees although this might not be the case. The objective of this study was to resolve the identity of the unknown *Ganoderma* species occurring on declining *A. cyclops* trees using DNA sequence comparisons for multiple gene regions as well as morphological observations.

MATERIALS AND METHODS

Fungal collection and isolation

Basidiomes and recently infected roots and basal sections were collected from dying *A. cyclops*. Collection sites included areas close to the towns of Heroldsbaai and Stillbaai in the Western Cape Province and the Nelson Mandela Bay Metropolitan area (Port Elizabeth, PE) in the Eastern Cape Province (Fig. 1). Isolations were performed on a basidiomycete selective medium composed of 2% Malt Extract Agar (MEA) [20 g/L malt extract and 15 g/L agar, Biolab, Midrand, South Africa] supplemented with benomyl, dichloran and streptomycin (BDS) as outlined in Worrall (1991). Small pieces of wood (approximately 2–3 mm³) from both infected basal sections and roots were first surface-disinfested in 7% bleach (NaClO) for ~ 90 s and rinsed twice with sterile distilled water before transferring them to the selective medium. Isolations from basidiomes were carried out by placing small sections (2–3 mm) obtained from the hymenophore onto the selective medium.

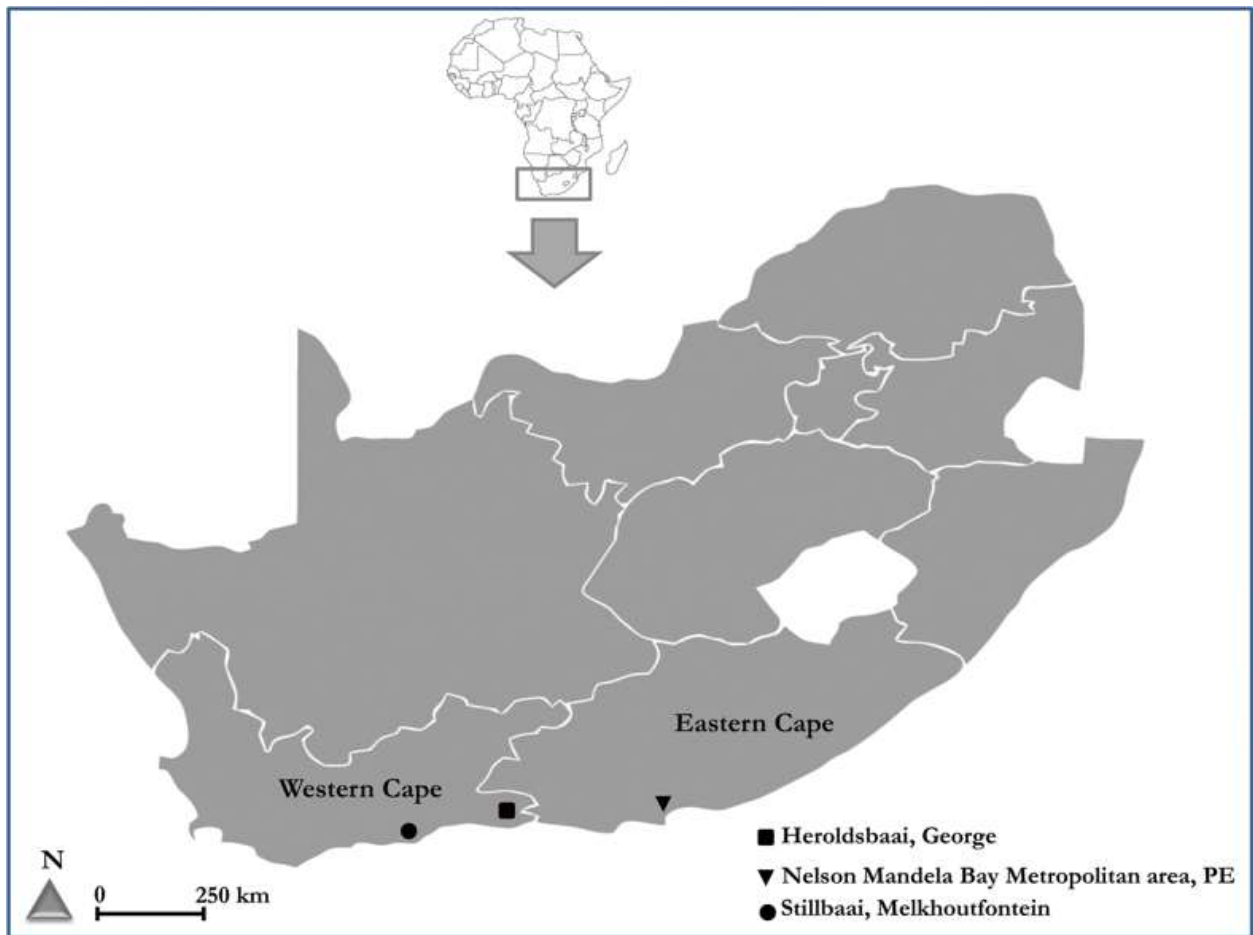


Fig. 1 South Africa map showing the sites in the Eastern and Western Cape Provinces where specimens of *Ganoderma* on *Acacia cyclops* were sampled. PE stands for Port Elizabeth

After 7-10 days of incubation at room temperature (22–24°C), pure cultures were obtained by aseptically transferring small pieces of the growing margin of the fungal isolates onto fresh 2% MEA. Two mature cultures of each isolate were preserved in the culture collection (CMW) of the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, South Africa. Representative isolates were deposited in the culture collection (CBS) of the Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands. Dried basidiomes were also deposited in the herbarium of the South African National Collection of Fungi (PREM), Roodeplaat, South Africa.

Genomic DNA extraction, amplification and sequencing

Genomic DNA was extracted from cultures (Table 1) following the Cetyltrimethylammonium Bromide (CTAB) extraction protocol described by Möller et al. (1992). DNA concentrations were determined with a NanoDrop® ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). Working DNA concentrations for polymerase chain reactions (PCRs) were obtained by adjusting the initial concentrations of the extracted genomic DNA to 100 ng/ μ L. The complete internal transcribed spacer (ITS) region, including ITS-1, ITS-2 and the 5.8S small subunit gene was amplified for all isolates using primers ITS1 and ITS4 (White et al. 1990). Amplifications of the partial translation elongation factor 1- α (TEF1- α) and partial β -tubulin gene regions were obtained using primer pairs EF595F/EF1160R (Kausrud and Schumacher 2001) and β -tubulin_F/ β -tubulin_R (Park et al. 2012), respectively.

All PCR reactions were carried out in 25 μ L total mixtures consisting of 17.5 μ L sterile SABAX water (Adcock Ingram Ltd, Bryanston, RSA), 1 μ L genomic DNA (100 ng/ μ L), 0.5 μ L (2.5 units) of MyTaq™ DNA polymerase (Bioline), 5 μ L 5 x MyTaq™ Reaction Buffer supplied with the enzyme and 0.5 μ L (10 mM) of each primer. The PCR conditions used with primers ITS1/ITS4 and EF595F/EF1160R were as follows: an initial denaturation step at 95°C for 4 min, followed by 35 cycles of denaturation at 95°C for 30 s, 30 s of annealing at 55°C and 60 s of extension at 72°C. The reactions were completed with a final elongation step at 72°C for 7 min. Amplifications with the β -tubulin_F/ β -tubulin_R primers were carried out with the same parameters as those used by Park et al. (2012). PCR products were electrophoresed on 2% agarose gels after staining with GelRed™ nucleic acid dye (Biotium Incorporation, USA) and visualized under UV illumination. Sephadex G-50 columns (Sigma, Steinheim, Germany) were then used to purify the PCR products following the recommendations of the manufacturer.

Table 1. Isolates used in the phylogenetic analyses

Species	Voucher no.	Geographical origin	GenBank accession numbers		
			ITS	β -tubulin	TEF1- α
<i>Ganoderma adspersum</i>	Yao34456	United Kingdom	AJ006685	-	-
<i>G. applanatum</i>	Dai 12483	China	KF494999	-	KF494977
<i>G. applanatum</i>	ATCC 44053	Japan	JQ520161	JQ675614	-
<i>G. aridicola</i>	Dai12588 (holotype)	South Africa	KU572491	-	KU572502
<i>G. austroafricanum</i>	CMW 41454	South Africa	KM507324	-	-
<i>G. boninense</i>	WD 2028	Japan	KJ143905	-	KJ143924
<i>G. boninense</i>	WD 2085	Japan	KJ143906	-	KJ143925
<i>G. carnosum</i>	MJ 21/08	Czech Republic	KU572492	-	-
<i>G. carnosum</i>	JV 8709/8	Czech Republic	KU572493	-	-
<i>G. carnosum</i>	CBS 516.96	Netherlands	-	JQ675616	-
<i>G. cupreum</i>	GanoTK4	Cameroon	JN105701	-	-
<i>G. cupreum</i>	GanoTK7	Cameroon	JN105702	-	-
<i>G. curtisii</i>	CBS 100131	United States of America (USA)	JQ781848	-	KJ143926
<i>G. curtisii</i>	CBS 100132	USA	JQ781849	-	KJ143927
<i>G. curtisii</i>	CBS 100132	Netherlands	-	JQ675617	-
<i>G. destructans</i>	CBS 139793 (type)	South Africa	NR132919	MG020151*	MG020213*
<i>G. destructans</i>	CMW 43671	South Africa	KR183857	MG020156*	MG020220*
<i>G. destructans</i>	CMW42129	South Africa	MG020232	MG020158	MG020191
<i>G. destructans</i>	CMW42130	South Africa	MG020233	MG020159	MG020192
<i>G. destructans</i>	CMW42131	South Africa	MG020234	MG020160	MG020193
<i>G. destructans</i>	CMW42134	South Africa	MG020235	MG020161	MG020216
<i>G. destructans</i>	CMW42135	South Africa	MG020236	MG020162	MG020214
<i>G. destructans</i>	CMW42136	South Africa	MG020237	MG020163	MG020194
<i>G. destructans</i>	CMW42137	South Africa	MG020238	MG020164	MG020195
<i>G. destructans</i>	CMW42138	South Africa	MG020239	MG020165	MG020196
<i>G. destructans</i>	CMW42139	South Africa	MG020240	MG020166	MG020197
<i>G. destructans</i>	CMW42140	South Africa	MG020241	MG020167	MG020215
<i>G. destructans</i>	CMW42141	South Africa	MG020242	MG020168	MG020198
<i>G. destructans</i>	CMW42142	South Africa	MG020243	MG020169	MG020199

Table 1. (continued)

Species	Voucher no.	Geographical origin	GenBank accession numbers		
			ITS	β -tubulin	TEF1- α
<i>G. destructans</i>	CMW42143	South Africa	MG020244	MG020170	MG020221
<i>G. destructans</i>	CMW42146	South Africa	MG020245	MG020171	MG020200
<i>G. destructans</i>	CMW42147	South Africa	MG020246	MG020172	MG020201
<i>G. destructans</i>	CMW42148	South Africa	MG020247	MG020173	MG020202
<i>G. destructans</i>	CMW42151	South Africa	MG020250	MG020174	MG020203
<i>G. destructans</i>	CMW42152	South Africa	MG020251	MG020175	MG020204
<i>G. destructans</i>	CMW42153	South Africa	MG020252	MG020176	MG020205
<i>G. destructans</i>	CMW42154	South Africa	MG020253	MG020177	MG020206
<i>G. destructans</i>	CMW42155	South Africa	MG020254	MG020178	MG020217
<i>G. destructans</i>	CMW42158	South Africa	MG020256	MG020179	MG020224
<i>G. destructans</i>	CMW42159	South Africa	MG020257	MG020180	MG020223
<i>G. destructans</i>	CMW42160	South Africa	MG020258	MG020181	MG020207
<i>G. destructans</i>	CMW42161	South Africa	MG020259	MG020182	MG020208
<i>G. destructans</i>	CMW42162	South Africa	MG020260	MG020183	MG020209
<i>G. destructans</i>	CMW42163	South Africa	MG020261	MG020184	MG020218
<i>G. destructans</i>	CMW42164	South Africa	MG020262	MG020185	MG020210
<i>G. destructans</i>	CMW42165	South Africa	MG020263	MG020186	MG020211
<i>G. destructans</i>	CMW45109	South Africa	MG020266	MG020187	MG020225
<i>G. destructans</i>	CMW45110	South Africa	MG020267	MG020188	MG020222
<i>G. destructans</i>	CMW45113	South Africa	MG020268	MG020189	MG020212
<i>G. destructans</i>	CMW45114	South Africa	MG020269	MG020190	MG020219
<i>G. dunense</i> sp. nov	CMW42149	South Africa	MG020248	MG020153	MG020226
<i>G. dunense</i> sp. nov	CMW42150	South Africa	MG020249	MG020154	MG020228
<i>G. dunense</i> sp. nov	CMW42157 (Type)	South Africa	MG020255	MG020150	MG020227
<i>G. gibbosum</i>	XSD-35	Unknown	EU273514	-	-
<i>G. gibbosum</i>	AS5.624 type 3	China	AY593856	-	-
<i>G. enigmaticum</i>	Dai 15970	Africa	KU572486	-	KU572496
<i>G. enigmaticum</i>	Dai 15971	Africa	KU572487	-	KU572497
<i>G. enigmaticum</i>	CBS 139792 (type)	South Africa	NR132918	MG020157*	MG020231*

Table 1. (continued)

Species	Voucher no.	Geographical origin	GenBank accession numbers		
			ITS	β -tubulin	TEF1- α
<i>G. leucocontextum</i>	Dai 15601	China	KU572485	-	KU572495
<i>G. leucocontextum</i>	GDGM 44489	China	KM396271	-	-
<i>G. lingzhi</i>	Wu 1006-38 (holotype)	China	JQ781858	-	JX029976
<i>G. lingzhi</i>	Dai 12574	China	KJ143908	-	JX029977
<i>G. lingzhi</i>	Dai 12479	China	JQ781864	-	JX029975
<i>G. lobatum</i>	JV 0409/13J	USA	KF605675	-	-
<i>G. lobatum</i>	JV 1212/10J	USA	KF605676	-	KU572501
<i>G. lobatum</i>	ATCC 42985	Canada	-	JQ675618	-
<i>G. lobatum</i>	ASI 7061	USA	-	JQ675619	-
<i>G. lucidum</i>	Cui 9207	China	KJ143910	-	KJ143928
<i>G. lucidum</i>	K 175217	UK	KJ143911	-	KJ143929
<i>G. lucidum</i>	ASI 7117	Korea	-	JQ675633	-
<i>G. lucidum</i>	IUM 4303	Bangladesh	-	JQ675635	-
<i>G. lucidum</i>	IUM 0047	Korea	-	JQ675627	-
<i>G. meredithae</i>	ATCC 64492	USA	-	JQ675643	-
<i>G. meredithae</i>	ASI 7140	Unknown	-	JQ675644	-
<i>G. mirabile</i>	CBS 218.36	Philippines	-	JQ675645	-
<i>G. multipileum</i>	CWN 04670	China	KJ143913	-	KJ143931
<i>G. multipileum</i>	Dai 9447	China	KJ143914	-	KJ143932
<i>G. multiplicatum</i>	Dai 12320	China	-	-	KU572500
<i>G. multiplicatum</i>	Dai 13710	China	-	-	KU572499
<i>G. mutabile</i>	Yuan 2289 (type)	China	JN383977	-	-
<i>G. neojaponicum</i>	ASI 7032	Unknown	-	JQ675646	-
<i>G. oerstedii</i>	GO138	Unknown	-	DQ288098	-
<i>G. oregonense</i>	CBS 265.88	USA	JQ781875	NS	KJ143933
<i>G. oregonense</i>	ASI 7049	USA	-	JQ675647	-
<i>G. oregonense</i>	ASI 7067	USA	-	JQ675650	-
<i>G. oregonense</i>	CBS 266.88	USA	JQ781876	-	-
<i>G. pfeifferi</i>	CBS 747.84	Netherlands	-	JQ675651	-

Table 1. (continued)

Species	Voucher no.	Geographical origin	GenBank accession numbers		
			ITS	β -tubulin	TEF1- α
<i>G. philippii</i>	E7098	Indonesia	AJ536662	-	-
<i>G. philippii</i>	E7092	Indonesia	AJ608710	-	-
<i>G. resinaceum</i>	BR 4150	France	KJ143915	-	-
<i>G. resinaceum</i>	CBS 194.76	Netherlands	KJ143916	-	KJ143934
<i>G. resinaceum</i>	ATCC 52416	Argentina	-	JQ675652	-
<i>G. resinaceum</i>	IUM 3651	Czech Republic	-	JQ675657	-
<i>G. ryvardenii</i>	HKAS 58055	Cameroon	HM138670	-	-
<i>G. ryvardenii</i>	HKAS 58053 (type)	Cameroon	HM138671	-	-
<i>G. sessile</i>	JV 1209/9	USA	KF605629	-	KJ143936
<i>G. sessile</i>	LDW 20121017	USA	KJ143917	-	KJ143935
<i>G. sinense</i>	Wei 5327	China	KF494998	-	KF494976
<i>Ganoderma</i> sp. A	CMW45100	South Africa	MG020264	MG020152	MG020229
<i>Ganoderma</i> sp. A	CMW45101	South Africa	MG020265	MG020155	MG020230
<i>Ganoderma</i> sp.	ASI 7150	Unknown	-	JQ675665	-
<i>Ganoderma</i> sp.	ASI 7151	Unknown	-	JQ675666	-
<i>G. subamboinens</i>	GSUB136	Unknown	-	DQ288096	-
<i>G. subamboinens</i>	GSUB137	Unknown	-	DQ288097	-
<i>G. tenue</i>	GTEN24	Unknown	-	DQ288074	-
<i>G. tornatum</i>	CBS 109679	Netherlands	-	JQ675670	-
<i>G. tornatum</i>	BAFC1172	Argentina	AH008096	-	-
<i>G. tornatum</i>	BAFC1139	Argentina	AH008098	-	-
<i>G. tropicum</i>	He 1232	China	KF495000	-	KF494975
<i>G. tropicum</i>	Yuan 3490	China	JQ781880	-	KJ143938
<i>G. tsugae</i>	Dai 12751b	USA	KJ143919	-	KJ143939
<i>G. tsugae</i>	Dai 12760	USA	KJ143920	-	KJ143940
<i>G. tsugae</i>	ATCC 64795	Canada	-	JQ675668	-
<i>G. tsugae</i>	ASI 7064	USA	-	JQ675669	-
<i>G. valesiacum</i>	CBS 428.84	USA	-	JQ675671	-
<i>G. webeianum</i>	CBS 219.36	Philippines	-	JQ675672	-

Table 1. (continued)

Species	Voucher no.	Geographical origin	GenBank accession numbers		
			ITS	β -tubulin	TEF1- α
<i>G. zonatum</i>	FL-02	USA	KJ143921	-	KJ143941
<i>G. zonatum</i>	FL-03	USA	KJ143922	-	KJ143942
<i>Tomophagus colossus</i>	TC-02	Vietnam	KJ143923	-	KJ143943
<i>Trametes suaveolens</i>	-	Unknown	-	FJ410378	-

Reference sequences with star (*) are those generated in this study. In bold are isolates of the newly proposed species.

Purified PCR products were sequenced in 12 μ L reaction volumes using the same primer pairs as in the amplification reactions. DNA sequencing reactions were done using Big Dye (Perkin-Emmer, Warrington, UK), following the protocol outlined by the manufacturer. Purification of the sequencing PCR products followed the same approach as for the PCR products. DNA sequencing was carried out on a DNA Analyzer ABI PRISM™ 3100 (Applied BioSystems, Foster City, CA, USA) at the sequencing facility of the University of Pretoria. CLC Main Workbench v7.6.1 was used to assess the quality of the electropherograms and to construct consensus sequences. DNA sequences of the β -tubulin and TEF1- α gene regions of *G. destructans* and *G. enigmaticum*, previously described from South Africa (Coetzee et al. 2015), were also generated for comparative purposes. These sequences as well as those of the novel species were deposited in GenBank.

DNA sequence datasets and multi-gene phylogenetic analysis

Preliminary identification based on BLASTn searches of ITS sequences against those of reference sequences in GenBank was made to confirm that the sequences of the fungal isolates from *A. cyclops* were species of *Ganoderma*. Phylogenetic analyses were subsequently performed to determine the phylogenetic placement of the isolates from *A. cyclops*. For this purpose, four sequence datasets were generated. Of these, three represented the ITS, β -tubulin and TEF1- α gene regions alone, and one consisted of the combined sequences for all three loci. In addition to sequences generated in this study, the datasets also included reference sequences used in previous studies, including those of Glen et al. (2009), Douanla-Meli et al. (2009), Kinge et al. (2012), Park et al. (2012), Zhou et al. (2015), Coetzee et al. (2015), Xing et al. (2016) (Table 1). *Tomophagus colossus* (Fr.) C.F. Baker and *Trametes suaveolens* (L.) Fr. were selected as outgroup taxa. Sequence alignments were performed using the online version of MAFFT

(<http://mafft.cbrc.jp/alignment/server/index.html>) v7 (Katoh and Standley 2013), and the phylogenetic relationships between taxa were evaluated using Maximum Likelihood (ML), Maximum Parsimony (MP) and Bayesian Inference (BI) analyses. The GTR+G+I substitution model determined by JModeltest v2.1.6 (Darriba et al. 2012) on CIPRES (Miller et al. 2012) under the Akaike Information Criterion (AIC) was selected as the best-fit nucleotide substitution model and incorporated in the phylogenetic analyses.

Maximum likelihood analyses were performed in RaxML (Stamatakis 2006) using raxmlGUI 1.3 (Silvestro and Michalak 2012) with 10 parallel runs. Maximum likelihood bootstrap (MLB) analysis was done using 1000 replications. For BI analyses, four parallel runs, each with four Monte Carlo Markov Chains (MCMC) were performed using MrBayes v3.2 (Ronquist et al. 2012). Sampling of trees was carried out for four million generations with trees sampled at every 100th generation. The first 25% of the sampled trees were discarded as burn-in and the Bayesian posterior probabilities (BPP) were calculated for the remaining trees. Maximum Parsimony analyses were performed in PAUP* version 4.0b10 (Swofford 2002). A heuristic search option with a tree bisection-reconnection (TBR) branch swapping algorithm, involving 100 random stepwise additions of sequences, was used to generate the most parsimonious trees. All characters were of equal weight, gaps were treated as missing data and tree branches of zero length were collapsed. The robustness of clades at the branch nodes was estimated using bootstrap (MPB) with 1000 replicates and using the same settings to obtain the fundamental tree but with the addition of sequences set to closest. Other estimated parameters included tree length (TL), consistency index (CI), retention index (RI), rescaled consistency index (RC), and homoplasy index (HI). The generated phylogenetic trees were visualized using MEGA 5.05 (Tamura et al. 2011).

Morphology

Microscopic observations were made from fine sections of basidiomes mounted on microscope slides in 5% KOH (potassium hydroxide) with and without 1% phloxine and Melzer's reagent (IKI). The observations were made using a Nikon Eclipse Ni compound microscope (Nikon, Japan) at a magnification of up to 100x under an oil-immersion objective, and on a Zeiss Stemi SV6 stereo microscope. Images of the microscopic structures were captured with a Nikon DS-Ri2 camera fitted on the microscope and the measurements were made using the program NIS-Elements BR (Nikon Instruments Software-Elements Basic research).

RESULTS

Signs and Symptoms

Typical signs and symptoms of Ganoderma root rot disease (Glen et al. 2009; Coetzee et al. 2011) were observed on the declining *A. cyclops* trees. These included crown wilt and die-back, fresh basidiomes of *Ganoderma* attached to the bases of the affected trees (Fig. 2a), basal cankers and butt rot (Fig. 2b), and roots with red rhizomorphic encrusting sheets covering a whitish mycelial mat (Fig. 2c).



Fig. 2 Signs and symptoms of *Ganoderma* species infecting *Acacia cyclops* in South Africa. (a) Basidiome of *G. destructans* with reddish brown (laccate shiny) pilear surface attached to the base of wilting tree, (b) Collar rot, (c) Reddish brown to dark brown rhizomorphic sheet of a *Ganoderma* sp. covering root of dying tree

Fungal Isolates

In total, 38 isolates resembling species of *Ganoderma* were obtained from a total of 55 diseased and dying trees. Of these, three were isolated from roots covered with red-coloured rhizomorphic sheets; three from roots without rhizomorphic sheets; five from tree bases and the remaining 27 from fresh basidiomes (Table 1). The isolates in culture were either white to yellowish or completely yellow and ranged from having flat to fluffy mycelial growth.

DNA sequence comparisons

The ITS, β -tubulin and TEF1- α gene regions of the 38 isolates were successfully amplified using PCR, and DNA sequences were obtained for all the amplicons. Results of BLASTn searches based on ITS revealed that 36 isolates had a high level of DNA sequence similarity with *Ganoderma destructans*. Two others (CMW45100, CMW45101) had a high level of sequence similarity with *G. applanatum* (Pers.) Pat. and *G. gibbosum* (Blume & T. Nees) Pat. Besides the 38 *Ganoderma* isolates, a single isolate that was obtained from roots without rhizomorphic sheets was identified as *P. acaciicola*, and it was excluded from the rest of the study.

Sequence data and multi-gene phylogenetic analyses

The ITS dataset comprised of 89 ingroup taxa and one outgroup taxon. The dataset resulted in an alignment length of 651 characters, of which 357 were constant, 61 parsimony-uninformative and 233 parsimony-informative. The heuristic search resulted in eight parsimonious trees with TL = 540, CI = 0.622, RI = 0.899 and RC = 0.559. Since the tree topologies resulting from the BI, ML and MP analyses were nearly identical, only the ML tree is presented, along with the statistical values (BPP and Bootstrap values) of the two other analyses (Fig. 3). In the phylogenetic tree of the ITS sequence data, the isolates generated from *A. cyclops* formed two groups, clustering at distant positions. The first group, composed of 36 isolates, formed a monophyletic clade with isolates representing *G. destructans* and *G. multipileum*. However, this clade had statistical support only in the ML analysis (75% bootstrap support). The second group, comprised of isolates CMW45100 and CMW45101, nested in a strongly supported clade (MLB = 93%, MPB = 100% and BPP = 1) with isolates representing *G. applanatum*, *G. gibbosum* and

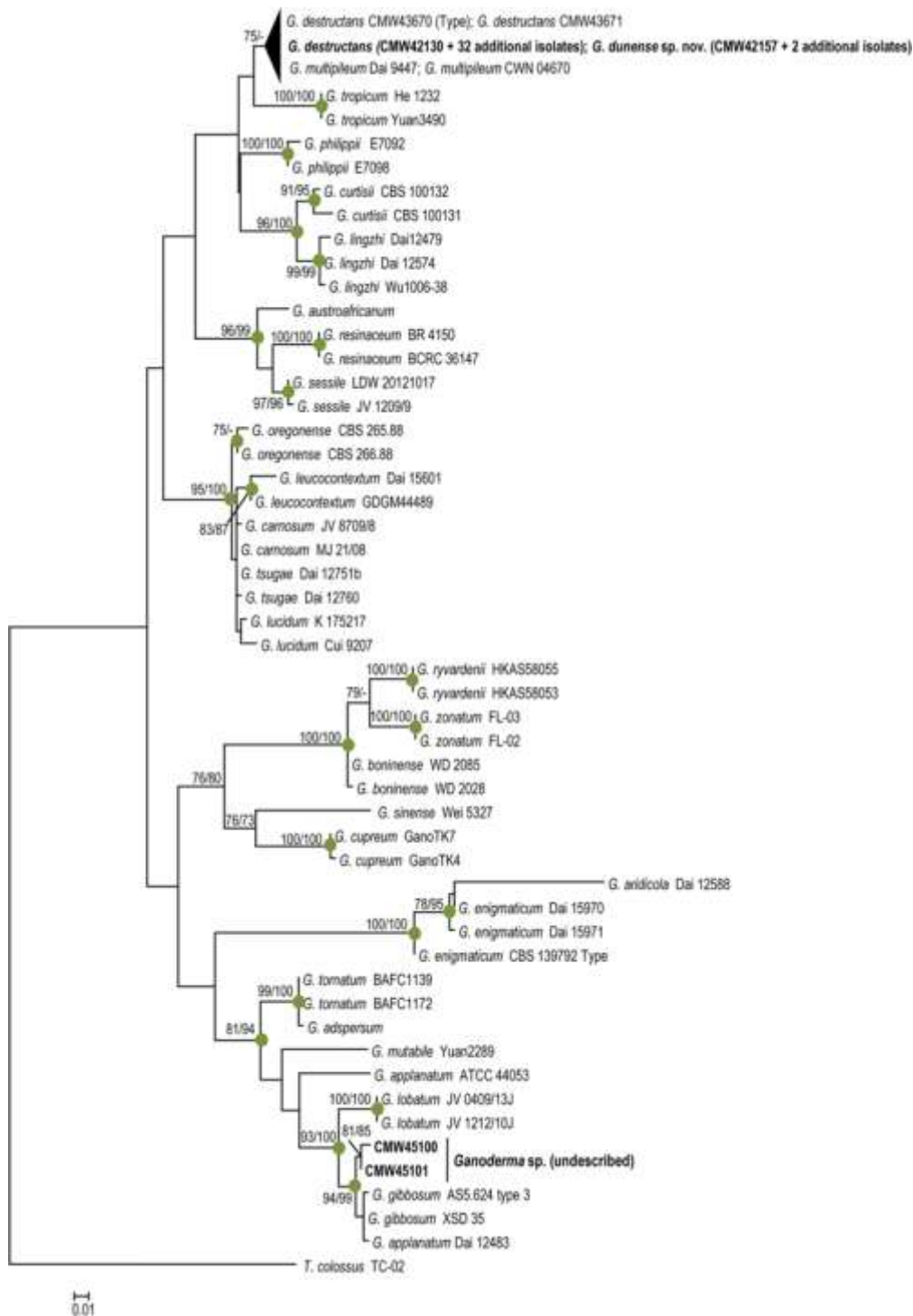
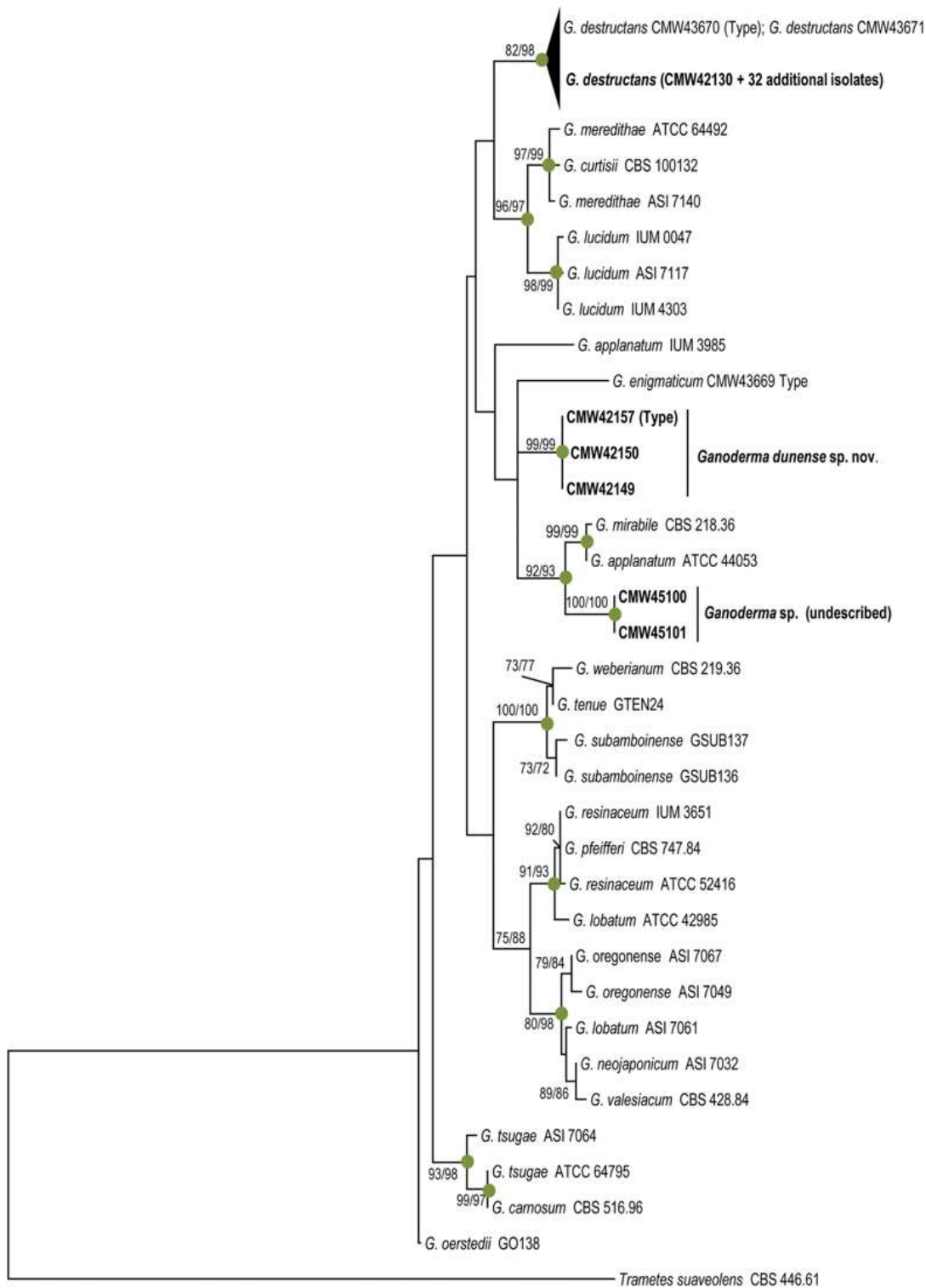


Fig. 3 Maximum likelihood tree based on ITS sequences, depicting the phylogenetic position of isolates obtained from *Acacia cyclops* (in bold) and related species within *Ganoderma*. Bootstrap values $\geq 70\%$ from 1000 replicates of ML and MP analyses are displayed at the nodes. Posterior probability ≥ 0.95 from Bayesian Inference analysis are indicated at nodes with green circles

G. lobatum (Schwein.) G.F. Atk. However, the two *A. cyclops* isolates formed a well-resolved sub-clade, although not sufficiently supported by BI analysis (BPP \leq 0.95).

The β -tubulin sequence dataset consisted of 68 taxa, which resulted in an alignment length of 401 total characters. Of the total number of characters, 277 were constant, 44 were parsimony-uninformative and 80 parsimony-informative. The heuristic search yielded 100 parsimonious trees with TL = 206, CI = 0.544, RI = 0.887 and RC = 0.482. Only the ML consensus tree incorporating the statistical values of BI and MP is presented (Fig. 4) since the three analyses yielded almost congruent topologies. In the β -tubulin phylogeny, unlike that for the ITS dataset, the isolates obtained from *A. cyclops* clustered at three different positions (Fig. 4). Thirty-three isolates (generated in this study) formed a monophyletic clade having strong statistical support with sequences representing *G. destructans* (MLB = 82%, MPB = 98%, BPP = 0.96). Three other isolates (CMW42149, CMW42150 and CMW42157) grouped in a well-resolved and strongly supported monophyletic cluster (MLB and MPB = 99%, BPP = 1), while the third group comprising isolates CMW45100 and CMW45101 formed another strongly supported clade (MLB and MPB = 100%, BPP = 1) that was closely related to the lineage including *G. mirabile* and *G. applanatum*.

The dataset from TEF1- α contained 71 ingroup taxa and one outgroup taxon. The total number of characters after alignment of the dataset was 579, of which 375 were constant, 35 were parsimony-uninformative and 169 parsimony-informative. The heuristic search resulted in 100 parsimonious trees with TL = 435, CI = 0.584, RI = 0.885 and RC = 0.517. The three analyses resulted in almost identical tree topologies. Hence, only the ML tree is shown alongside with BI and MP statistical values. As with β -tubulin, the TEF1- α phylogeny placed the isolates from *A. cyclops* at three different positions (Fig. 5). The first group, consisting of the same 33



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0.01

Fig. 4 Maximum likelihood tree based on β -Tubulin sequences, depicting the phylogenetic position of isolates obtained from *Acacia cyclops* (in bold) and related species within *Ganoderma*. Bootstrap values $\geq 70\%$ from 1000 replicates of ML and MP analyses are displayed at the nodes. Posterior probability ≥ 0.95 from Bayesian Inference analysis are indicated at nodes with green circles

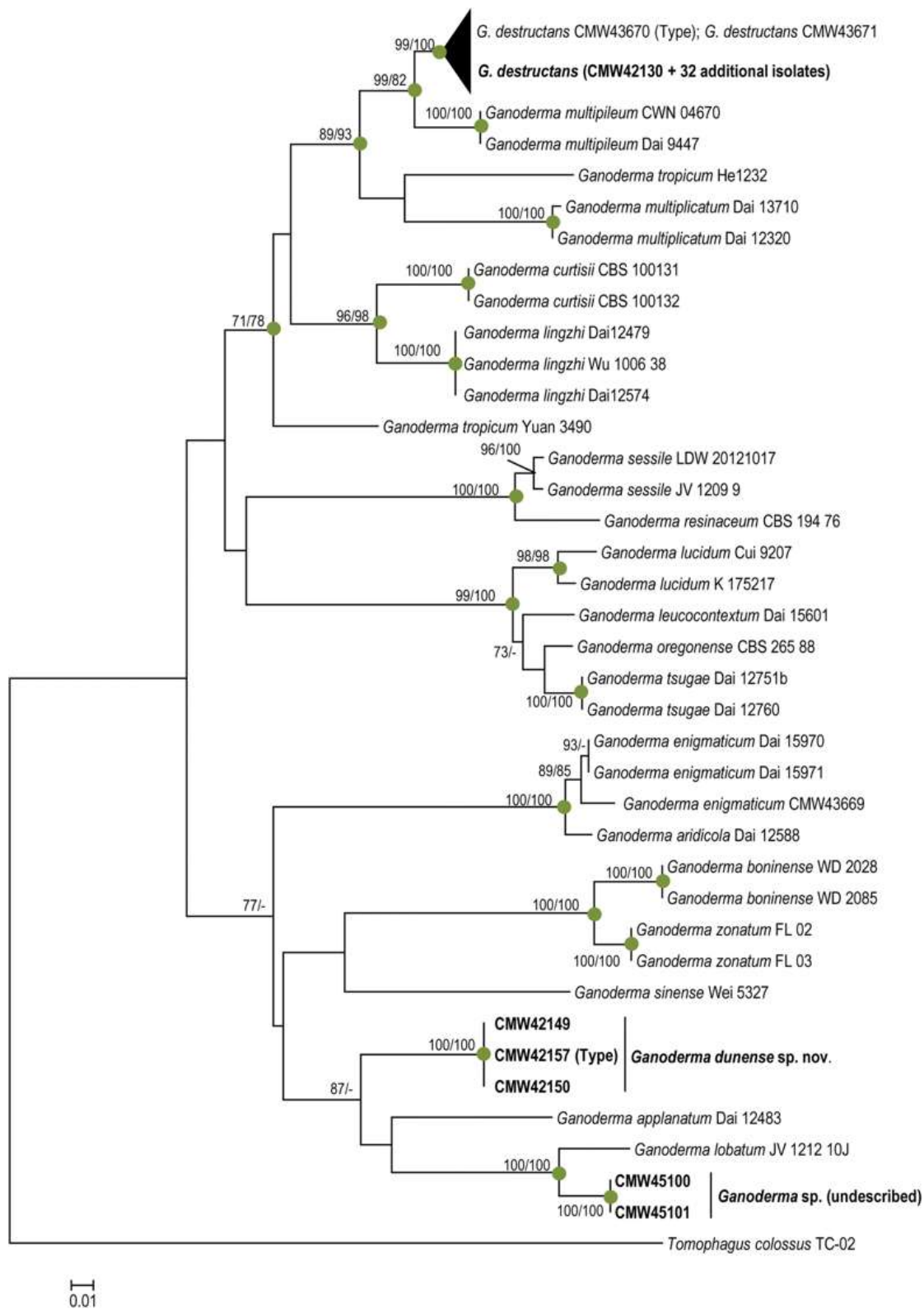


Fig. 5 Maximum likelihood tree based on TEF1- α gene sequences, depicting the phylogenetic position of isolates obtained from *Acacia cyclops* (in bold) and related species within *Ganoderma*. Bootstrap values $\geq 70\%$ from 1000 replicates of ML and MP analyses are displayed at the nodes. Posterior probability ≥ 0.95 from Bayesian Inference analysis are indicated at nodes with green circles

isolates as in the β -tubulin gene tree, formed a strongly supported monophyletic clade (MLB = 99%, MPB = 100%, BPP = 0.98) with sequences representing *G. destructans* (generated in this study). This group was closely related to the lineage representing *G. multipileum*. The second group, including isolates CMW42149, CMW42150 and CMW42157, formed a well-supported monophyletic clade (MLB and MPB = 100%, BPP = 1). The third group of isolates, including CMW45100 and CMW45101, formed a cluster with *G. lobatum*, but represented a distinct lineage with high bootstrap values (MLB and MPB = 100%, BPP=1).

The combined dataset of the ITS, β -tubulin and TEF1- α sequences comprised 115 taxa and 1631 total characters. In this dataset the ITS, β -tubulin and TEF1- α respectively contributed 651, 401 and 579 characters. Of the total number of characters, 1152 were constant, 94 parsimony-uninformative and 385 were parsimony-informative. The heuristic search generated 100 parsimonious trees with TL = 1005, CI = 0.541, RI = 0.878, and RC = 0.475. Since the three analyses resulted in similar topologies, only the topology of the ML, incorporating BPP and MP bootstrap values, is presented (Fig. 6). The phylogeny for the combined dataset supported that of β -tubulin and TEF1- α , placing the isolates obtained from *A. cyclops* at three different positions. A first group, consisting of the same 33 isolates as in the β -tubulin and TEF1- α phylogenies, formed a well-resolved monophyletic clade with strong statistical support (MLB = 88% and MPB = 98%) in ML and MP analyses with isolates representing *G. destructans*. This clade was closely related to *G. multipileum*, but both formed distinct lineages. The second group, comprised of isolates CMW42149, CMW42150 and CMW42157, clustered in a clearly resolved monophyletic clade with strong statistical support (MLB and MPB =100%, BPP = 1) in all three analyses. Finally, the third group, composed of isolates CMW45100 and CMW45101, formed a

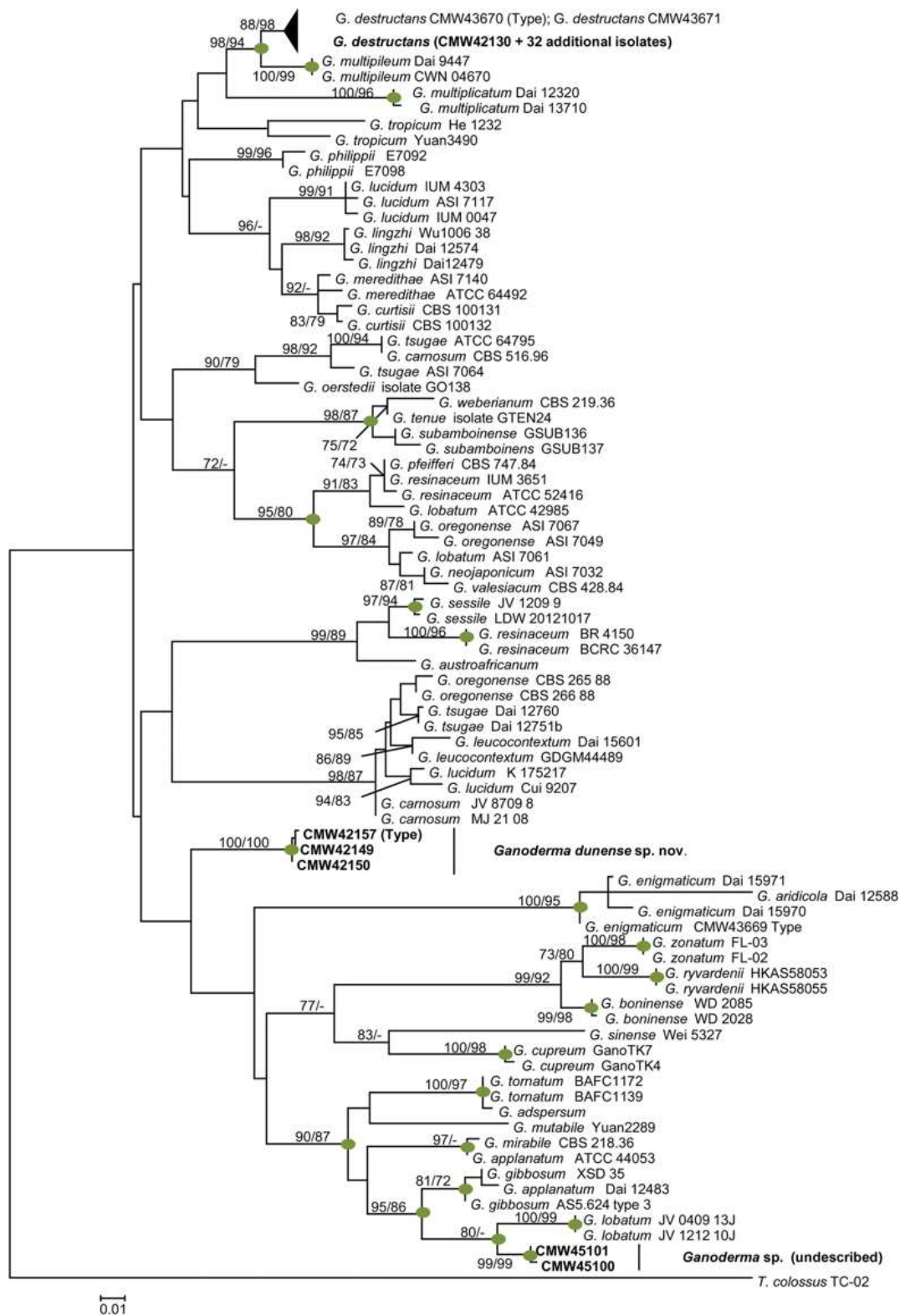


Fig. 6 Maximum likelihood tree based on combined dataset of ITS, β -Tubulin and TEF1- α gene sequences, depicting the phylogenetic position of the isolates obtained from *Acacia cyclops* (in bold) and related species within *Ganoderma*. Bootstrap values $\geq 70\%$ from 1000 replicates of ML and MP analyses are displayed at the nodes. Posterior probability ≥ 0.95 from Bayesian Inference analysis are indicated at nodes with green circles

highly supported clade (MLB and MPB = 99%, BPP = 1), which was closely related to *G. lobatum* but with marginal support.

Taxonomy

Based on a phylogenetic species recognition concept, the 38 isolates of *Ganoderma* from *A. cyclops* in South Africa, represent three distinct taxa. One of these was *G. destructans* M.P.A. Coetzee, Marinc. & M.J. Wingf. (Coetzee et al. 2015) and two others represented novel species. One of these novel taxa (CMW45100 and CMW45101) is not described because only a rudimentary and immature basidiome (Fig. 7) could be found for it. The other new species, represented by isolates CMW42149, CMW42150 and CMW42157 is described as follows:

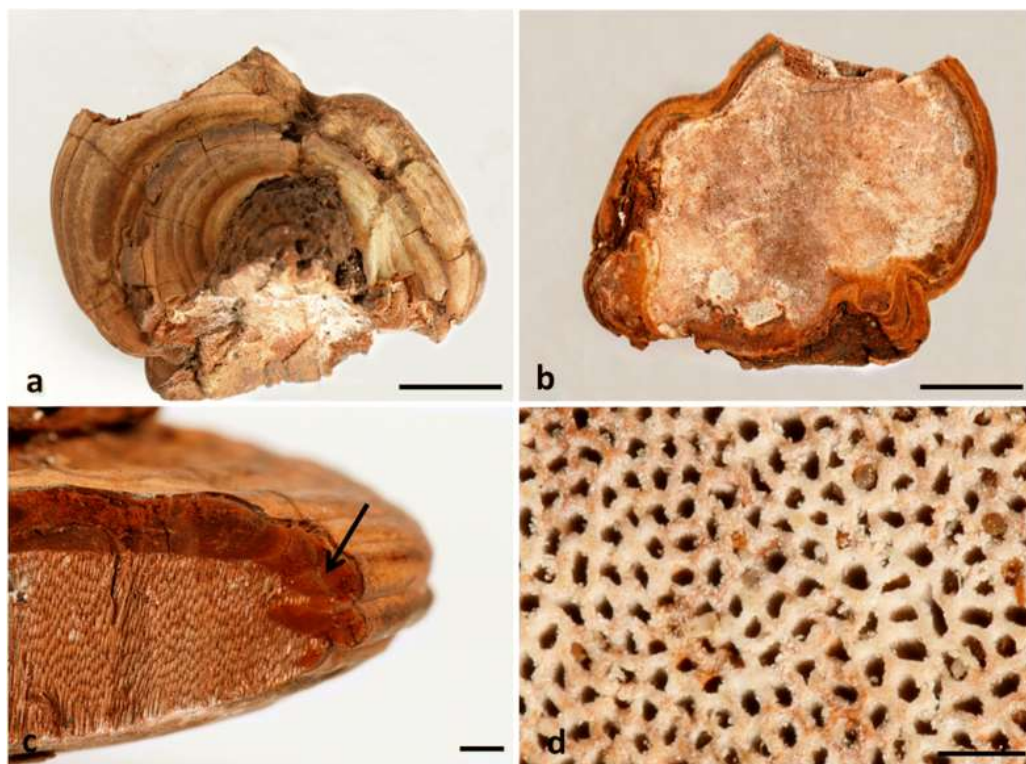


Fig. 7 *Ganoderma* sp. (a-b) pilear and hymenial surfaces of an immature basidiome; (c) homogenous chocolate brown context with discrete resin-like incrustations/deposits (arrowed); (d) pore surface. Bars: a-b = 2 cm; c = 2 mm; d = 1 mm



Fig. 8 *Ganoderma dunense* sp. nov. (Type, PREM 61936). (a-b) mature reniform-like basidiome showing a pileus surface partially covered with a dull brown deposit of basidiospores and a poroid dark brown hymenial surface; (c) pores; (d) duplex context with a light brown upper layer and a chocolate brown lower layer towards the tubes; (e) basidiospores; (f) palisade of clavi-form cutis cells; (g) skeletal hyphae; (h) skeletal hypha showing a distinct lumen (arrowed). Bars: a-b = 5 cm; c = 1 mm; d = 1 cm; e-h = 10 μ m

Ganoderma dunense Tchetet, Rajchenb., & Jol. Roux sp. nov.

MB823686

Fig. 8

Etymology: Name refers to the coastal sand dunes where this species was collected on dying *A. cyclops* trees.

Diagnosis: *Ganoderma dunense* is morphologically similar to species in the *Ganoderma lucidum* complex. It is characterised by a light and spongy, perennial, applanate, dimidiate or reniform, mucronate basidiome, a laccate shiny pilear surface and a thick, soft-spongy duplex context. *Hymenial surface* poroid, white when fresh. *Hyphal system* trimitic and *basidiospores* ovoid, $10.9\text{--}12.3 \times 7.3\text{--}8.5 \mu\text{m}$, double walled with hyaline exosporium and yellowish brown coarse echinulae endosporium, IKI-.

Type: **South Africa**, Western Cape Province, George, Heroldsbaai (S34° 03.242' E22° 22.711'), at the base of dying *Acacia cyclops* A. Cunn. ex G. Don (Fabaceae), 13 April 2014, J.M. Tchetet Tchoumi and J. Roux, JMT137 (holotype-PREM61936, culture ex-type CMW42157 = CBS142831). GenBank accession number: ITS = MG020255, β -tubulin = MG020150, EF1- α = MG020227.

Description: **Basidiomes** light, with spongy consistency, perennial, applanate, dimidiate or reniform, up to 14–20 cm wide, 11–15 cm radius, and 2–2.5 cm thick at base, mucronate; mucro lateral, more or less ellipsoid, 2.3–5 cm long, 1.8–4 cm wide. Margins more or less circular, slightly lobate and round. **Pileus surface** laccate shiny, dark brown to dark brown chestnut, mostly covered by a dull brown deposit of basidiospores, glabrous, sulcate with wide furrows

that become narrow towards the margins, and with irregular protuberances and a thin crust. **Hymenial surface** poroid, white when fresh turning brown to dark brown upon bruising; pores round to somewhat angular and elongated, 3–4 per mm; dissepiments thin. **Context** soft-spongy, 4–8 mm thick, duplex, with a light brown upper layer and a chocolate brown lower layer towards the tubes, absence of black resin-like deposits. **Tubes** light greyish brown, not stratified, up to 1.5 cm long. **Hyphal system** trimitic; generative hyphae not easily found, colourless, thin-walled, bearing clamp septa; skeletal hyphae poorly branched, brown to brown chestnut, thick-walled, 3.1–8 μm thick, lumen 1.4–2.2 μm wide; skeleto-binding hyphae present, branched at the apex with long flagelliform branches, basal stem unbranched. **Cutis** a hymeniodermis composed of a palisade of vertical and closely packed clavate cells; cells yellowish brown, thick-walled, some inflated at the apex and with rare growths of protuberances on the lateral parts, amyloid, 22.5–34.4 \times 5.4–10.6 μm . **Cystidia** and **basidia** not seen. **Basidiospores** ovoid with indistinct truncate apex, 10.9–12.3 \times 7.3–8.5 μm (11.6 \pm 0.7 \times 7.9 \pm 0.6 μm), IKI-, double-walled, exosporium hyaline with inter-wall pillars, endosporium yellowish brown with coarse echinulae.

Additional specimen: South Africa, Western Cape Province, George, Heroldsbaai (S34° 03.249' E22° 22.720'), at the base of dying *A. cyclops*. 13 April 2014, J.M. Tchetet Tchoumi and J. Roux, JMT124 (paratype-PREM 61937, culture ex-type CMW42149 = CBS142945). GenBank accession number: ITS = MG020248, β -tubulin = MG020153, EF1- α = MG020226.

Remarks: *Ganoderma dunense* and *G. destructans* cannot be distinguished from each other based on ITS phylogeny (Fig. 3). They segregate only in phylogenies for the β -tubulin and TEF1- α gene regions as well as in the combined genes (Figs 4, 5, 6). This phylogenetic divergence based on these three analyses is also reflected in some morphological characters. Morphologically, *G. destructans* differs from *G. dunense* in having a stipitate basidiome, a laccate shiny reddish

brown pilear surface with well-developed protuberances, a not completely homogenous context, slightly smaller pores (3–5 per mm) and larger basidiospores ($11\text{--}14 \times 7\text{--}9 \mu\text{m}$, Coetzee et al. 2015; Table 2).

Ganoderma dunense is macroscopically similar to *G. enigmaticum* M.P.A. Coetzee, Marinc. & M.J. Wingf., *G. aridicola* J.H. Xing & B.K. Cui and *G. austroafricanum* M.P.A. Coetzee, M.J. Wingf., Marinc. & Blanchette, with which it is genetically related (Fig. 6). They share the same origin (South Africa), a white poroid hymenophore surface when fresh, irregular pores and absence of black melanoid bands in the context (Crous et al. 2014; Coetzee et al. 2015; Xing et al. 2016; Table 2). However, *G. enigmaticum* and *G. aridicola* differ from *G. dunense* in their ecology. *Ganoderma enigmaticum* occurred as a parasite at the base of the trunk of *Ceratonia siliqua* L. in Pretoria, Gauteng Province (Coetzee et al. 2015), while *G. aridicola* was recovered on charred wood of *Ficus* in Durban, KwaZulu-Natal Province (Xing et al. 2016). In addition, *G. enigmaticum* differs morphologically from *G. dunense* by having a stipitate basidiome, homogenous context and narrower ellipsoid basidiospores ($8\text{--}11 \times 3.5\text{--}6 \mu\text{m}$, Coetzee et al. 2015; Table 2). Similarly, *G. aridicola* differs from *G. dunense* by having a sessile basidiome, a fuscous homogenous context, distinctly stratified tube layers, smaller pores (6–8 per mm) and broadly ellipsoid basidiospores ($9.7\text{--}11.2 \times 7\text{--}7.8 \mu\text{m}$, Xing et al. 2016; Table 2). *G. austroafricanum* has an annual sessile basidiome, a dimitic hyphal system and smaller ($8\text{--}11 \times 5.5\text{--}7 \mu\text{m}$) subglobose basidiospores (Crous et al. 2014; Table 2).

When compared morphologically with other laccate *Ganoderma* species from South Africa and tropical Africa based on published descriptoins (Steyaert 1961, 1962, 1967, 1972, 1980; Table 2), *G. dunense* shares few characteristics with most of these species. Exceptions are *G. namutambalaense* Steyaert (1962), collected in Uganda and *G. megalosporum* Steyaert (1962)

Table 2. Morphological comparison of *G. dunense* sp. nov. with other laccate *Ganoderma* taxa found in South Africa and tropical Africa

Taxon	Shape	Pores/mm	Spores (μm)	Cutis	Other	Reference
<i>Ganoderma africanum</i>	NI	NI	NI	NI	type not found	Moncalvo and Ryvarden 1997
<i>G. aridicola</i>	sessile dimidiate, pilear surface fuscous black when fresh, reddish brown to black upon drying	6–8	9.7–11.2 \times 7–7.8, broadly ellipsoid	amyloid elements 30–55 \times 5–8 μm	skeletal hyphae up to 2.5–5 μm wide	Xing et al. 2016
<i>G. austroafricanum</i>	sessile, dimidiate	3–4	8–11 \times 5.5–7 broadly ellipsoid	NI	NI	Crous et al. 2014
<i>G. capensis</i>	sessile	5–6	9.75–10.5 \times 5.75–6.2	elements 50 \times 5–10 μm up to 9–12 μm when capitate	dimitic hyphal system	Reid 1975
	-	6	10–12 \times 6–7 ovoid to ellipsoid	amyloid elements 35–45(–55) \times 7–9 μm	trimitic with skeleto-binding hyphae, skeletal hyphae 5.5–10 μm diam.	This publication
<i>G. destructans</i>	stipitate globular pileus	3–5	11–14 \times 7–9	amyloid elements 13–35 \times 4.5–7.5	homogeneous context	Coetzee et al. 2015
	laccate shiny reddish brown, stipe lateral to eccentric and circular to ellipsoid, pileus dimidiate to circular, with small to wide protuberances	-	-	-	not completely homogeneous context	This publication

Table 2. (continued)

Taxon	Shape	Pores/mm	Spores (μm)	Cutis	Other	Reference
<i>G. dunense</i> sp. nov.	sessile but with a lateral mucro, pileus dimidiate or reniform	3–4	10.9-12.3 \times 7.3-8.5	amyloid elements clavate, 22.5–34.4 \times 5.4–10.6 μm	duplex context present	This publication
<i>G. enigmaticum</i>	stipitate globular pileus	3–5	8–11 \times 3.5–6	amyloid elements 20–46 \times 5.5–9 μm	-	Coetzee et al. 2015
<i>G. hildebrandii</i>	small basidiome centrally stipitate	7–8	7–8(8.5) \times 4.5–5.5(–6)	IKI–elements similar but wider up to 15 μm	dextrinoid skeletal hyphae	Moncalvo and Ryvarden 1995, 1997
<i>G. reticulatosporum</i>	centrally stipitate pileus, umbrella shaped	3–4	25–26 \times 15.5–16	NI	perhaps <i>Humphreya</i> sp.	Reid 1973 Moncalvo and Ryvarden 1997
<i>G. namutambalaense</i>	substipitate flabellate	5–6	11.5–14 \times 8–9.5	elements sphaeropedunculate, 50 \times 4–7(–10) μm , sustaining hyphae 2-3 μm	from Uganda	Steyaert 1962
<i>G. megalosporum</i>	sessile dimidiate	3–4	11.5–13 \times 8–9	elements subcylindric 30 \times 4-6 μm	from Kenya	Steyaert 1962

*NI refers to no information in the literature

Table 3. Morphological comparison of *G. dunense* with its close relatives in the global multi-locus phylogeny (Fig. 6)

Taxon	Shape	Pores/mm	Spores (μm)	Cutis	Other	Reference
<i>G. carnosum</i> .	laterally to rarely eccentrically stipitate, dimidiate, orbicular to reniform	3–4	10–13 \times 7–8.5	amyloid elements up to 75 μm from clamp to the apex	crust becoming duller and darker with age, presence of protuberances	Patouillard 1889; Ryvardeen and Melo 2014
<i>G. dunense</i> sp. nov.	sessile but with a lateral mucro, pileus dimidiate or reniform	3–4	10.9–12.3 \times 7.3–8.5	amyloid elements clavate, 22.5–34.4 \times 5.4–10.6 μm	duplex context present	This publication
<i>G. leucocontextum</i>	stipitate, reniform, dimidiate to flabelliform	4–6	9.5–12.5 \times 7–9	amyloid elements 30–60 \times 8–10 μm	thinner context of white to slightly cream colour	Li et al. 2015
<i>G. lucidum</i>	fan-shaped to dimidiate; centrally to laterally stipitate, applanate	4–5	7–11 \times 6–8	amyloid hyphal end cells up to 7–11 μm diam	orange red to bay upper surface; deeper colour context	Ryvardeen and Gilbertson 1993; Ryvardeen and Melo 2014
<i>G. oregonense</i>	sessile to substipitate basidiome	2–3	13–17 \times 8–10	-	grow on conifers	Gilbertson and Ryvardeen, 1986
<i>G. resinaceum</i>	stipitate, dimidiate to broadly attached	3–4	9–11.5 \times 4.5–7	amyloid elements up to 50 μm from clamp to the apex	excretes a resinous layer on the pileus edge	Ryvardeen 2000; Ryvardeen and Melo 2014
<i>G. sessile</i>	dimidiate, sessile pileus sometimes imbricate, conchate to flabelliform	4–5	12–16 \times 6–8	cylindric, smooth elements, 60–75 \times 7–10 μm	context thin, soft corky or woody, radially fibrous, concentrically zonate, ochraceous	Murrill 1902; Gottlieb and Wright 1999
<i>G. tsugae</i>	fan-shaped to dimidiate; centrally to laterally stipitate	4–6	13–15 \times 7.5–8.5	-	whitish and soft-corky context; grow on conifers	Murrill 1902; Gilbertson and Ryvardeen 1986

collected in Kenya, which share minor similarities with *G. dunense*. They can, however, easily be distinguished from *G. dunense* in that *G. namutambalaense* has a substipitate basidiome, smaller pores and larger basidiospores, while *G. megalosporum* has larger basidiospores and much smaller sub-cylindric cuticular cells (Table 2).

Other *Ganoderma* spp. phylogenetically closely related to *G. dunense* (Fig. 6) are compared in Table 3. They differ from *G. dunense* either in their ecology, features of the pilear surface, pore size, basidiospore shape and size, and/or their distribution.

DISCUSSION

Three distinct *Ganoderma* species were found associated with dying *Acacia cyclops* trees in the Eastern and Western Cape Provinces of South Africa. Two of these species represent novel taxa but a name was provided only for one due to absence of mature basidiomes needed to describe the second species. The description of the new species brings to 11 the *Ganoderma* spp. recorded from South Africa (Van del Bijl 1921; Reid 1973; 1974; 1975; Moncalvo and Ryvarden 1997; Crous et al. 2014; Coetzee et al. 2015 and Xing et al. 2016). Of these, only five have been described based on DNA sequence data.

The majority of *Ganoderma* isolates obtained in this study (33 = ~ 87%) represented *G. destructans*. This fungus was recently described as the causal agent of a serious root disease of non-native Jacaranda (*Jacaranda mimosifolia* D. Don) trees in the city of Pretoria (Tshwane Metropolitan area, Gauteng Province) in South Africa (Coetzee et al. 2015). The common occurrence of *G. destructans* on *A. cyclops* in the Western Cape, together with its occurrence on *J. mimosifolia* more than 1000 km further north in the country suggests that it does not only have

a wide host range, but also wide geographical distribution. Based on the assertion of Coetzee et al. (2015) that this fungus might be a native species, *A. cyclops* thus represents the second non-native host that it has successfully colonized. Its wide geographic distribution also supports the contention that it is a native South African fungus.

The two novel species of *Ganoderma* obtained from *A. cyclops* in this study were represented by only five isolates. Two were of the *Ganoderma* that was not describe and three were of the newly described *G. dunense*. Based on the appearance of the pilear surface, *G. dunense* could be linked to the *G. lucidum* complex, while the undescribed *Ganoderma* sp. was related to species in the *G. applanatum* complex (Imazeki 1952; Richter et al. 2015).

As has been found in many other studies (i.e. Gottlieb et al. 2000 and Zhou et al. 2015) sequence data for the ITS gene regions were not sufficient to delineate between *G. dunense* and *G. destructans*. Based on the ITS phylogeny, *G. dunense* would be treated as the same as *G. destructans*. But phylogenies inferred from the β -tubulin and TEF1- α data sets, as well as that of the combined genes, showed that this taxon represented a novel species. Morphological characterization also revealed very distinct macro- and microscopic characters that could further distinguish between these two taxa.

In the global multi-locus phylogeny, *G. dunense* formed a well-resolved monophyletic clade. However, it shared a genetic affinity with *G. enigmaticum*, *G. aridicola*, *G. carnosum*, *G. lucidum* (s. lat.), *G. leucocontextum*, *G. tsugae*, *G. oregonense*, *G. austroafricanum*, *G. resinaceum* and *G. sessile*. These species all reside in the *G. lucidum* complex subgenus *Ganoderma* (Li et al. 2015; Zhou et al. 2015; Richter et al. 2015). The phylogenetic relatedness

of *G. dunense* to these species confirms its initial assignment to this complex based on morphology.

The extent of infection by *Ganoderma* species in this study raises an intriguing question about the role of *P. acaciicola* in the death of *A. cyclops* and whether it is the only causal agent of the deaths of these trees as reported by Wood and Ginns (2006). This question is worth noting because most of the isolates recovered from recently infected trees in the current study were of *G. dunense* and only a single isolate was identified as *P. acaciicola*. Moreover, although *Ganoderma* was included in the pathogenicity trials conducted by Wood and Ginns (2006), results of the present study show that at least three species of *Ganoderma* are associated with the dying *A. cyclops* in these areas. It is not known which of the three *Ganoderma* species was used in the aforementioned trials. Additionally, most of the sampled trees exhibited symptoms typical of *Ganoderma* root rot similar to those observed in previous studies (Glen et al. 2009; Coetzee et al. 2011). It is, therefore, clear that further investigations should be undertaken to determine the role of *P. acaciicola* and *Ganoderma* species in the death of *A. cyclops* trees in the coastal areas of South Africa.

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