RESEARCH ARTICLE



Two novel species of *Calonectria* isolated from soil in a natural forest in China

QianLi Liu¹, ShuaiFei Chen¹

I China Eucalypt Research Center (CERC), Chinese Academy of Forestry (CAF), ZhanJiang 524022, GuangDong Province, China

Corresponding author: ShuaiFei Chen (shuaifei.chen@gmail.com)

Academic editor: G.Mugambi | Received 24 June 2017 | Accepted 3 August 2017 | Published 22 August 2017

Citation: Liu QL, Chen SF (2017) Two novel species of *Calonectria* isolated from soil in a natural forest in China. MycoKeys 26: 25–60. https://doi.org/10.3897/mycokeys.26.14688

Abstract

Species of *Calonectria* include important pathogens of numerous agronomic and forestry crops worldwide, and they are commonly distributed in soils of tropical and subtropical regions of the world. Previous research results indicated that species diversity of *Calonectria* in China is relatively high. Most *Calonectria* spp. reported and described from China were obtained from diseased *Eucalyptus* tissues or soils in *Eucalyptus* plantations established in tropical and subtropical areas in southern China. Recently, a number of *Calonectria* isolates were isolated from soils in a natural forest in the temperate region of central China. These isolates were identified by DNA sequence comparisons for the translation elongation factor 1-alpha (*tef1*), histone H3 (*his3*), calmodulin (*cmdA*) and β -tubulin (*tub2*) gene regions, combined with morphological characteristics. Two novel species of *Calonectria* were identified and described, and are named here as *Calonectria lichi* and *Ca. montana*, which reside in the Prolate Group and Sphaero-Naviculate Group, respectively. This study revealed that more species of *Calonectria* may occur in natural forests in central China than previously suspected.

Key words

Cylindrocladium, pathogen, phylogeny, taxonomy

Introduction

Calonectria species include many notorious plant pathogens and are widely distributed in tropical and subtropical areas of the world (Crous 2002, Lombard et al. 2010d, Aiello et al. 2013, Vitale et al. 2013, Alfenas et al. 2015). These species can cause serious plant

epidemics on a wide range of plant hosts (Peerally 1991, Schoch et al. 2001, Crous 2002), and result in considerable economic losses to agriculture and forestry. Example include shoot blight on *Pinus* spp. in South African nurseries (Crous et al. 1991), root rot on Myrtus communis in Tunisia (Lombard et al. 2011), and leaf blight on Buxus sempervirens in Iran (Mirabolfathy et al. 2013). In addition, members of the genus Calonectria are responsible for red crown rot of *Glycine max* (soybean) in Japan (Yamamoto et al. 2017), fruit rot of Nephelium lappaceum (rambutan) in Puerto Rico (Serrato-Diaz et al. 2013) and root rot of Arbutus unedo (strawberry) in Italy (Vitale et al. 2009). As an important fast-growing tree species, Eucalyptus plays a significant role in the global pulpwood supply. Previous research showed that Calonectria leaf blight (CLB), associated with several species of *Calonectria*, is considered to be one of the most prominent Eucalyptus leaf diseases that has occurred in numerous countries such as Brazil (Alfenas et al. 2015, Lombard et al. 2016), China (Zhou et al. 2008, Chen et al. 2011), Colombia (Rodas et al. 2005), India (Sharma et al. 1984) and Vietnam (Old et al. 1999). Other fungal diseases of Eucalyptus spp. caused by Calonectria species include dampingoff, shoot blight, and root rot, which have been observed in Brazil (Ferreira 1989) and South Africa (Crous et al. 1991), and these diseases have received considerable attention.

Calonectria spp. are soil-borne fungi, they can form microsclerotia in soil and infected plant roots, stem and leaves as primary inoculum. After diseased tissues decompose or the plants are harvested, microsclerotia are released into the soil, which allows them to survive for extended periods even up to 15 years or more (Sobers and Littrell 1974, Crous 2002). Species of *Calonectria* are also rapidly dispersed via aerial dissemination and water movement, which leads to the transmission of *Calonectria* disease (Vitale et al. 2013). Based on previous studies, at least 145 *Calonectria* species have been identified using molecular data and have been described worldwide (Crous 2002, Crous et al. 2004, 2006, 2012, 2013, 2015, Lombard et al. 2010a, b, c, 2011, 2015, 2016, Chen et al. 2011, Xu et al. 2012, Alfenas et al. 2013a, b, 2015, Gehesquière et al. 2015). Sixty species were isolated from soil samples collected in subtropical or tropical regions (Crous 2002, Crous et al. 2004, Lombard et al. 2010a, b, c, 2015, 2016, Chen et al. 2011, Xu et al. 2012, Alfenas et al. 2015).

In China, *Calonectria* has a relatively high species diversity, and to date, 28 *Calonectria* species have been identified and described. Based on previous studies, *Calonectria* species have been reported in nine provinces and one Special Administrative Region (SAR), which with the exception of LiaoNing and ShanDong Provinces belong to temperate regions (Luan et al. 2006, Li et al. 2010). Most *Calonectria* have been isolated from agronomic crops or forestry plantations in subtropical and tropical regions, including FuJian, GuangDong, GuangXi, GuiZhou, HaiNan, JiangXi and YunNan Provinces, as well as Hong Kong SAR (Crous et al. 2004, Lombard et al. 2010a, 2015, Chen et al. 2011, Gai et al. 2012, Xu et al. 2012, Pei et al. 2015).

China has large areas of plantation and natural forests. To date 27 *Calonectria* species have been isolated from *Eucalyptus* tissues with CLB/leaf rot symptoms or from soils originating from *Eucalyptus* plantations in tropical or subtropical areas in Fu-Jian, GuangDong, GuangXi and HaiNan Provinces (Crous et al. 2004, Lombard et

al. 2010a, 2015, Chen et al. 2011). However, little information is known about the species diversity of *Calonectria* in natural forests. In this study, a number of soil samples were collected from a natural forest in the temperate region of central China, and baited with alfalfa seeds for *Calonectria*. The aim of the current study was to identify these isolates using a combination of phylogenetic analyses and morphological characteristics and to gain a preliminary understanding of the species diversity of *Calonectria* in natural forests in China.

Materials and methods

Fungal isolates

In April 2016, 17 soil samples were collected from a natural forestry area in central China. The collected soils were baited with surface-disinfested (30 s in 75% ethanol and washed several times with sterile water) *Medicago sativa* (alfalfa) seeds using the method described by Crous (2002). After one week, sporulating conidiophores were produced on infected alfalfa tissue. Using a dissection microscope AxioCam Stemi 2000C (Carl Zeiss, Germany), conidial masses were selected and scattered onto 2 % malt extract agar (MEA) (20 g malt extract powder and 20 g agar powder per liter of water: malt extract powder was obtained from Beijing Shuangxuan microbial culture medium products factory, Beijing, China; the agar powder was obtained from Beijing Solarbio Science & Technology Co., Ltd., Beijing, China) using sterile needles. After incubation at 25 °C for one day, germinated spores were individually transferred onto fresh MEA under the dissection microscope and were incubated at 25 °C for one week.

Single conidial cultures were deposited in the Culture Collection of the China Eucalypt Research Centre (CERC), Chinese Academy of Forestry (CAF), ZhanJiang, GuangDong Province, China. Representative isolates were stored in the China General Microbiological Culture Collection Center (CGMCC), Beijing, China. The specimens (pure fungal cultures) were deposited in the Collection of Central South Forestry Fungi of China (CSFF), GuangDong Province, China.

DNA extraction, PCR and sequence reactions

Single conidial cultures grew on MEA for one week at 25 °C, after which actively growing mycelium was scraped using a sterilized scalpel and transferred into 2 mL Eppendorf tubes. Total genomic DNA was extracted following the protocols "Extraction method 5: grinding and CTAB" described by Van Burik et al. (1998). The extracted DNA was dissolved in 30 μ L TE buffer (1 M Tris-HCl and 0.5 M EDTA, pH 8.0), and a Nano-Drop 2000 spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) was used to quantify the concentration.

Based on previous research (Lombard et al. 2010d, Alfenas et al. 2015), partial gene regions including translation elongation factor 1-alpha (*tef1*), histone H3 (*his3*), calmodulin (*cmdA*) and β -tubulin (*tub2*), were used as successful DNA barcodes at species, being able to clearly distinguish between intra- and inter-specific divergence. The primer pairs EF1-728F/EF2, CYLH3F/CYLH3R, CAL-228F/CAL-2Rd and T1/CYLTUB1R were used to amplify the fragments of the respective *tef1*, *his3*, *cmdA* and *tub2* genes (Lombard et al. 2010d).

The PCR reaction mixture used to amplify the different loci consisted of TopTaqTM Master Mix 12.5 μ L (Qiagen Inc., Hilden, Germany), forward primer 1 μ L, 10 μ M (Invitrogen, Shanghai, China), reverse primer 1 μ L, 10 μ M (Invitrogen, Shanghai, China), and RNase-Free H₂O 8.5 μ L (Qiagen Inc., Hilden, Germany), and 2 μ L (100 ng/ μ L) of the DNA samples was added as the template to each PCR reaction. The amplifications were performed in 25 μ L reaction volumes on an MJ Mini Cycler (BIO-RAD, Hercules, CA, USA) under the conditions described by Groenewald et al. (2013). The amplification products were separated by 1.5% agarose gel electrophoresis and visualized with SYBR Safe DNA gel stain (Thermo Fisher Scientific Inc., USA).

Amplified fragments were sequenced in both directions using the same primer pairs used for amplification by the Beijing Genomics Institute, Guangzhou, China. Sequences were edited using MEGA v. 6.0.5 software (Tamura et al. 2013). All sequences of the isolates obtained in this study were submitted to GenBank (http://www.ncbi. nlm.nih.gov) (Table 1).

Phylogenetic analyses

The sequences generated from this study were added to other sequences of closely related *Calonectria* species downloaded from GenBank for phylogenetic analyses. All sequences used in this study were aligned using the online MAFFT v. 7 (http://mafft. cbrc.jp/alignment/server) with the alignment strategy FFT-NS-i (Slow; interactive refinement method). The aligned sequences were manually edited using MEGA v. 6.0.5 and were deposited in TreeBASE (http://treebase.org).

Phylogenetic analyses were conducted on individual *tef1*, *his3*, *cmdA* and *tub2* sequence datasets and on the combined datasets for the four gene regions, depending on the sequence availability. Two methods, maximum parsimony (MP) and maximum likelihood (ML) were used for phylogenetic analyses.

MP analyses were performed using PAUP v. 4.0 b10 (Swofford 2003), gaps were treated as a fifth character, and characters were unordered and of equal weight with 1000 random addition replicates. A partition homogeneity test (PHT) was conducted to determine whether data for the four genes could be combined. The most parsimonious trees were acquired using the heuristic search option with stepwise addition, tree bisection, and reconstruction branch swapping. MAXTREES was set to 5,000, and zero-length branches were collapsed. A bootstrap analysis (50% majority rule, 1,000 replicates) was carried out to determine statistical support for internal nodes in trees.

Cnariae	Icolota No †#	Cubetworta	Somuling cita	Collector		GenBank acc	cession No. ^{5,1}		Dafamura
opene	TOTALC INO.	Outparte	oampung are		tef1	his3	cmdA	tub2	
	CBS 114813	Pinus radiata	New Zealand	H. Pearson	GQ267292	DQ190693	GQ267360	DQ190591	Gadgil and Dick 2004
Calonectria acteola	CBS 114812	P. radiata	New Zealand	H. Pearson	GQ267291	DQ190692	GQ267359	DQ190590	Gadgil and Dick 2004
Ca. aconidialis	CBS 136086	Soil in Eucalyptus plantation	HaiNan, China	X. Mou & S.F. Chen	KJ462785	KJ463133	KJ463017	N/A ⁵	Lombard et al. 2015
Ca. arbusta	CBS 136079	Soil in Eucalyptus plantation	GuangXi, China	X. Zhou & G. Zhao	KJ462787	KJ463135	KJ463018	KJ462904	Lombard et al. 2015
	CBS 114073	Leaf litter	Thailand	N.L. Hywel-Jones	AY725705	AY725658	AY725741	AY725616	Crous et al. 2004
Ca. astatica	CBS 112711	Leaf litter	Thailand	N.L. Hywel-Jones	AY725702	AY725655	AY725738	AY725613	Crous et al. 2004
Ca. australiensis	CBS 112954	Ficus pleurocarpa	Australia	C. Pearce & B. Paulu	GQ267293	DQ190699	GQ267363	DQ190596	Crous et al. 2006
Ca. brassicicola	CBS 112841	Brassica sp.	Indonesia	M.J. Wingfield	KX784689	N/A	KX784561	KX784619	Lombard et al. 2016
Ca. canadiana	CBS 110817	Picea sp.	Canada	S. Greifenhagen	GQ267297	AF348228	AY725743	AF348212	Lombard et al. 2010b
	CBS 114827	Soil	Hong Kong	E.C.Y. Liew	AY725710	AY725661	AY725747	AY725619	Lombard et al. 2010b
Ca. comensis	CBS 112744	Soil	Hong Kong	E.C.Y. Liew	AY725709	AY725660	AY725746	AY725618	Lombard et al. 2010b
:	CBS 293.79	Camellia sinensis	Indonesia	N/A	GQ267301	DQ190639	GQ267373	DQ190564	Lombard et al. 2010b
Ca. comounti	CBS 114704	Arachis pintoi	Australia	D. Hutton	GQ267300	DQ190638	GQ267372	DQ190563	Lombard et al. 2010b
	CBS 112220	Eucalyptus grandis	Colombia	M.J. Wingfield	AY725711	AY725662	AY725748	GQ267207	Lombard et al. 2010b
Ca. colomotensis	CBS 112221	E. grandis	Colombia	M.J. Wingfield	AY725712	AY725663	AY725749	AY725620	Lombard et al. 2010b
	CBS 127198	E. grandis	FuJian, China	M.J. Wingfield	HQ285822	HQ285808	MF527084	HQ285794	Chen et al. 2011; This study
Ca. croustana	CBS 127199	E. grandis	FuJian, China	M.J. Wingfield	HQ285823	HQ285809	MF527085	HQ285795	Chen et al. 2011; This study
Ca. curvispora	CBS 116159	Soil	Madagascar	P.W. Crous	GQ267302	AY725664	GQ267374	AF333394	Lombard et al. 2010b
	CBS 125275	E. grandis	Sumatra Utara	M.J. Wingfield	GQ267338	GQ267267	GQ267430	GQ267218	Lombard et al. 2010b
Ca. eucatypti	CBS 125276	E. grandis	Sumatra Utara	M.J. Wingfield	GQ267339	GQ267268	GQ267431	GQ267219	Lombard et al. 2010b
	CBS 136247	Soil in Eucalyptus plantation	Guangxi, China	X. Zhou & G. Zhao	KJ462798	KJ463146	KJ463029	KJ462914	Lombard et al. 2015
Ca. expansa	CBS 136078	Soil in <i>Eucalyptus</i> plantation	Guangdong, China	X. Zhou & G. Zhao	KJ462797	KJ463145	KJ463028	KJ462913	Lombard et al. 2015
	CBS 127201	E. grandis	FuJian, China	M.J. Wingfield	HQ285820	HQ285806	MF527089	HQ285792	Chen et al. 2011; This study
Ca. Jujuanensis	CBS 127200	E. grandis	FuJian, China	M.J. Wingfield	HQ285819	HQ285805	MF527088	HQ285791	Chen et al. 2011; This study

Table 1. The species of Calonectria used in this study.

Charies	Isolate No †#	Currente	Samulina cite	Collector		GenBank acc	cession No. ^{5/1}		Reference
emide	1201410 1 10.	Outonary	oampinig are		tefl	his3	cmdA	tub2	
	CBS 136092	Soil in Eucalyptus plantation	Guangxi, China	X. Mou & R. Chang	KJ462803	KJ463151	KJ463034	KJ462919	Lombard et al. 2015
La. guangxiensis	CBS 136094	Soil in Eucalyptus plantation	Guangxi, China	X. Mou & R. Chang	KJ462804	N/A	KJ463035	KJ462920	Lombard et al. 2015
Ca. hainanensis	CBS 136248	Soil in Eucalyptus plantation	Hainan, China	X. Mou & S.F. Chen	KJ462805	KJ463152	KJ463036	N/A	Lombard et al. 2015
	CBS 114828	Soil	Hong Kong	E.C.Y. Liew	AY725717	AY725667	AY725755	AY725622	Lombard et al. 2010b
Ca. hongkongensis	CBS 114711	Soil	Hong Kong	M.J. Wingfield	AY725716	AY725666	AY725754	AY725621	Lombard et al. 2010b
Ca. ilicicola	CBS 190.50	Solanum tuberosum	Indonesia	K.B. Boedijn & J. Reitsma	AY725726	AY725676	AY725764	AY725631	Lombard et al. 2010b
	CBS 112215	A. hypogaea	U.S.A.	Beute	AY725726	AY725684	AY725765	AY725639	Crous et al. 2004
	CBS 112823	Syzygium aromaticum	Indonesia	M.J. Wingfield	AY725718	AY725668	AY725756	AY725623	Lombard et al. 2010b
Ca. maonestae	CBS 112840	S. aromaticum	Indonesia	M.J. Wingfield	AY725720	AY725670	AY725758	AY725625	Lombard et al. 2010b
C. indonesiana	CBS 112936	Soil	Indonesia	M.J. Wingfield	KX784701	N/A	KX784573	KX784631	Lombard et al. 2016
	CBS 144.36	N/A	N/A	N/A	GQ267332	GQ267262	GQ267453	GQ267239	Lombard et al. 2010b
Ca. maustata	CBS 114684	Rhododendron sp.	U.S.A.	N.E. El-Gholl	GQ267333	DQ190653	GQ267454	AF232862	Lombard et al. 2010b
	CBS 170.77	Idesia polycarpa	New Zealand	N/A	GQ267308	GQ267249	GQ267380	GQ267209	Lombard et al. 2010b
Ca. Kyotensis	CBS 413.67	Paphiopedilum callosum	Celle, Germany	W. Gerlach	GQ267307	GQ267248	GQ267379	GQ267208	Lombard et al. 2010b
Ca. lateralis	CBS 136629	Soil in Eucalyptus plantation	Guangxi,China	X. Zhou & G. Zhao	KJ462840	KJ463186	KJ463070	KJ462955	Lombard et al. 2015
	CERC 8866	Soil	Central China	S.F. Chen	MF527039	MF527055	MF527071	MF527097	This study
	CERC 8841	Soil	Central China	S.F. Chen	MF527036	MF527052	MF527068	MF527094	This study
	CERC 8848	Soil	Central China	S.F. Chen	MF527037	MF527053	MF527069	MF527095	This study
	CERC 8850	Soil	Central China	S.F. Chen	MF527038	MF527054	MF527070	MF527096	This study
Ca. lichi	CERC 8871	Soil	Central China	S.F. Chen	MF527040	MF527056	MF527072	MF527098	This study
	CERC 8890	Soil	Central China	S.F. Chen	MF527041	MF527057	MF527073	MF527099	This study
	CERC 8900	Soil	Central China	S.F. Chen	MF527042	MF527058	MF527074	MF527100	This study
	CERC 8906	Soil	Central China	S.F. Chen	MF527043	MF527059	MF527075	MF527101	This study
	CERC 8928	Soil	Central China	S.F. Chen	MF527044	MF527060	MF527076	MF527102	This study
Ca. macroconidialis	CBS 114880	E. grandis	South Africa	P.W. Crous	GQ267313	DQ190655	GQ267393	AF232855	Lombard et al. 2010b
Ca. magnispora	CBS 136249	Soil in Eucalyptus plantation	Guangxi, China	X. Mou & R. Chang	KJ462841	KJ463187	KJ463071	KJ462956	Lombard et al. 2015
	CBS 112752	Soil	Indonesia	M.J. Wingfield	AY725722	AY725672	AY725760	AY725627	Lombard et al. 2010b
Ca. malesiana	CBS 112710	Debris	Thailand	N.L. Hywel-Jones	AY725721	AY725671	AY725759	AY725626	Lombard et al. 2010b

Charies	Icolate No 14	Currente	Samuling eite	Collector		GenBank ao	cession No. [%]		Reference
opposed			an guiding		tef1	his3	cmdA	tub2	
	CBS 136638	<i>E. urophylla</i> \times <i>E. grandis</i> clone seedling leaf	Guangdong, China	G. Zhao	KJ462845	KJ463191	KJ463075	KJ462960	Lombard et al. 2015
Ca. microcomataus	CBS 136633	<i>E. urophylla</i> \times <i>E. grandis</i> clone seedling leaf	Guangdong, China	G. Zhao	KJ462842	KJ463188	KJ463072	KJ462957	Lombard et al. 2015
	CERC 8952	Soil	Central China	S.F. Chen	MF527049	MF527065	MF527081	MF527107	This study
	CERC 8930	Soil	Central China	S.F. Chen	MF527045	MF527061	MF527077	MF527103	This study
	CERC 8932	Soil	Central China	S.F. Chen	MF527046	MF527062	MF527078	MF527104	This study
Ca. montana	CERC 8936	Soil	Central China	S.F. Chen	MF527047	MF527063	MF527079	MF527105	This study
	CERC 8938	Soil	Central China	S.F. Chen	MF527048	MF527064	MF527080	MF527106	This study
	CERC 8957	Soil	Central China	S.F. Chen	MF527050	MF527066	MF527082	MF527108	This study
	CERC 8966	Soil	Central China	S.F. Chen	MF527051	MF527067	MF527083	MF527109	This study
7	CPC 28835	Soil	Thailand	P.W. Crous	KT964773	N/A	KT964771	KT964769	Crous et al. 2015
Ca. monucota	CPC 28836	Soil	Thailand	P.W. Crous	KT964774	N/A	KT964772	KT964770	Crous et al. 2015
Ca. multiseptata	CBS 112682	Eucalyptus sp.	Indonesia	M.J. Wingfield	FJ918535	DQ190659	GQ267397	DQ190573	Lombard et al. 2010b
	CBS 131802	Nymphaea tetragona	Guiyang, Guizhou	S.Y. Qin	KC555273	N/A	N/A	JN984864	Xu et al. 2012
Ca. nympnaeae	HGUP 100004	N. tetragona	Guiyang, Guizhou	Y. Wang	KC555274	N/A	N/A	JN984865	Xu et al. 2012
e U	CBS 109063	Araucaria heterophylla	Hawaii, USA	M. Aragaki	AY725724	GQ267255	AY725762	GQ267213	Lombard et al. 2010b
Ca. pacifica	CBS 114038	Ipomoea aquatica	New Zealand	C.F. Hill	GQ267320	AY725675	GQ267402	AY725630	Lombard et al. 2010b
	CBS 114679	N/A	USA	A.Y. Rossman	KX784714	N/A	KX784582	KX784644	Lombard et al. 2016
ca. paracomounn	CBS 114705	Annona reticulata	Australia	D. Hutton	KX784715	N/A	N/A	KX784645	Lombard et al. 2016
Ca. parakyotensis	CBS 136085	Soil in Eucalyptus plantation	Guangdong, China	X. Mou & R. Chang	KJ462851	KJ463197	KJ463081	N/A	Lombard et al. 2015
,	CBS 136095	Soil in Eucalyptus plantation	Guangxi, China	X. Mou & R. Chang	KJ462852	KJ463198	KJ463082	N/A	Lombard et al. 2015
Ca. parva	CBS 110798	Eucalyptus grandis roots	South Africa	P.W. Crous	KX784716	N/A	KX784583	KX784646	Lombard et al. 2016
Ca. pauciramosa	CMW 5683	E. grandis	South Africa	P.W. Crous	FJ918565	FJ918531	GQ267405	FJ918514	Lombard et al. 2010b
	CMW 30823	E. grandis	South Africa	P.W. Crous	FJ918566	FJ918532	GQ267404	FJ918515	Lombard et al. 2010b
Ca. penicilloides	CBS 174.55	Prunus sp.	Japan	Tubaki	GQ267322	GQ267257	GQ267406	AF333414	Lombard et al. 2010b
Ca. pluriramosa	CBS 136976	Soil in Eucalyptus plantation	Guangxi, China	X. Zhou & G. Zhao	KJ462882	KJ463228	KJ463112	KJ462995	Lombard et al. 2015
Ca. pseudokyotensis	CBS 137332	Soil in Eucalyptus plantation	Guangxi,China	X. Zhou & G. Zhao	KJ462881	KJ463227	KJ463111	KJ462994	Lombard et al. 2015

Sneries	Icolate No 14	Culvetrate	Samuling site	Collector		GenBank acc	cession No. ^{5,}		Reference
smile	POLICIC TAG	oup name	ampine guide		tefl	his3	cmdA	tub2	
	CBS 127195	E. dumnii	FuJian, China	M.J. Wingfield	HQ285816	HQ285802	MF527091	HQ285788	Chen et al. 2011; This study
Ca. pseudocounounu	CBS 127196	E. dumnii	FuJian, China	M.J. Wingfield	HQ285817	HQ285803	MF527092	HQ285789	Chen et al. 2011; This study
	CBS 123694	E. urophylla × E. grandis cutting	Guangdong, China	M.J. Wingfield	FJ918541	FJ918519	GQ267411	FJ918504	Lombard et al. 2010b
Ca. pseudoreteaudu	CBS 123696	E. urophylla × $E.$ grandis cutting	Guangdong, China	M.J. Wingfield	FJ918542	FJ918520	GQ267410	FJ918505	Lombard et al. 2010b
	CBS 112146	E. urophylla	Australia	B. Brown	FJ918543	FJ918521	GQ267415	AF389835	Lombard et al. 2010b
La. queenstanatca	CBS 112155	E. pellita	Australia	K.M. Old	FJ918544	DQ190667	GQ267416	AF389834	Lombard et al. 2010b
	CBS 112144	E. camaldulensis	Vietnam	M.J. Dudzinski	FJ918537	DQ190661	GQ267417	AF389833	Lombard et al. 2010b
Ca. reteauati	CBS 112143	E. camaldulensis	Vietnam	M.J. Dudzinski	FJ918536	DQ190660	GQ267418	GQ240642	Lombard et al. 2010b
Ca. sphaeropendun- culata	CBS 136081	Soil in Eucalyptus plantation	Guangxi, China	X. Zhou & G. Zhao	KJ462890	KJ463236	KJ463120	KJ463003	Lombard et al. 2015
Ċ	CBS 112829	Soil	Indonesia	M.J. Wingfield	AY725733	AY725696	AY725771	AY725649	Lombard et al. 2010b
Ca. sumarrensis	CBS 112934	Soil	Indonesia	M.J. Wingfield	AY725735	AY725698	AY725773	AY725651	Lombard et al. 2010b
Ca. syzygiicola	CBS 112831	Soil	Indonesia	M.J. Wingfield	KX784736	N/A	N/A	KX784663	Lombard et al. 2016
	CBS 112151	E. urophylla	Australia	C. Hanwood	FJ918545	FJ918522	GQ267451	FJ918506	Lombard et al. 2010b
Ca. terrae-reginae	CBS 112634	Xanthorrhoea australis	Australia	T. Baigent	FJ918546	DQ190668	GQ267452	FJ918507	Lombard et al. 2010b
	CBS 136077	Soil in Eucalyptus plantation	Guangxi, China	X. Zhou & G. Zhao	KJ462900	KJ463246	N/A	KJ463013	Lombard et al. 2015
Ca. mangroua	CBS 136093	Soil in <i>Eucalyptus</i> plantation	Guangxi, China	X. Mou & R. Chang	KJ462901	KJ463247	KJ463130	KJ463014	Lombard et al. 2015
	-		- - F		- -	(: 7	(י ק נ

CBS: Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands, CERC: China Eucalypt Research Centre, Zhanjiang, GuangDong Province, China, CMW: culture collection of the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, Pretoria, South Africa; CPC: Pedro Crous working collection housed at CBS; HGUP: Plant Pathology Herbarium of Gui Zhou University, GuiYang 550025, China.

Isolates represent ex-type and indicated in bold.

⁵ tef1: translation elongation factor 1-alpha; his3: histone H3; cmdA: calmodulin; tub2: β-tubulin.

GenBank accession no. indicated in bold were generated in this study

⁹ "N/A" represents information that are not available.

The tree length (TL), consistency index (CI), retention index (RI) and homoplasy index (HI) were used to assess phylogenetic trees (Hillis and Huelsenbeck 1992).

ML analyses were performed using PHYML v. 3.0 (Guindon and Gascuel 2003), and the best evolutionary model was obtained using JMODELTEST v. 2.1.5 (Posada 2008). In PHYML, the maximum number of retained trees was set to 1,000, and nodal support was determined by non-parametric bootstrapping with 1,000 replicates.

Based on the morphological characteristics, datasets were separated into two groups: the Prolate Group and the Sphaero-Naviculate Group (Lombard et al. 2010b), and therefore phylogenetic analyses were performed with two separate sequence datasets. *Calonectria hongkongensis* (CBS 114711 and CBS 114828) and *Ca. pauciramosa* (CMW 5683 and CMW 30823) represented the outgroup taxa for the Prolate Group and Sphaero-Naviculate Group, respectively. The phylogenetic trees were viewed using MEGA v. 6.0.5 for both MP and ML analyses.

Sexual compatibility

Based on multi-gene phylogenetic analyses, isolates of each identified *Calonectria* species were crossed with each other in all possible combinations. Crosses were performed on minimal salt agar (MSA; Guerber and Correll 2001) on the surface of the medium using three sterile toothpicks. Isolates crossed with themselves were regarded as controls. These crosses were used to determine whether the identified species had a heterothallic or a homothallic mating system. The cultures were incubated at 25 °C for six weeks. When isolate combinations produced extruding viable ascospores, crosses were considered successful.

Morphology

To determine the morphological characteristics of the asexual morphs, representative isolates identified by DNA sequence comparisons were selected. Agar plugs from the periphery of actively growing single conidial cultures were transferred onto synthetic nutrient-poor agar (SNA; Nirenburg 1981) and incubated at 25 °C for one week (there were five replicates per isolate). Asexual structures that emerged on the surface of the SNA medium were mounted in one drop of 80% lactic acid on glass slides and examined under an Axio Imager A1 microscope (Carl Zeiss Ltd., Munchen, Germany) and an AxioCam ERc 5S digital camera with Zeiss Axio Vision Rel. 4.8 software (Carl Zeiss Ltd., Munchen, Germany). Sexual morphs were studied by transferring perithecia obtained from the sexual compatibility tests into a tissue-freezing medium (Leica Biosystems, Nussloch, Germany) and were hand-sectioned using an HM550 Cryostat Microtome (Microm International GmbH, Thermo Fisher Scientific, Wall-

dorf, Germany) at -20 °C. The 10-µm sections were mounted in 80% lactic acid and 3% KOH.

Fifty measurements were made for each morphological structure of the isolates selected as the holotype specimen, 30 measurements were made for the isolates selected as the paratype specimen. Minimum, maximum and average (mean) values were determined and presented as follows: (minimum–) (average – standard deviation) – (average + standard deviation) (–maximum).

The optimal growth temperature of the *Calonectria* species was determined by transferring the representative isolates to fresh 9 mm MEA Petri dishes, which were incubated under temperatures ranging from 5 to 35 °C at 5 °C intervals in the dark (there were five replicates per isolate). Colony colors were determined by inoculating the isolates on fresh MEA at 25 °C in the dark, after seven days incubation, a comparison was performed using the colour charts of Rayner (1970).

Results

Fungal isolates

A total of 40 isolates with the typical morphological of *Calonectria* species were obtained from the infected alfalfa tissue cultivated in the soil samples. Based on preliminary phylogenetic analysis of the *tef1* gene region (data not shown), 16 isolates from all soil samples were selected for further study (Table 1).

Phylogenetic analyses

Sequences for the 78 ex-type and other strains of 48 *Calonectria* species closely related to isolates obtained in this study were downloaded from GenBank (Table 1). For the 16 isolates collected in this study, nine resided in the Prolate Group, and seven were clustered in the Sphaero-Naviculate Group. Phylogenetic analyses of individual *tef1*, *his3*, *cmdA* and *tub2* and the combined sequence datasets were conducted using both MP and ML method. For both the Prolate and Sphaero-Naviculate Groups, although the related position of some *Calonectira* species were slightly different between the MP and ML trees, the overall topologies were similar, and the ML trees were exhibited.

For the Prolate and Sphaero-Naviculate Groups, the PHT comparing the combined *tef1*, *his3*, *cmdA* and *tub2* gene datasets generated P values of 0.141 and 0.333, respectively, which indicated that no significant difference existed between these datasets. These datasets were consequently combined and subjected to phylogenetic analyses. For each of the two groups, the sequence alignments of *tef1*, *his3*, *cmdA*, *tub2* and the combination of the four genes were deposited in TreeBASE (TreeBASE No. 21357). The number of parsimony informative characters, the statistical values for the

	,					Maxim	um narsin	VUOL		
Dataset	Phylogenetic group	No. of taxa	No. of bp ^T	PIC [‡]	No. of trees	Tree length	CI§	RI	RC	HI#
tefl	Prolate	45	515	210	8	448	0.7054	0.8847	0.6240	0.2946
his3	Prolate	38	449	140	6	340	0.6941	0.9176	0.6369	0.3059
cmdA	Prolate	42	476	152	792	245	0.7591	0.9295	0.7056	0.2408
tub2	Prolate	45	579	204	18	350	0.8085	0.9395	0.7597	0.1914
tef1 his3 cmdA tub2	Prolate	45	2019	706	1	1484	0.6880	0.8940	0.6150	0.3120
tefl	Sphaero-Naviculate	51	522	159	33	330	0.7030	0.9056	0.6367	0.2969
his3	Sphaero-Naviculate	47	455	138	11	386	0.6632	0.9110	0.6042	0.3367
cmdA	Sphaero-Naviculate	49	473	138	48	228	0.7763	0.9406	0.7302	0.2236
tub2	Sphaero-Naviculate	47	534	174	4	401	0.7107	0.9216	0.6550	0.2892
tef1 his3 cmdA tub2	Sphaero-Naviculate	51	1984	609	1350	1535	0.6190	0.8790	0.6047	0.3810
Dataset	Phylogenetic groun				Maximuı	m likelihood				
	Anna Bronch Bronk	Subst. model ^{††}	NST ^{##}			Rate ma	atrix			Rates
tefl	Prolate	TIM2+G	6	1.6588	2.3553	3 1.65	88	1.0000	4.4652	Gamma
his3	Prolate	GTR+G	6	1.8190	7.5654	é 4.62	281	1.4320	15.6259	Gamma
cmdA	Prolate	HKY+G	2							Gamma
tub2	Prolate	TPM3uf+G	6	1.5151	4.2112	2 1.00	000	1.5151	4.2112	Gamma
tef1 his3 cmdA tub2	Prolate	TIM2+I+G	9	1.3725	3.6221	1.37	725	1.0000	5.1226	Gamma
tefl	Sphaero-Naviculate	GTR+G	6	2.3612	2.5155	0.62	27	0.7074	5.0226	Gamma
his3	Sphaero-Naviculate	HKY+I+G	2							Gamma
cmdA	Sphaero-Naviculate	TrN+G	6	1.0000	3.8308	3 1.00	000	1.0000	6.4755	Gamma
tub2	Sphaero-Naviculate	TPM3uf+G	6	1.5714	4.605	5 1.00	000	1.5714	4.6055	Gamma
tef1 his3 cmdA tub2	Sphaero-Naviculate	GTR+I+G	6	1.6318	3.813() 1.08	888	1.1609	5.2579	Gamma
† bp = base pairs.			${}^{\circ}RC = r_{0}$	escaled consist	tency index.					

Table 2. Statistics resulting from phylogenetic analyses.

[§] CI = consistency index. RI = retention index.

[‡] PIC = number of parsimony informative characters.

^{††} Subst. model = best fit substitution model. # HI = homoplasy index.

 $^{\text{\tiny \#}}$ NST = number of substitution rate categories.

phylogenetic trees of the MP analyses, and the parameters for the best-fit substitution models of ML analyses are shown in Table 2.

Phylogenetic analyses of each of the individual and combined sequence datasets indicated that in the Prolate Group, the nine isolates resided in the Ca. colhounii species complex and were closely related to Ca. colhounii, Ca. eucalypti, Ca. fujianensis, Ca. nymphaeae, Ca. paracolhounii and Ca. pseudocolhounii. In the his3 and cmdA phylogenetic trees, the nine isolates and Ca. fujianensis were clustered in the same clade (Suppl. materials 2, 3), while in the trees based on the *tef1* and *tub2* sequences, the nine isolates formed an independent clade (Supplementary Figures 1, 4). Based on the phylogenetic analyses of the combined sequences of the four genes, the nine isolates formed a new, strongly defined phylogenetic clade that was distinct from other Calonectria species and was supported by high bootstrap values (ML = 94%, MP = 93%) (Figure 1). Fixed unique single nucleotide polymorphisms (SNPs) were identified in the new phylogenetic clades of the nine isolates and their phylogenetically closed Calonectria species (Table 3). The total number of SNP differences between the new clade and the other closely related species varied between 10-34 for all four gene regions combined (Table 4). The results of these phylogenetic and SNP analyses indicate that the nine isolates in the Prolate Group represent a distinct, undescribed species.

Phylogenetic analyses of each of the individual and combined datasets indicated that in the Sphaero-Naviculate Group, the seven isolates were clustered in the *Ca. kyotensis* species complex and were closely related to *Ca. canadiana*. In the *tef1* phylogenetic trees, the seven isolates were grouped in the same clade with *Ca. canadiana* (Suppl. material 5). In the phylogenetic trees based on the *his3, cmdA* and *tub2* sequences, the seven isolates formed an independent clade distinct from *Ca. canadiana* and other species in the *Ca. kyotensis* species complex (Suppl. materials 6, 7 and 8). Based on the combined sequences of the four genes, the seven isolates formed a strongly defined phylogenetic clade that was distinct from *Ca. canadiana* and was supported by high bootstrap values (ML = 100%, MP = 100%) (Figure 2). The seven isolates obtained in this study were distinguished from *Ca. canadiana* using SNP analyses for each of the *tef1, his3, cmdA* and *tub2* gene region sequences (Tables 5). The total number of SNP differences between the seven isolates and *Ca. canadiana* for all four genes was 51 (Table 6). The results indicate that the seven isolates in the Sphaero-Navivulate Group represent a novel species.

Sexual compatibility

After a six-week mating test on MSA, all 16 isolates and the crosses of isolates of each identified species failed to yield sexual structures, indicating that they were either self-sterile (heterothallic) or had retained the ability to recombine to produce fertile progeny.



Figure 1. Phylogenetic tree of *Calonectria* species in the Prolate group based on maximum likelihood (ML) analysis of combined DNA dataset of *tef1*, *his3*, *cmdA* and *tub2* gene sequences. ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in blue. The tree was rooted to *Ca. hongkongensis* (CBS 114711 and CBS 114828).

	ci.
•	
	5
	8
	Ã,
_	S
	2
	ž
-	3
	Ľ
	÷
	õ
	S
-	3
	5
2	£`
	g
	Ĕ.
	Ð
	8
	ക
	<u> </u>
	2
-	5
	ĕ
	-
-	g
	an a
	1
~	9
	μı
	7
•	11.1
	Ct
	1CI
	03
-	21
(3
	2
	e
	9
	B
	ē
-	
	S
	ō
•	50
	<u>5</u>
	N A
	ă
	gene
	r gene
	ur gene
,	tour gene
ر	n tour gene
ر.	in tour gene
ر	ns in four gene
,	ons in four gene
	isons in four gene
,	arisons in four gene
	parisons in four gene
ر	mparisons in four gene
	comparisons in four gene
	comparisons in four gene
ر	m comparisons in four gene
	ism comparisons in four gene
	ohism comparisons in four gene
	rphism comparisons in four gene
	torphism comparisons in four gene
· · ·	morphism comparisons in four gene
· ·	lymorphism comparisons in four gene
· · ·	olymorphism comparisons in four gene
	: polymorphism comparisons in four gene
· · ·	le polymorphism comparisons in four gene
· · · · · · · · · · · · · · · · · · ·	ide polymorphism comparisons in four gene
	otide polymorphism comparisons in four gene
	leotide polymorphism comparisons in four gene
· · · · · ·	icleotide polymorphism comparisons in four gene
J	nucleotide polymorphism comparisons in four gene
· · · · · · · · · · · · · · · · · · ·	e nucleotide polymorphism comparisons in four gene
· · · · · · · · · · · · · · · · · · ·	sle nucleotide polymorphism comparisons in four gene
	ngle nucleotide polymorphism comparisons in tour gene
	single nucleotide polymorphism comparisons in tour gene
	 Single nucleotide polymorphism comparisons in four gene
· · · · · · · · · · · · · · · · · · ·	5. Single nucleotide polymorphism comparisons in four gene
· · · · · · · · · · · · · · · · · · ·	e 3. Single nucleotide polymorphism comparisons in tour gene
· · · · · · · · · · · · · · · · · · ·	JIE 3. Mugle nucleotide polymorphism comparisons in four gene

										tej	T^{\dagger}								
opecies	Isolate no.	28 [‡]	81	89	90	91	92	93	100	120	121	124	184	185	186	243	418	425	432
	CERC 8866[§]	Α	A	I	I	Ι	I	I	C	Τ	Τ	Α	I	I	I	Α	C	I	A
	CERC 8841	Α	Α	I	Ι	Ι	Ι	I	U	Н	Н	A	I	I	I	Α	U	I	Α
	CERC 8848	Α	Α	Ι	Ι	Ι	Ι	I	C	Ц	Τ	Α	I	I	I	Α	C	I	Α
	CERC 8850	Α	Α	I	Ι	Ι	Ι	I	С	Τ	Т	Α	I	I	I	Α	С	I	Α
Ca. lichi	CERC 8871	Α	Α	I	Ι	Ι	Ι	I	U	Н	Ц	A	I	I	I	Α	U	I	A
	CERC 8890	Α	Α	Ι	Ι	Ι	Ι	Ι	C	Ļ	Τ	Α	I	I	I	Α	С	I	Α
	CERC 8900	Α	Α	Ι	Ι	Ι	I	Ι	U	H	Т	Α	I	I	I	Α	С	I	Α
	CERC 8906	Α	Α	Ι	Ι	Ι	Ι	I	С	Т	Τ	Α	I	I	I	Α	С	I	Α
	CERC 8928	А	Α	I	I	I	I	I	С	Τ	Т	А	I	I	I	А	С	I	А
	CBS 293.79	C	Ţ	Υ	C	Α	Α	C	C	I	I	Α	I	I	I	G	С	I	Α
Ca. comounu	CBS 114704	С	Τ	Α	С	Α	Α	С	C	I	I	Α	I	I	I	G	С	I	Α
	CBS 125275	Α	H	I	Ι	Ι	Ι	I	U	Н	I	A	I	I	I	Α	U	I	A
Ca. eucarypu	CBS 125276	Α	Τ	I	Ι	Ι	Ι	I	С	Τ	I	Α	I	I	I	Α	С	I	Α
	CBS 127201	А	Τ	I	I	I	I	I	С	Τ	I	G	Α	Α	Α	А	С	I	А
Ca. Jujtanensis	CBS 127200	А	Τ	Ι	Ι	Ι	I	Ι	C	Г	I	G	Α	Α	Α	А	C	I	Α
	CBS 131802	А	Τ	I	I	I	I	I	С	Τ	I	G	I	I	I	А	С	I	А
ca. nympnaeae	HGUP 100004	А	Τ	Ι	Ι	Ι	Ι	Ι	С	Τ	I	G	I	I	I	А	С	I	А
Ca. paracolhounii	CBS 114679	Α	Τ	I	Ι	Ι	I	I	Т	Н	Т	Α	I	I	I	Α	IJ	C	C
	CBS 127195	Α	H	Ι	I	I	I	I	U	H	I	Α	I	I	I	Α	U	I	Α
ca. pseudocomounti	CBS 127196	A	H	Ι	Ι	I	I	I	U	T	I	A	I	I	I	A	U	I	Α

ntinue.
õ
ň
e
ab

Table 3. Continue.																		
										tefl								
opecies	Isolate no.	433	435	436	437	438	441	443	444	446	447	448	450	452	453	457	473	483
	CERC 8866[§]	Τ	Τ	С	Τ	С	Т	Τ	Α	С	Τ	Α	С	Τ	Τ	Τ	G	I
	CERC 8841	Н	H	U	H	U	Н	H	Α	U	Н	Α	υ	H	H	Н	IJ	I
	CERC 8848	Н	Н	υ	Τ	υ	Г	Н	Α	U	Н	Α	υ	Г	Г	Н	IJ	I
	CERC 8850	Н	H	C	H	U	Г	H	Α	C	Н	Α	υ	H	H	Н	IJ	I
Ca. lichi	CERC 8871	Н	H	υ	H	υ	H	H	Α	U	H	Α	υ	H	H	H	IJ	I
	CERC 8890	Н	H	C	H	U	Г	H	Α	C	Н	Α	υ	H	H	Н	IJ	I
	CERC 8900	Н	H	υ	H	υ	H	H	Α	υ	H	Α	υ	H	H	Н	IJ	I
	CERC 8906	Г	Г	C	T	U	Г	Г	Α	U	Н	Α	υ	Τ	Г	Ч	IJ	I
	CERC 8928	Н	H	U	H	U	Н	H	Α	U	Н	Α	υ	H	H	Н	IJ	I
	CBS 293.79	Τ	Ţ	C	C	U	С	Ţ	Α	С	Ţ	Α	U	U	Τ	C	G	C
Ca. comount	CBS 114704	Τ	Τ	С	С	С	С	Τ	Α	С	Τ	А	С	С	Τ	С	G	С
	CBS 125275	Н	H	υ	H	υ	H	H	Α	U	H	Α	υ	H	H	H	G	U
Ca. eucarypu	CBS 125276	Τ	Τ	С	Т	С	Т	Τ	Α	С	Τ	А	С	Τ	Τ	Τ	G	С
	CBS 127201	Τ	Τ	С	Τ	С	Т	Τ	А	С	Τ	А	С	Τ	Τ	Τ	G	I
Ca. Jujianensis	CBS 127200	Н	Τ	C	Τ	C	Τ	Н	Α	С	T	А	U	Н	Г	Ţ	G	I
	CBS 131802	Τ	Τ	С	Τ	С	Τ	Τ	А	С	Τ	А	С	I	Τ	Τ	I	N/A ⁵
ca. nympnaeae	HGUP 100004	Н	Η	C	Τ	U	Τ	Н	А	С	T	А	U	I	Г	Ţ	I	N/A
Ca. paracolhounii	CBS 114679	Α	IJ	I	ı	Т	Н	C	Т	IJ	IJ	IJ	T	IJ	IJ	N/A	N/A	N/A
Ca providence lance	CBS 127195	H	Г	С	Τ	C	H	Г	Α	C	Н	А	С	Г	H	H	G	I
ca. pseudocomounii	CBS 127196	L	H	U	Τ	υ	T	L	A	υ	Τ	A	υ	Ţ	Ţ	H	IJ	I

tinue.
. Con
m a
Table

															'							
	T1-1-				His3										cmdA							
opecies	Isolate no.	45	234	272	293	344	353	368	169	204 2	205	010	238 2	2 44 2	66 2	93 3	25 3	34 4]	11 4	29 43	2 47	4
	CERC 8866[§]	A	Г	A	C	U	C	А	IJ	A	U	U	IJ	۔ ن	י ניז	A	ں ں		0	0	L	L
	CERC 8841	A	H	Α	U	υ	U	А	IJ	A	U	υ	IJ	ں ن	י ניז	Ā	ں ں	() ()	0	0	L	
	CERC 8848	A	Н	A	C	U	U	А	IJ	A	U	U	IJ	۔ ن	() ()	A	ں ں		0		L	
	CERC 8850	A	Н	A	C	U	U	А	IJ	A	C	U	IJ	۔ ن	י ניז	A	ں ں	() ()	0	0	L	L
Ca. lichi	CERC 8871	Α	H	Α	U	U	U	A	IJ	A	υ	U	IJ	۔ ن	י ניז	A	ں ں	() ()	0	0	L	
	CERC 8890	A	Н	A	C	U	U	А	IJ	A	U	U	IJ	۔ ن	י ניז	A	U	() ()	0	0	L	г.,
	CERC 8900	A	H	Α	U	υ	U	А	IJ	A	υ	υ	IJ	ں ن	י ניז	Ā	ں ں	() ()	0	0	L	
	CERC 8906	A	Н	A	C	U	U	А	IJ	A	U	U	IJ	۔ ن	י ניז	A	ں ں		0		L	
	CERC 8928	A	H	Α	U	υ	U	А	IJ	A	U	υ	IJ	ں ن	י ניז	Ā	ں ں	() ()	0	0	L	
	CBS 293.79	Α	H	Α	H	C	L	C	IJ	A	C	U	IJ	U U	י ניז	- V	U	о Сл	0	L U		()
Ca. comounti	CBS 114704	A	H	Α	H	U	H	U	IJ	A	U	U	IJ	۔ ن	י ניז	A	ں ں	() ()	0	L		()
	CBS 125275	I	H	H	H	H	H	υ	IJ	A	υ	υ	IJ	۔ ن	י ניז	۲ ۲	U U	() ()	0	L D		()
Ca. eucarypti	CBS 125276	Ι	H	Г	Τ	Т	L	С	G	A	С	С	G	G G	י ניז	A	י ט) ()	0	T D		(1)
	CBS 127201	A	H	Α	U	υ	U	А	IJ	A	υ	υ	IJ	ں ن	י ניז	Ā	ں ں	() ()	0	0	L	
Ca. Jujianensis	CBS 127200	A	Н	A	C	U	U	А	IJ	A	U	U	IJ	۔ ن	י ניז	A	ں ں		0	0	L	
	CBS 131802	N/A	N/A	N/A	N/A	N/A	N/A	V/A	N/A	N/A	V/A	V/A	V/A	V/A N	A N	A N	I/A N	/A N	A N	/A N/	A N/	A
Ca. nympnaeae	HGUP 100004	N/A	N/A	N/A	N/A	N/A	N/A 1	V/A	N/A 1	N/A 1	V/A 1	V/A 1	V/A 1	V/A N	A N	A N	I/A N	/A N	A N	/A N/	A N/	Ā
Ca. paracolhounii	CBS 114679	N/A	N/A	N/A	N/A	N/A	N/A 1	N/A	T	T	G	A	С	C	\ ۲	Г	C 7	1 (()		N N	A/
	CBS 127195	A	C	Α	H	C	H	U	IJ	A	C	C	IJ	ں ن	י ניז	A	U		()	T D	,	L
са. р <i>seuaocomounu</i>	CBS 127196	А	C	Α	H	С	H	C	G	A	С	С	G	IJ	י ניז	A	י ט	0	()	C T	Ľ	L

Table 3. Continue.

	-									tul	52								
opecies	Isolate no.	24	28	33	68	98	103	427	442	446	455	534	535	536	537	541	547	550	551
	CERC 8866 [§]	U	G	C	C	А	U	C	U	Г	Α	IJ	Н	IJ	U	Г	C	Н	C
	CERC 8841	υ	IJ	C	υ	А	υ	υ	υ	Н	Α	IJ	Н	IJ	υ	Н	U	н	0
	CERC 8848	U	IJ	C	υ	А	υ	U	υ	Г	Α	IJ	Г	IJ	U	Г	U	Г	C
	CERC 8850	υ	IJ	U	υ	А	υ	υ	υ	H	Α	IJ	Н	IJ	υ	H	υ	H	U
Ca. lichi	CERC 8871	υ	G	С	С	А	C	C	C	Τ	Α	IJ	H	IJ	U	Ţ	U	H	0
	CERC 8890	U	IJ	C	υ	А	υ	υ	υ	Ţ	A	IJ	Н	IJ	υ	Ţ	U	Н	C
	CERC 8900	υ	IJ	C	C	Α	υ	υ	υ	Ţ	Α	IJ	H	IJ	U	H	U	H	U
	CERC 8906	C	IJ	C	C	А	U	C	U	Ţ	Α	IJ	H	IJ	U	H	U	H	C
	CERC 8928	υ	IJ	C	υ	А	υ	υ	υ	Н	Α	IJ	Н	IJ	υ	Н	U	н	0
:	CBS 293.79	N/A	N/A	N/A	υ	C	υ	U	υ	Г	Α	IJ	Г	IJ	U	Г	U	Г	C
Ca. colhounti	CBS 114704	N/A	N/A	N/A	υ	C	υ	υ	υ	Н	Α	IJ	Н	IJ	U	Н	U	Н	U
	CBS 125275	Τ	Α	Τ	υ	C	υ	υ	υ	C	G	IJ	Г	IJ	υ	Г	U	Н	0
ca. eucatypti	CBS 125276	H	Α	H	U	C	υ	υ	υ	c	G	IJ	Н	IJ	υ	H	U	Н	C
:	CBS 127201	υ	Α	U	υ	C	T	υ	υ	Н	Α	IJ	Г	IJ	υ	Н	U	Н	0
Ca. Jujianensis	CBS 127200	U	Α	C	U	С	T	U	U	Г	A	IJ	Н	IJ	U	Ţ	U	Н	C
	CBS 131802	υ	Α	U	υ	А	υ	υ	υ	H	Α	I	I	I	I	I	T	U	J
ca. nympnaeae	HGUP 100004	U	Α	C	υ	А	υ	υ	υ	H	A	I	I	I	I	I	T	U	G
Ca. paracolhounii	CBS 114679	N/A	N/A	N/A	Α	C	υ	υ	G	H	Α	IJ	H	IJ	υ	Ţ	υ	H	U
	CBS 127195	Т	Α	Τ	С	С	C	Т	С	Τ	Α	G	Ţ	IJ	C	Ţ	C	Ţ	C
ca. pseuaocomoumu	CBS 127196	Т	Α	Т	С	С	С	Т	С	Т	Α	G	Τ	G	С	Τ	С	Τ	С
[†] Polymorphic nucleo	tides occurring or	aly in a	ull of th	e isolat	es are s	hown,	not alle	eles tha	t partia	lly occi	ur in in	dividu	als per	phylog	enetic g	group.			

ά -L 4 2 [‡] Numerical positions of the nucleotides in the DNA sequence alignments are indicated.

[§]Ex-type isolates are indicated in bold.

Fixed polymorphisms for each group are shaded and in bold, those fixed but shared between two or more groups are only shaded. ⁴ "N/A" represents sequences that are not available.

	Ca. colhounii	Ca. eucalypti	Ca. fujianensis	Ca. nymphaeae	Ca. paracolhounii	Ca. pseudocolhounii
Ca. lichi	$22(16/3/2/1)^{\dagger}$	19(4/6/2/7)	10(6/0/4)	14(5/NA [‡] /NA/9)	34(19/NA/11/4)	13(3/4/1/5)
Ca. colhounii		19(12/3/2/2)	24(18/3/2/1)	24(15/NA/NA/9)	42(28/NA/12/2)	18(13/1/3/1)
Ca. eucalypti			22(6/6/2/8)	18(4/NA/NA/14)	45(26/NA/12/7)	11(1/4/1/5)
Ca. fujianensis				16(5/NA/NA/11)	37(23/NA/11/3)	15(5/4/1/5)
Ca. nymphaeae					32(20/NA/NA/12)	16(4/NA/NA/12)
Ca. paracolhounii						36(20/NA/12/4)
[†] The order of the four ₃	genes: total (tef1, his3, ci	ndA and $tub2$).				
[‡] "NA" represents seque	ences that are not availab	ole.				

Table 4. Number of unique alleles found in Calonectria lichi and the phylogenetically closest related species in total and in the four gene regions.

[‡] "NA" represents sequences that are not available.



Figure 2. Phylogenetic tree of *Calonectria* species in the Sphaero-Naviculate group based on maximum likelihood (ML) analysis of combined DNA dataset of *tef1*, *his3*, *cmdA* and *tub2* gene sequences. ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in orange. The tree was rooted to *Ca. pauciramosa* (CMW 5683 and CMW 30823).

canadiana.
Ч Сa.
<i>ia</i> and
i montari
alonectrie
between C
regions l
ır gene
n foi
ns ir
mparisor
rphism co
: polymo
otide
: nucle
ingle
S.
с С
Ìq
Ha

$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	Species	Isolate no.	tef.	γ [†]	ő	٥٢	34	47	0,		02		csid			20		1		00	00	90
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$		CEDC 6053%		49/	07	67	7	4/				5	7		67 (°C U	1 /0		י כ	ΩΩ □ 19	- 00	2
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$		CFRC 8930							ן טע		ן טע								ן סיט			ار
CENC 8936 C		CERC 8932		1				1	, U		, 0						1		, U			0
CERC 8938 C	nontana	CERC 8936	υ	1	υ	υ	υ	1	IJ	U U	IJ	0	U	υ	U	υ		0 0	U	A	L	U
$ \begin{array}{ $		CERC 8938	υ	I	υ	υ	U	1	IJ	U U	ڻ ن	U U	U U	U	U	U U	1	0 0	U	A	Г	U
CERC 8966 C - C		CERC 8957	υ	1	υ	υ	υ	1	IJ	U U	ۍ ا	U U	U U	U	U	U U	1	0 0	U	A	Ц	U
emaine CBS 110817 - T G T A T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A C T A T T G C		CERC 8966	υ	1	υ	υ	U		IJ	U	J	C	U	U	C	U		0 0	U	A	Г	IJ
pecies Iolate no. isit it it it	anadiana	CBS 110817	I	Ļ	ს	IJ	Ţ	A	Ţ	J	A	Ţ	Ţ	IJ	Ļ	A	A	۹ ۲	U	ر ۲	A	F
Ppcdrs Mondre IIO 199 202 213 213 220 237 330 331 337 339 372 378 397 400 403 418 421 CERC 8932 C												bis3										
CENC 8952 C	opecies	Isolate no.	199	202	205	212	213	220	27 2	29 2	57 3	00	121	336	339 3	372 3	378 3	97 4	00 4	03 4	18 4	21
CERC 8930 C		CERC 8952	υ	υ	υ	U	Τ	U	C	U U	U U	U U	U	U	L	U	1	0 0	U U	U U	L	b
Instructional conditional conditionale conditinal conditional conditional conditional conditional condi		CERC 8930	υ	υ	υ	υ	Ţ	U	C	C	U U	U U	U	U	L	U U	1	0 0	U U	U U	Г	U
nontana CERC 8936 C		CERC 8932	υ	υ	υ	υ	T	υ	U U	U U	U U	U U	U U	U	L	U U	1	0 0	U U	U U	Г	υ
CERC 8938 C	nontana	CERC 8936	υ	υ	U	υ	Ţ	U	C	C	U U	C	U U	U	н	U U	1	0 0	U U	U U	Г	U
CERC 8957 C		CERC 8938	υ	υ	υ	υ	T	υ	U U	U U	U U	U U	U U	U	L	U U	1	0 0	U U	U U	Г	0
CERC 8966 C C C T C		CERC 8957	υ	υ	υ	υ	Ţ	U	U	U	U U	U	U	U	Ļ	U	1	0 0	U	U U	Ц	U
anadiana CBS 110817 T A T G G T G G T G G T A C G G T A C G G T A C G G T C G T C G T G T C G T G G T G T G G T G G T G G T G		CERC 8966	υ	υ	υ	υ	Ţ	с С	с С	C	U U	U U	U U	U U	Ļ	с U	1	- 	U U	U U	Ц	υ
pecies Isolate no. cm/d1	anadiana	CBS 110817	Ţ	A	Ţ	IJ	IJ	A	IJ	Ţ	IJ	G	A	A	С	G	G	, T	Ţ	-	(7	A
pectes Isolate Ito. 470 3 4 7 10 174 181 336 403 439 502 CERC 8952 T G T C C T C C C C C C C C C C			cmdA					tu	b2													
CERC 8952 T G T C	pecies	Isolate no.	470	æ	4	~	10	174	181	336	403	439	502	I								
CERC 8930 T G T C		CERC 8952	Ţ	ს	T	C	L	C	L	C	C	Τ	C	I								
Description CERC 8932 T G T C		CERC 8930	H	ს	Ţ	υ	H	υ	H	υ	υ	Ţ	υ	I								
nontana CERC 8936 T G T C		CERC 8932	Ţ	IJ	Τ	C	H	C	T	C	c	Ţ	C									
CERC 8938 T G T C	nontana	CERC 8936	Τ	IJ	T	C	L	C	Τ	C	C	Ţ	C									
CERC 8957 T G T C		CERC 8938	T	IJ	H	U	H	U	H	U	U	H	U									
CERC 8966 T G T C		CERC 8957	Τ	IJ	T	C	L	C	Τ	C	C	T	C									
anadiana CBS 110817 C A C G C T C T A C T A		CERC 8966	Τ	IJ	Τ	C	L	C	Τ	C	C	Τ	C									
	anadiana	CBS 110817	C	A	U	IJ	U	H	υ	Ţ	Α	υ	H									

Polymorphic nucleotides occurring only in all of the isolates are shown, not alleles that partially occur in individuals per phylogenetic group. ‡ Numerical positions of the nucleotides in the DNA sequence alignments are indicated. [§]Ex-type isolates are indicated in bold. **Table 6.** Number of unique alleles found in *Calonectria montana* and *Ca. canadiana* in total and in the four gene regions.

	Ca. canadiana
Ca. montana	51(2/38/1/10)†

[†] The order of the four genes: total (*tef1*, *his3*, *cmdA* and *tub2*).

Taxonomy

Based on DNA sequence comparisons, the 16 isolates collected in this study presented two strongly defined phylogenetic clades in both the Prolate Group and the Sphaero-Naviculate Group. Morphological differences were observed between each phylogenetic clade and its phylogenetically closed species, especially with respect to the size of the macroconidia (Table 7). Based on the phylogenetic analyses, as well as morphological characteristics, the fungi isolated from the soil in this study represent two novel species of *Calonectria*, they are described as follows:

Calonectria lichi Q.L. Liu & S.F. Chen, sp. nov.

MycoBank MB821348 Figure 3

Etymology. *lichi*, which is *Calonectria* in Chinese.

Diagnosis. *Calonectria lichi* differs from the phylogenetically closely related species *Ca. colhounii*, *Ca. eucalypti*, *Ca. fujianensis*, *Ca. nymphaeae*, *Ca. paracolhounii* and *Ca. pseudocolhounii* with respect to the macroconidia dimensions.

Type. CHINA. From soil under a natural forest in central China, 07 April 2016, ShuaiFei Chen, CSFF 2019 – holotype, CERC 8866 = CGMCC 3.18733 – ex-type culture.

Description. Sexual morph unknown. Macroconidiophores consisting of a stipe, a suite of penicillate arranged fertile branches, a stipe extension, and a terminal vesicle; stipe septate, hyaline, smooth, $(39.5-)78.5-160.5(-206.5) \times (4.5-)5.5-7.5(-8.5)$ µm; stipe extension septate, straight to flexuous, (124-)139.5-187.5(-218) µm long, 2.5–5 µm wide at the apical septum, terminating in a clavate vesicle, (3.5-)4-5(-5.5) µm diam, lateral stipe extensions (90° to main axis) absent. Conidiogenous apparatus (44–)56–92(–108.5) µm long, (35-)52-82.5(-94) µm wide; primary branches aseptate to 1–septate, $(12-)16.5-33.5(-46.5) \times (4-)4.5-6.5(-9)$ µm; secondary branches aseptate, $(7-)9.5-16(-21) \times (3-)3.5-5(-6)$ µm; tertiary branches aseptate, $(7.5-)9-12.5(-14.5) \times (3-)3.5-4.5(-6)$ µm; each terminal branch producing 2–4 phialides; phialides doliiform to reniform, hyaline, aseptate, $(6-)8-12(-14.5) \times (2.5-)3-4(-5)$ µm, apex with minute periclinal thickening and inconspicuous collarette. Macroconidia cylindrical, rounded at both ends, straight, $(53-)60.5-70.5(-79) \times (5-)5.5-6.5(-7)$ µm (av. = 65.7×6 µm), 3–septate, lacking a visible abscission scar,

Sneices	Macroconidia	Macroconidia average	Macroconidia	Vesicle	Vecicle chane	Reference
operation	$(\mathbf{L} \times \mathbf{W})^{\dagger,\pm}$	$(\mathbf{L} \times \mathbf{W})^{\dagger,\pm}$	septation	(Min. – Max.) ^{†,§}	Acateric attache	
Ca. lichi	$(53-)60.5-70.5(-79) \times (5-)5.5-6.5(-7)^{4}$	65.7 × 6	3	(3.5-)4-5(-5.5)	clavate	This study
Ca. colhounii	$(45-)60-70(-80) \times (4-)5(-6)$	65 × 5	(1-)3	3-4	clavate	Crous 2002
Ca. eucalypti	$(66-)69-75(-80) \times (5-)6$	72 × 6	<i>c</i> c	4–6	broadly clavate	Lombard et al. 2010b
Ca. fujianensis	$(48-)50-55(-60) \times (2.5-)3.5-4.5(-5)$	52.5×4	(1-)3	(3-)3.5-4.5(-5)	clavate	Chen et al. 2011
Ca. nymphaeae	$55-63 \times 5.3-6.3$	61×5.9	3-4	3-5	clavate	Xu et al. 2012
Ca. paracolhounii	$(37-)39-43(-45) \times 4-5$	41×5	3	3-5	narrowly clavate	Lombard et al. 2016
Ca. pseudocolhounii	$(49-)55-65(-74) \times (3.5-)4-5(-5.5)$	60×4.5	(1-)3	(3.5-)4-5(-6)	clavate	Chen et al. 2011
Ca. montana	$(37.5-)40.5-45.5(-51.5) \times 4-5(-5.5)$	43.2×4.6	1	(4-)7-11(-12.5)	sphaeropedunculate	This study
Ca. canadiana	(38-)48-55(-65) × 4(-5)	50×4	1	6-10	pyriform to sphaeropedunculate	Kang et al. 2001; Lechat et al. 2010

Table 7. Morphological comparisons of Calonectria lichi, Ca. montana and their phylogenetically closely related species.

^{\dagger} All measurements are in μ m.

[§] Min.–Max. = minimum–maximum. ‡ L × W = length × width.

Species indicated in bold are described in this study.

⁴ Measurements are presented in the format [(minimum-) (average – standard deviation) – (average + standard deviation) (-maximum)].



Figure 3. *Calonectria lichi.* **a–c** Macroconidiophore **d–f** Clavate vesicles **g–i** Conidiogenous apparatus with conidiophore branches and doliiform to reniform phialides **j–k** Macroconidia Scale bars: **a–c** = 50 μ m; **d–f** = 5 μ m; **g–k** = 10 μ m.

held in parallel cylindrical clusters by colorless slime. Megaconidia and microconidia not observed.

Culture characteristics. Colonies forming abundant white aerial mycelium on MEA at 25 °C after seven days, with feathery, irregular margins at the edges, moderate sporulation. Surface with white to buff outer margins, and salmon (13'd) inner region, becoming ochreous (44) towards the center, reverse sienna (8) to umber (9) with abundant chlamydospores throughout the medium, forming microsclerotia. Optimal

growth temperature at 25 °C, no growth at 5 °C and 35 °C, after seven days, colonies at 10 °C, 15 °C, 20 °C, 25 °C and 30 °C reached 21.9 mm, 30.8 mm, 41.5 mm, 54.4 mm and 37.2 mm, respectively.

Substratum. Soil in a natural forest.

Distribution. Central China.

Other specimens examined. CHINA. From soil in a natural forest in central China, 07 April 2016, ShuaiFei Chen, CSFF 2020, culture CERC 8850 = CGMCC 3.18732; CHINA. From soil under a natural forest in central China, 07 April 2016, ShuaiFei Chen, CSFF 2021, culture CERC 8890 = CGMCC 3.18734; CHINA. From soil in a natural forest in central China, 07 April 2016, ShuaiFei Chen, culture CERC 8841, CERC 8848, CERC 8871, CERC 8900, CERC 8906 and CERC 8928.

Notes. *Calonectria lichi* is a new species in the *Ca. colhounii* complex and is closely related to *Ca. colhounii*, *Ca. eucalypti*, *Ca. fujianensis*, *Ca. nymphaeae*, *Ca. paracolhounii* and *Ca. pseudocolhounii* (Crous 2002, Lombard et al. 2010b, 2016, Chen et al. 2011, Xu et al. 2012, Crous et al. 2015). The macroconidia of *Ca. lichi* (av. 65.7 × 6.0 µm) are longer and wider than those of *Ca. colhounii* (av. 65 × 5 µm), *Ca. fujianensis* (av. 52.5 × 4 µm), *Ca. nymphaeae* (av. 61 × 5.9 µm), *Ca. paracolhounii* (av. 41 × 5 µm) and *Ca. pseudocolhounii* (av. 60 ×4.5 µm), but narrower than those of *Ca. eucalypti* (av. 72 × 6µm).

Calonectria montana Q.L. Liu & S.F. Chen, sp. nov.

MycoBank MB821349 Figure 4

Etymology. *montis*, meaning mountain in Latin, referring to the location where this fungus was collected.

Diagnosis. *Calonectria montana* can be distinguished from the phylogenetically closely related species *Ca. canadiana* by the size of macroconidia.

Type. CHINA. From soil under a natural forest in central China, 07 April 2016, ShuaiFei Chen, holotype CSFF 2022, ex-type culture CERC 8952 = CGMCC 3.18735.

Description. Sexual morph unknown. Macroconidiophores consisting of a stipe, a suite of penicillate arranged fertile branches, a stipe extension, and a terminal vesicle; stipe septate, hyaline, smooth, $(30-)52-91(-123.5) \times (4-)5.5-8(-9.5)$ µm; stipe extension septate, straight to flexuous (76.5–)107–168(–211.5) µm long, (2.5–)3–4.5(–5.5) µm wide at the apical septum, terminating in a pyriform to sphaeropedunculate vesicle, (4–)7–11(–12.5) µm diam, lateral stipe extensions (90° to main axis) absent. Conidiogenous apparatus (40–)49–87.5(–102.5) µm long, (44–)62–91(–104) µm wide; primary branches aseptate to 1–septate, (14.5–)19.5–34(–55.5) × (4–)4.5–6(–7) µm; secondary branches aseptate, (11–)13.5–23(–33) × (3–)4–5(–6) µm; tertiary branches aseptate, (9–)11–15(–16.5) × (3.5–)3.5–4.5(–5) µm; each terminal branch producing 2–6 phialides; phialides doliiform to reniform, hyaline, aseptate, (8–)10.5–



Figure 4. *Calonectria montana.* **a–c** Macroconidiophores **d–f** Sphaeropedunculate vesicles **g–h** Conidiogenous apparatus with conidiophore branches and doliiform to reniform phialides **i–j** Macroconidia Scale bars: **a–c** = 20 μ m; **d–j** = 10 μ m.

12.5(–15.5) × (2.5–)3.5–4.5(–5) µm, apex with minute periclinal thickening and inconspicuous collarette. Macroconidia cylindrical, rounded at both ends, straight, (37.5–)40.5–45.5(–51.5) × 4–5(–5.5) µm (av. = 43.2 × 4.6 µm), 1–septate, lacking a visible abscission scar, held in parallel cylindrical clusters by colorless slime. Megaconidia and microconidia not observed.

Culture characteristics. Colonies forming abundant buff and wooly aerial mycelium on MEA at 25 °C after seven days, with feathery, irregular margins at the edges, sporulation moderate and more concentrated in the colony centre. Surface with buff to sienna (8) outer margins, reverse sienna (8) to umber (9), and chesnut (9'm) inner region, abundant chlamydospores throughout the medium, forming microsclerotia. Optimal growth temperature at 30 °C, no growth at 5 °C and 35 °C, after seven days, colonies at 10 °C, 15 °C, 20 °C, 25 °C and 30 °C reached 22.9 mm, 31.5 mm, 51.1 mm, 61.9 mm and 77.2 mm, respectively, this is a high-temperature species.

Substratum. Soil under the natural forest.

Distribution. Central China.

Other specimens examined. CHINA. From soil in a natural forest in central China, 07 April 2016, ShuaiFei Chen, CSFF 2023, culture CERC 8957 = CGMCC 3.18736; From soil in a natural forest in central China, 07 April 2016, ShuaiFei Chen, CSFF 2024, culture CERC 8966 = CGMCC 3.18737; From soil in a natural forest in central China, 07 April 2016, ShuaiFei Chen, culture CERC 8930, CERC 8932, CERC 8936 and CERC 8938.

Notes. *Calonectria montana* is a new addition to the *Ca. kyotensis* complex and is phylogenetically closely related to *Ca. canadiana* (Crous 2002, Crous et al. 2004, Lombard et al. 2015, 2016). The macroconidia of *Ca. montana* (av. $43.2 \times 4.6 \mu$ m) are shorter and wider than those of *Ca. canadiana* (av. $50 \times 4 \mu$ m).

Discussion

This study identified two novel species of *Calonectria* from soil in a natural forest in the temperate region of central China. The identification of the fungi was supported by DNA sequence comparisons and morphological features. The two species were named *Calonectria lichi* and *Ca. montana*.

Calonectria lichi is a new addition to the *Ca. colhounii* complex that belongs to the Prolate Group. Based on phylogenetic analyses of four gene sequences, *Ca. lichi* formed a distinct and well-supported phylogenetic clade closely related to *Ca. fujianensis, Ca. nymphaeae* and *Ca. paracolhounii*, but it can be distinguished from these species by its larger macroconidia. To date, 10 species in the *Ca. colhounii* complex have been identified and described. Other than *Ca. lichi* described in this study, the other species include *Ca. colhounii*, *Ca. eucalypti, Ca. fujianensis, Ca. macroconidialis, Ca. monticola, Ca. nymphaeae*, *Ca. paracolhounii*, *Ca. fujianensis, Ca. macroconidialis, Ca. monticola, Ca. nymphaeae, Ca. paracolhounii, Ca. parva* and *Ca. pseudocolhounii* (Crous 2002, Lombard et al. 2010b, 2016, Chen et al. 2011, Xu et al. 2012, Crous et al. 2015). Of these species, *Ca. colhounii, Ca. eucalypti, Ca. fujianensis, Ca. nymphaeae* and *Ca. pseudocolhounii* have been shown to be homothallic and always produce bright yellow perithecia (Crous 2002, Lombard et al. 2010b, Chen et al. 2010b, Chen et al. 2011, Xu et al. 2012). In China, four species in the *Ca. colhounii* complex have been reported: except for *Ca. lichi*, which was isolated from a natural forest in the temperate zone in central China, the other species, including *Ca. fujianensis, Ca. pseudocolhounii* and *Ca. nym*. *phaeae*, were previously isolated from tropical or subtropical regions in southern China (Chen et al. 2011, Xu et al. 2012).

Calonectria montana adds a new species to the *Ca. kyotensis* complex that belongs to the Sphaero-Naviculate Group. Phylogenetic analyses showed that *Ca. montana*, which formed an independent clade with a high bootstrap value, is closely related to *Ca. canadiana*. Morphological differences were observed between *Ca. montana* and *Ca. canadiana*, especially with respect to the size of the macroconidia and the shape of the vesicles (Kang et al. 2001, Crous 2002). Species in the *Ca. kyotensis* complex are characterized by having sphaeropedunculate vesicles with lateral stipe extensions on a conidiogenous apparatus (Crous et al. 2004, Lombard et al. 2010b, 2015, 2016). No lateral stipe extensions were produced by *Ca. montana*, indicating that this species is different from other species in the *Ca. kyotensis* complex. In this study, *Ca. montana* was isolated from soil in central China, 14 species residing in the *Ca. kyotensis* complex were previously reported in China, and all of them were isolated from soil in southern China (Crous et al. 2004, Lombard et al. 2015). The results from this study suggest that more species in *Ca. kyotensis* complex have yet to be discovered from China.

Species of Calonectria are important plant pathogens that can cause devastating diseases on various plant hosts worldwide, especially on horticultural, agronomic and forestry crops (Polizzi et al. 2001, 2009, Crous 2002, Saracchi et al. 2008, Chen et al. 2011, Pan et al. 2012). In China, Calonectria species have been reported as pathogens of various important agronomic and forestry crops. In agriculture, the Fabaceae and Arecaceae plant families are susceptible to infection by Calonectria species, including Ca. ilicicola, which causes black rot (CBR) of Arachis hypogaea (peanut) and Medicago sativa (Gai et al. 2012, Pan et al. 2012, Pei et al. 2015), Ca. ilicicola causes red crown rot of Glycine max (soybean) (Guan et al. 2010), and Ca. colhounii and Ca. pteridis cause leaf spot on Phoenix canariensis and Serenoa repens, respectively (Luo et al. 2009, Yang et al. 2014). In forestry, leaf blight caused by Calonectria species is considered as one of the most serious threats to Eucalyptus plantations and nurseries in southern China (Zhou et al. 2008, Lombard et al. 2010a, Chen et al. 2011). The leaf inoculations showed that all tested Calonectria species were pathogenic to the tested *Eucalyptus* clones, including the clones that are widely planted in southern China (Chen et al. 2011, Li et al. 2014a, b). These research results suggest that species of *Calonectria* need to be monitored carefully, both in agronomic crops and forests.

Accurate diagnosis of plant diseases and identification of their casual agents provide the foundation for developing effective disease management strategies (Booth et al. 2000, Crous 2002, Old et al. 2003, Vitale et al. 2013, Wingfield et al. 2015). Based on previous research results, the majority of *Calonectria* species identified and described in China were isolated from diseased plant tissues or soil under forestry plantations in subtropical and tropical regions (Crous et al. 2004, Lombard et al. 2010a, 2015, Chen et al. 2011). In this study, two novel *Calonectria* species were described, and they were isolated from soil in a natural forest in the temperate zone. The results from this study suggest that more extensive surveys need to be conducted to collect *Calonectria* in more geographic regions with different climate zones, which will help to clarify the species diversity of *Calonectria* in China.

Acknowledgments

This study was supported by the Fundamental Research Funds for the Central Non-Profit Research Institution of CAF (Project No. CAFYBB2014MA018) and the National Natural Science Foundation of China (NSFC) (Project numbers: 31622019 and 31400546). We thank LetPub (www.letpub.com) for linguistic assistance during the preparation of this manuscript.

References

- Aiello D, Cirvilleri G, Polizzi G, Vitale A (2013) Effects of fungicide treatments for the control of epidemic and exotic *Calonectria* diseases in Italy. Plant Disease 97: 37–43. https://doi. org/10.1094/PDIS-03-12-0266-RE
- Alfenas RF, Pereira OL, Jorge VL, Crous PW, Alfenas AC (2013a) A new species of *Calonectria* causing leaf blight and cutting rot of three forest tree species in Brazil. Tropical Plant Pathology 38: 513–521. https://doi.org/10.1590/S1982-56762013000600007
- Alfenas RF, Pereira OL, Ferreira MA, Jorge VL, Crous PW, Alfenas AC (2013b) *Calonectria metrosideri*, a highly aggressive pathogen causing leaf blight, root rot, and wilt of *Metrosideros* spp. in Brazil. Forest Pathology 43: 257–265. https://doi.org/10.1111/efp.12035
- Alfenas RF, Lombard L, Pereira OL, Alfenas AC, Crous PW (2015) Diversity and potential impact of *Calonectria* species in *Eucalyptus* plantations in Brazil. Studies in Mycology 80: 89–130. https://doi.org/10.1016/j.simyco.2014.11.002
- Booth TH, Jovanovic T, Old KM, Dudzinski MJ (2000) Climatic mapping to identify highrisk areas for *Cylindrocladium quinqueseptatum* leaf blight on eucalypts in mainland South East Asia and around the world. Environmental Pollution 108: 365–372. https://doi. org/10.1016/S0269-7491(99)00215-8
- Chen SF, Lombard L, Roux J, Xie YJ, Wingfield MJ, Zhou XD (2011) Novel species of *Calonectria* associated with *Eucalyptus* leaf blight in Southeast China. Persoonia 26: 1–12. https://doi.org/10.3767/003158511X555236
- Crous PW, Phillips AJL, Wingfield MJ (1991) The genera Cylindrocladium and Cylindrocladiella in South Africa, with special reference to forest nurseries. South African Forestry Journal 157: 69–85. https://doi.org/10.1080/00382167.1991.9629103
- Crous PW (2002) Taxonomy and pathology of *Cylindrocladium* (*Calonectria*) and allied genera. APS Press, St. Paul, Minnesota, USA.
- Crous PW, Groenewald JZ, Risède JM, Simoneau P, Hywel-Jones NL (2004) *Calonectria* species and their *Cylindrocladium* anamorphs: species with sphaeropedunculate vesicles. Studies in Mycology 50: 415–430.
- Crous PW, Groenewald JZ, Risède JM, Simoneau P, Hyde KD (2006) *Calonectria* species and their *Cylindrocladium* anamorphs: species with clavate vesicles. Studies in Mycology 55: 213–226. https://doi.org/10.3114/sim.55.1.213
- Crous PW, Shivas RG, Wingfield MJ, Summerell BA, Rossman AY, et al (2012) Fungal Planet description sheets: 128–153. Persoonia 29: 146–201. https://doi.org/10.3767/003158512X661589

- Crous PW, Wingfield MJ, Guarro J, Cheewangkoon R, Van der Bank M, et al (2013) Fungal Planet description sheets: 154–213. Persoonia 31: 188–296. https://doi.org/10.3767/00-3158513X675925
- Crous PW, Wingfield MJ, Le Roux JJ, Richardson DM, Strasberg D, et al (2015) Fungal Planet description sheets: 371–399. Persoonia 35: 264–327. https://doi.org/10.3767/00-3158515X690269
- Ferreira FA (1989) Patologia florestal. Principais doenças florestais no Brasil. Viçosa, Sociedade de Investigações Florestais, Viçosa, MG, Brazil. 570 pp.
- Gadgil PD, Dick MA (2004) Fungi silvicolae novazelandae: 5. New Zealand Journal of Forestry Science 34: 316–323.
- Gai Y, Deng Q, Pan R, Chen X, Deng M (2012) First Report of Cylindrocladium Black Rot of Peanut Caused by *Cylindrocladium parasiticum* (Teleomorph *Calonectria ilicicola*) in Jiangxi Province, China. Plant Disease 96: 586. https://doi.org/10.1094/PDIS-11-11-1010
- Gehesquière B, Crouch JA, Marra RE, Van Poucke K, Rys F, et al (2015) Characterization and taxonomic reassessment of the box blight pathogen *Calonectria pseudonaviculata*, introducing *Calonectria henricotiae* sp. nov. Plant Pathology 65: 37–52. https://doi.org/10.1111/ppa.12401
- Groenewald JZ, Nakashima C, Nishikawa J, Shin HD, Park JH, Jama AN, Groenewald M, Braun U, Crous PW (2013) Species concepts in *Cercospora*: spotting the weeds among the roses. Studies in Mycology 75: 115–170. https://doi.org/10.3114/sim0012
- Guan M, Pan R, Gao X, Xu D, Deng Q, Deng M (2010) First report of red crown rot caused by *Cylindrocladium parasiticum* on soybean in Guangdong, southern China. Plant Disease 94: 485. https://doi.org/10.1094/PDIS-94-4-0485B
- Guerber JC, Correll JC (2001) Characterization of *Glomerella acutata*, the teleomorph of *Colle-totrichum acutatum*. Mycologia 93: 216–229. https://doi.org/10.2307/3761619
- Guindon S, Gascuel O (2003) A simple, Fast, and Accurate Algorithm to Estimate Large Phylogenies by Maximum Likelihood. Systematic biology 52: 696–704. https://doi. org/10.1080/10635150390235520
- Hillis DM, Huelsenbeck JP (1992) Signal, noise and reliability in molecular phylogenetic analyses. Journal of Heredity 83: 189–195. https://doi.org/10.1093/oxfordjournals.jhered. a111190
- Kang J, Crous PW, Schoch CL (2001) Species concepts in the Cylindrocladium floridanum and Cy. Spathiphylli complexes (Hypocreaceae) based on multi-allelic sequence data, sexual compatibility and morphology. Systematic and Applied Microbiology 24: 206–217. https://doi.org/10.1078/0723-2020-00026
- Lechat C, Crous PW, Groenewald JZ (2010) The enigma of *Calonectria* species occurring on leaves of *Ilex aquifolium* in Europe. IMA fungus 1: 101–108. https://doi.org/10.5598/ imafungus.2010.01.02.01
- Li N, Zhao X, Liu AX, Liu H (2010) Brown spot disease of tree peony caused by *Cylindrocla-dium canadense* in China. Journal of General Plant Pathology 76: 295–298. https://doi.org/10.1007/s10327-010-0245-2
- Li GQ, Chen SF, Wu ZH, Zhou XD, Xie YJ (2014a) Preliminary Analyses on Diversity and Pathogenicity of *Calonectria* spp. on *Eucalyptus* in China. Chinese Journal of Tropical Crops 35: 1183–1191. [In Chinese]

- Li GQ, Li JQ, Liu FF, Li TH, Chen SF (2014b) Preliminary Analyses on Pathogenicity of Twelve *Calonectria* spp. on Ten *Eucalyptus* Clones in China. Eucalypt Science & Technology 31: 1–7. https://doi.org/10.13987/j.cnki.askj.2014.04.001 [In Chinese]
- Lombard L, Zhou XD, Crous PW, Wingfield BD, Wingfield MJ (2010a) *Calonectria* species associated with cutting rot of *Eucalyptus*. Persoonia 24: 1–11. https://doi.org/10.3767/003158510X486568
- Lombard L, Crous PW, Wingfield BD, Wingfield MJ (2010b) Phylogeny and systematics of the genus *Calonectria*. Studies in Mycology 66: 31–69. https://doi.org/10.3114/ sim.2010.66.03
- Lombard L, Crous PW, Wingfield BD, Wingfield MJ (2010c) Multigene phylogeny and mating tests reveal three cryptic species related to *Calonectria pauciramosa*. Studies in Mycology 66: 15–30. https://doi.org/10.3114/sim.2010.66.02
- Lombard L, Crous PW, Wingfield BD, Wingfield MJ (2010d) Species concepts in *Calonectria* (*Cylindrocladium*). Studies in Mycology 66: 1–14. https://doi.org/10.3114/sim.2010.66.01
- Lombard L, Polizzi G, Guarnaccia V, Vitale A, Crous PW (2011) *Calonectria* spp. causing leaf spot, crown and root rot of ornamental plants in Tunisia. Persoonia 27: 73–79. https://doi. org/10.3767/003158511X615086
- Lombard L, Chen SF, Mou X, Zhou XD, Crous PW, Wingfield MJ (2015) New species, hyperdiversity and potential importance of *Calonectria* spp. from *Eucalyptus* in South China. Studies in Mycology 80: 151–188. https://doi.org/10.1016/j.simyco.2014.11.003
- Lombard L, Wingfield MJ, Alfenas AC, Crous PW (2016) The forgotten *Calonectria* collection: pouring old wine into new bags. Studies in Mycology 85: 159–198. https://doi. org/10.1016/j.simyco.2016.11.004
- Luan YS, Feng L, An LJ (2006) First Report of Blueberry Leaf Spot Caused by *Cylindrocladium colhounii* in China. Plant Disease 90: 1553. http://dx.doi.org/10.1094/PD901553A
- Luo JS, Wang MS, Lin XX, Zhang YY (2009) Pathogenic identification of *Phoenix canariensis* leaf spot disease. Chinese journal of tropical crops 30: 104–107. [In Chinese]
- Mirabolfathy M, Ahangaran Y, Lombard L, Crous PW (2013) Leaf blight of *Buxus sempervirens* in northern forests of Iran caused by *Calonectria pseudonaviculata*. Studies in Mycology 85: 159–198. http://dx.doi.org/10.1094/PDIS-03-13-0237-PDN
- Nirenburg HI (1981) A simplified method for identifying *Fusarium* spp. occurring on wheat. Canadian Journal of Botany 59: 1599–1609. https://doi.org/10.1139/b81-217
- Old KM, Pham QT, Dudzinski MJ, Gibbs RJ (1999) *Eucalyptus* pathology in Vietnam. In: Proceedings of the workshop on eucalypt diseases, ACIAR, Ho Chi Minh City, Vietnam. CSIRO Forestry and Forest Products, Canberra and Forest Science Institute of Vietnam, Hanoi, 5.
- Old KM, Wingfield MJ, Yuan ZQ (2003) A manual of diseases of eucalypts in South-East Asia. Centre for International Forestry Research, Indonesia.
- Pan R, Deng Q, Xu D, Ji C, Deng M, Chen W (2012) First Report of Peanut Cylindrocladium Black Rot Caused by Cylindrocladium parasiticum in Fujian Province, Eastern China. Plant Disease 99: 890. http://dx.doi.org/10.1094/PDIS11110982
- Peerally A (1991) The classification and phytopathology of *Cylindrocladium* species. Mycotaxon 40: 323–366.

- Pei WH, Cao JF, Yang MY, Zhao ZJ, Xue SM (2015) First report of black rot of *Medicago sativa* caused by *Cylindrocladium parasiticum* (teleomorph *Calonectria ilicicola*) in Yunnan Province, China. Plant Disease 99: 890. http://dx.doi.org/10.1094/PDIS11141171PDN
- Polizzi G, Catara V (2001) First report of leaf spot caused by Cylindrocladium pauciramosum on Acacia retinodes, Arbutus unedo, Feijoa sellowiana and Dodonaea viscosa in southern Italy. Plant Disease 85: 803. https://doi.org/10.1094/PDIS.2001.85.7.803C
- Polizzi G, Vitale A, Aiello D, Castello I, Guarnaccia V, Parlavecchio G (2009) First record of crown and root rot caused by *Cylindrocladium pauciramosum* on brush cherry in Italy. Plant Disease 93: 547. http://dx.doi.org/10.1094/PDIS9350547A
- Posada D (2008) jModelTest: phylogenetic model averaging. Molecular biology and Evolution 25: 1253–1256. https://doi.org/10.1093/molbev/msn083
- Rayner RW (1970) A mycological colour chart. Commonwealth Mycological Institute and British Mycological Society. Kew, Surrey, UK.
- Rodas CA, Lombard L, Gryzenhoinf M, Slippers B, Wingfield MJ (2005) Cylindrocladium blight of Eucalyptus grandis in Colombia. Australasian Plant Pathology 34: 143–149. https://doi. org/10.1071/AP05012
- Saracchi M, Rocchi F, Pizzatti C, Cortesi P (2008) Box blight, a new disease of *Buxus* in Italy caused by *Cylindrocladium buxicola*. Journal of plant pathology 90: 581–584.
- Schoch CL, Crous PW, Wingfield BD, Wingfield MJ (2001) Phylogeny of *Calonectria* based on comparisons of β-tubulin DNA sequences. Mycological Research 105: 1045–1052. https://doi.org/10.1016/S0953-7562(08)61966-8
- Serrato-Diaz LM, Latoni-Brailowsky EI, Rivera-Vargas LI, Goenaga R, Crous PW, French-Monar RD (2013) First Report of *Calonectria hongkongensis* Causing Fruit Rot of Rambutan (*Nephelium lappaceum*). Plant Disease 97: 1117. http://dx.doi.org/10.1094/PDIS01-130008PDN
- Sharma JK, Mohanan C, Florence EJ (1984) Nursery diseases of *Eucalyptus* in Keralal. Forest Pathology 14: 77–89. http://doi.org/10.1111/j.14390329.1984.tb00156.x
- Sobers EK, Littrell RH (1974) Pathogenicity of three species of *Cylindrocladium* to select hosts. Plant Disease Reporter 58: 1017–1019.
- Swofford DL (2003) PAUP*. Phylogenetic Analysis Using Parsimony (*and other methods). V. 4.0b10. Sinauer Associates, Sunderland, Massachusetts, USA.
- Tamura K, Stecher G, Peterson D, Filipski A, Sudhir Kumar (2013) MEGA6: Molecular evolutionary genetics analysis v. 6.0. Molecular Biology and Evolution 30: 2725–2729. https:// doi.org/10.1093/molbev/mst197
- Van Burik JAH, Schreckhise RW, White TC, Bowden RA, Myerson D (1998) Comparison of six extraction techniques for isolation of DNA from filamentous fungi. Medical Mycology 36: 299–303. https://doi.org/10.1111/j.1365-280X.1998.00161.x
- Vitale A, Aiello D, Castello I, Dimartino MA, Parlavecchio G, Polizzi G (2009) Severe outbreak of crown rot and root rot caused by *Cylindrocladium pauciramosum* on strawberry tree in Italy. Plant Disease 93: 842. http://dx.doi.org/10.1094/PDIS9380842B
- Vitale A, Crous PW, Lombard L, Polizzi G (2013) *Calonectria* diseases on ornamental plants in Europe and the Mediterranean basin: an overview. Journal of Plant Pathology 95: 463–476. http://doi.org/10.4454/JPP.V95I3.007

- Wingfield MJ, Brockerhoff EG, Wingfield BD, Slippers B (2015) Planted forest health: the need for a global strategy. Science 349: 832–836. https://doi.org/10.1126/science.aac6674
- Xu JJ, Qin SY, Hao YY, Ren J, Tan P, Bahkali AH, Hyde KD, Wang Y (2012) A new species of *Calonectria* causing leaf disease of water lily in China. Mycotaxon 122: 177–185. https:// doi.org/10.5248/122.177
- Yamamoto R, Nakagawa A, Shimada S, Komatsu S, Kanematsu S (2017) Histopathology of red crown rot of soybean. Journal of General Plant Pathology 83: 23–32. https://doi. org/10.1007/s10327-016-0694-3
- Yang W, Zheng L, Wang C, Xie CP (2014) The First Report of *Calonectria pteridis* causing a Leaf Spot Disease on *Serenoa repens* in China. Plant Disease 98: 854–855. https://doi. org/10.1094/PDIS11131167PDN
- Zhou XD, Xie YJ, Chen SF, Wingfield MJ (2008) Diseases of eucalypt plantations in China: challenges and opportunities. Fungal Diversity 32: 1–7.

Phylogenetic tree of *Calonectria* species in the Prolate group based on maximum likelihood (ML) analysis of *tef1* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in blue. The tree was rooted to *Ca. hongkongensis* (CBS 114711 and CBS 114828).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Phylogenetic tree of *Calonectria* species in the Prolate group based on maximum likelihood (ML) analysis of *his3* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in blue. The tree was rooted to *Ca. hongkongensis* (CBS 114711 and CBS 114828).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/mycokeys.26.14688.suppl2

Supplementary material 3

Phylogenetic tree of *Calonectria* species in the Prolate group based on maximum likelihood (ML) analysis of *cmdA* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in blue. The tree was rooted to *Ca. hongkongensis* (CBS 114711 and CBS 114828).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Phylogenetic tree of *Calonectria* species in the Prolate group based on maximum likelihood (ML) analysis of *tub2* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in blue. The tree was rooted to *Ca. hongkongensis* (CBS 114711 and CBS 114828).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/mycokeys.26.14688.suppl4

Supplementary material 5

Phylogenetic tree of *Calonectria* species in the Sphaero-Naviculate group based on maximum likelihood (ML) analysis of *tef1* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in orange. The tree was rooted to *Ca. pauciramosa* (CMW 5683 and CMW 30823).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Phylogenetic tree of *Calonectria* species in the Sphaero-Naviculate group based on maximum likelihood (ML) analysis of *his3* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in orange. The tree was rooted to *Ca. pauciramosa* (CMW 5683 and CMW 30823).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/mycokeys.26.14688.suppl6

Supplementary material 7

Phylogenetic tree of *Calonectria* species in the Sphaero-Naviculate group based on maximum likelihood (ML) analysis of *cmdA* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in orange. The tree was rooted to *Ca. pauciramosa* (CMW 5683 and CMW 30823).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Phylogenetic tree of *Calonectria* species in the Sphaero-Naviculate group based on maximum likelihood (ML) analysis of *tub2* gene sequences

Authors: QianLi Liu, ShuaiFei Chen

Data type: molecular data

- Explanation note: ML and MP (maximum parsimony) bootstrap values (ML/MP) are shown above branches, with bootstrap values below 60 % marked with an *, and absent analysis values are marked with -. Isolates representing ex-type material are marked with "T", isolates highlighted in bold were sequenced in this study and novel species were covered in orange. The tree was rooted to *Ca. pauciramosa* (CMW 5683 and CMW 30823).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.