

New *Raffaelea* species (*Ophiostomatales*) from the USA and Taiwan associated with ambrosia beetles and plant hosts

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Abstract: *Raffaelea* (*Ophiostomatales*) is a genus of more than 20 ophiostomatoid fungi commonly occurring in symbioses with wood-boring ambrosia beetles. We examined ambrosia beetles and plant hosts in the USA and Taiwan for the presence of these mycosymbionts and found 22 isolates representing known and undescribed lineages in *Raffaelea*. From 28S rDNA and β -tubulin sequences, we generated a molecular phylogeny of *Ophiostomatales* and observed morphological features of seven cultures representing undescribed lineages in *Raffaelea* s. lat. From these analyses, we describe five new species in *Raffaelea* s. lat.: *R. aguacate*, *R. campbellii*, *R. crossotarsa*, *R. cyclohipidia*, and *R. xyleborina* spp. nov. Our analyses also identified two plant-pathogenic species of *Raffaelea* associated with previously undocumented beetle hosts: (1) *R. quercivora*, the causative agent of Japanese oak wilt, from *Cyclohipidion ohnoi* and *Crossotarsus emancipatus* in Taiwan, and (2) *R. lauricola*, the pathogen responsible for laurel wilt, from *Ambrosiodmus lecontei* in Florida. The results of this study show that *Raffaelea* and associated ophiostomatoid fungi have been poorly sampled and that future investigations on ambrosia beetle mycosymbionts should reveal a substantially increased diversity.

Key words:

entomogenous fungi
insect-fungus interactions
Japanese oak wilt
laurel wilt
molecular phylogenetics
mycosymbioses

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INTRODUCTION

Raffaelea (Arx & Hennebert 1965) is a genus of primarily asexual fungi including more than 20 species in *Ophiostomatales* (Harrington *et al.* 2010, de Beer *et al.* 2013, Musvuugwa *et al.* 2015). These fungi commonly occur in symbioses with wood-boring ambrosia beetles (*Coleoptera*: *Curculionidae*: *Scolytinae* and *Platypodinae*). Ambrosia beetles propagate these and other fungi, which obtain nutrients from plant tissues and provide the beetles with a food source, throughout galleries in their plant hosts. When female beetles leave the parental gallery to establish a new generation, they transport inocula of one or several mycosymbionts in specialized cavities in various parts of their bodies, to be grown in the subsequently developed galleries (Hubbard 1897, Beaver 1989).

The asexual morphological characteristics of *Raffaelea* are rather simple: hyaline, rarely-branching, commonly

single-celled conidiophores are arranged singly or aggregated in sporodochia; conidiogenous cells are precurently or sympodially proliferating, which may leave denticles, annellations, or inconspicuous scarring; conidia range from elliptical to globose, with some exceptions, and may reproduce by yeast-like budding (Harrington *et al.* 2010, Musvuugwa *et al.* 2015). De Beer & Wingfield (2013) recognized two sexually reproducing species of *Ophiostoma*, *O. seticolle* and *O. deltoideosporum*, in *Raffaelea* s. str. based on DNA sequence phylogenies, but they did not transfer these species to *Raffaelea*. Subsequently, Musvuugwa *et al.* (2015) described a *Raffaelea* species, *R. vaginata*, with an observed sexual morph, similar to those of *O. seticolle* and *O. deltoideosporum*. The latter authors consequently emended the circumscription of the genus to include both asexual and sexual morphs, and transferred the two *Ophiostoma* species to *Raffaelea* as *R. seticollis* and *R. deltoideospora*, consistent

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Table 1. Cultures of *Raffaelea* examined in our molecular phylogenetic analyses, with ex-type cultures of new species in bold.

Species	Isolate	CMW culture number (Other identifier)	Beetle host	Plant host	Country	Region (Locality)	28S rDNA	BTUB	18S rDNA	ITS DNA
<i>R. aguacate</i>	PL1004	CMW38067 (272)		<i>Persea americana</i>	USA	Florida (Miami-Dade Co.)	KJ909296	KJ909297	KF026302	
<i>R. campbellii</i>	103p2	CMW44800		<i>Persea americana</i>	USA	Florida (Miami-Dade Co., Homestead)	KR018414	KR018442	KR018402	
<i>R. campbellii</i>	110p2	CMW44801		<i>Persea americana</i>	USA	Florida (Miami-Dade Co., Homestead)	KR018430	KR018444	KR018403	
<i>R. cf. campbellii</i>	Hulcr7355		<i>Euplatypus compositus</i>		USA	Florida	KX267101	KX267112		
<i>R. cf. subalba</i>	Hulcr7375		<i>Euplatypus compositus</i>		USA	Florida	KX267102	KX267113		
<i>R. crosotarsa</i>	Hulcr7182	CMW44793	<i>Crossotarsus emancipatus</i>	<i>Lithocarpus</i> sp.	Taiwan	Fushan	KX267103	KX267114	KX267129	KX267135
<i>R. cyclorhipidia</i>	Hulcr7168	CMW44790	<i>Cyclorhipidion ohnoi</i>	<i>Lithocarpus</i> sp.	Taiwan	Fushan	KX267104	KX267115	KX267130	KX267136
<i>R. lauricola</i>	Hulcr4530	(PL1007)	<i>Ambrosiodimus lecontei</i>	<i>Persea borbonia</i>	USA	Florida (Lake Kissimmee)		KX267116		
<i>R. lauricola</i>	Hulcr7161		<i>Xyleborinus glabratus</i>		Taiwan		KX267105	KX267117		
<i>R. lauricola</i>	Hulcr7164		<i>Xyleborinus glabratus</i>		Taiwan		KX267106	KX267118		
<i>R. quercivora</i>	Hulcr7167		<i>Cyclorhipidion ohnoi</i>	<i>Lithocarpus</i> sp.	Taiwan	Fushan	KX267107	KX267119	KX267131	
<i>R. quercivora</i>	Hulcr7176		<i>Crossotarsus emancipatus</i>	<i>Lithocarpus</i> sp.	Taiwan	Fushan	KX267108	KX267120	KX267132	
<i>R. subfusca</i>	Hulcr4520	(C2335)	<i>Xyleborus glabratus</i>		USA	South Carolina (Hunting Island State Park)		KX267121		KX267137
<i>R. subfusca</i>	Hulcr4717		<i>Euwallacea validus</i>		USA	Virginia (Albemarle Co., Batesville)		KX267122	KX267133	KX267138
<i>R. subfusca</i>	Hulcr4719		<i>Euwallacea validus</i>		USA	Pennsylvania (Huntingdon Co. Raystown Lake)	KX267109	KX267123	KX267134	
<i>R. xyleborina</i>	Hulcr6099	CMW45859	<i>Xyleborinus andrewesii</i>		USA	Florida (Highlands Co., Venus)	KX267110	KX267124		
<i>R. xyleborina</i>	Hulcr6100		<i>Xyleborinus andrewesii</i>		USA	Florida (Highlands Co., Venus)	KX267111	KX267125		
<i>R. xyleborina</i>	Hulcr6406		<i>Xyleborinus andrewesii</i>		USA	Florida (Highlands Co., Venus)		KX267126		KX267139
<i>R. xyleborina</i>	Hulcr6408		<i>Xyleborinus andrewesii</i>		USA	Florida (Highlands Co., Venus)		KX267127		KX267140
<i>Raffaelea</i> sp.	Hulcr5951	(PL1635)	<i>Xyleborus pinicola</i>	<i>Pinus keysei</i>	Thailand	Mae Chaem	KJ909308	KJ909310	KJ909309	
<i>Raffaelea</i> sp.	Hulcr7507		<i>Xyleborinus gracilis</i>		USA	Florida		KX267128		KX267141
<i>Raffaelea</i> sp.	PL1001	CMW38062		<i>Persea americana</i>	USA	California	KJ909293	KJ909295	KJ909294	

with the one fungus-one name rule (Hawksworth 2011).

Some molecular phylogenies of *Raffaelea* and additional genera within *Ophiostomatales* have suggested that *Raffaelea* is monophyletic (e.g. Harrington *et al.* 2010). However, more recent and comprehensive analyses (de Beer & Wingfield 2013, Dreaden *et al.* 2014, Musvuugwa *et al.* 2015) have shown that *Raffaelea* species constitute three clades in the order, *Raffaelea s. str.*, the *R. lauricola* complex, and the *R. sulphurea* complex. Of ecological interest, the two clades exterior to *Raffaelea s. str.* each include plant pathogens that have been spread in the last decade by their respective insect vectors. *Raffaelea lauricola*, the causative agent of laurel wilt in the southeastern USA, is associated with the ambrosia beetle *Xyleborus glabratus* (Harrington *et al.* 2008, Ploetz *et al.* 2013), among others (Carrillo *et al.* 2014). *Raffaelea lauricola*, the eponymous member of the *R. lauricola* complex (de Beer & Wingfield 2013, Musvuugwa *et al.* 2015), is sometimes placed as sister to *Raffaelea s. str.* in molecular phylogenies of individual rDNA loci (Musvuugwa *et al.* 2015) and additional coding genes (Dreaden *et al.* 2014). *Raffaelea quercivora*, which is responsible for Japanese oak wilt and associated with *Platypus quercivorus* (Kubono & Ito 2002, Kusumoto *et al.* 2014), lies within the *R. sulphurea* complex in *Leptographium s. lat.* (de Beer & Wingfield 2013, Dreaden *et al.* 2014, Musvuugwa *et al.* 2015).

During domestic (Campbell *et al.* 2016) and international studies to investigate the diversity of ambrosia beetles and their fungal symbionts, *raffaelea*-like isolates from the southeastern USA and Taiwan were collected; preliminary molecular analyses indicated that some of these isolates represent novel lineages within *Raffaelea s. lat.* In this study, we use nine isolates to describe five new species in *Raffaelea* from collections of plant hosts and ambrosia beetles. We have also characterized 13 additional *Raffaelea* isolates based on DNA sequence data.

MATERIALS AND METHODS

DNA extraction, PCR amplification and sequencing

Twenty-two *Raffaelea* cultures and DNA extracts were aggregated from the Forest Entomology laboratory at the University of Florida (Gainesville, FL) and the University of Florida's Tropical Research and Education Center (Homestead, FL). Cultures from ambrosia beetle hosts were isolated by dilution plating of mycangial contents, as described by Li *et al.* (2015). Cultures of newly described species are deposited in the culture collection (CMW) of the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, South Africa (Table 1).

Fungal DNA was isolated with Extract-N-Amp PCR kits (Sigma-Aldrich), as described by Li *et al.* (2015). Final concentrations of PCR reagent solutions in 25 µL were: (1) 1× ClonTech-TaKaRa Ex Taq Buffer; (2) 5 % DMSO; (3) 0.2 mM each dNTP; (4) 0.5 µM each primer; (5) 0.625 U Ex Taq polymerase; and (6) 0.01–0.1 ng extracted DNA. Primer combinations used for amplifications were: (1) LR0R/LR5 (Vilgalys & Hester 1990, Rehner & Samuels 1994) for nuclear large subunit (28S) ribosomal DNA (rDNA); (2) T10 or Bt2a/

Bt2b (Glass & Donaldson 1995, O'Donnell & Cigelnik 1997) for β-tubulin (βT); (3) NS1/NS4 for nuclear small subunit (18S) rDNA; and (4) either ITS3/LR3 or ITS1F/ITS4 (White *et al.* 1990, Gardes & Bruns 1993) for portions of the ITS1–5.8S–ITS2 (ITS) rDNA locus. The PCR conditions for βT and ITS rDNA were the same as those used by Yin *et al.* (2015) and for 18S and 28S rDNA by Dreaden *et al.* (2014). Amplified products were visualized and purified as described by Li *et al.* (2015), and these were submitted to the University of Florida Interdisciplinary Center for Biotechnology Research for Sanger sequencing. Chromatograms were assembled and inspected with Geneious v. 9.0.5.

Phylogenetic analyses

Sequences of 28S rDNA and βT (introns 3/4/5 removed) were aligned and visually inspected in Geneious for phylogenetic reconstruction. The alignment was divided into four partitions for phylogenetic consideration: one partition for the 28S rDNA alignment and for each of the three codon positions in the protein encoding βT. The Akaike information criterion in jModeltest 0.1.1 (Guindon & Gascuel 2003, Posada 2008) was used to select the nucleotide substitution model for each partition. Maximum likelihood (ML) phylogenetic analyses were conducted in GARLI 2.01 (Zwickl 2006) with the recommended partition parameters to determine the best tree topology (Fig. 1) and bootstrap support values from 500 search replicates, which were summarized in SumTrees (Sukumaran & Holder 2010). Bayesian posterior probabilities (BPP) were estimated with the same partition parameters in an analysis conducted in MrBayes 3.1.2 (Ronquist & Huelsenbeck 2003), in which two runs of four chains each were executed simultaneously for 5 000 000 generations, with sampling every 500 generations. SumTrees was used to compute BPP from a summary of 7501 trees retained after a burn-in of the first 2500 trees collected.

Growth trials and morphological characterization

To determine optimal growth rates of each new species of *Raffaelea*, discs of agar (7 mm diam) covered with mycelium were aseptically removed from 1-wk-old cultures growing on BD Difco™ MEA and used to inoculate plates incubated at 10–35 °C, in 5 °C intervals. After 9 d, colony growth was calculated as by Musvuugwa *et al.* (2015). Morphological features were examined by inoculating sterile slide mounts of BD Difco™ MEA with propagules collected by running a sterile needle along the surface of cultures growing on BD Difco™ MEA. Once reproductive structures were observed using a dissecting microscope (24–48 h), slides were examined on an Olympus BX53 equipped with a Canon EOS Rebel T3i using EOS Utility 2 software. For each new species, measurements of conidiophores ($n=5$) and conidia ($n=10$) were made to the nearest 0.5 µm, and means ($\pm$ standard deviation) were calculated to the nearest 0.1 µm.

RESULTS

All isolates we examined resided in *Raffaelea s. str.*, the *R. lauricola* complex, or the *R. sulphurea* complex in the

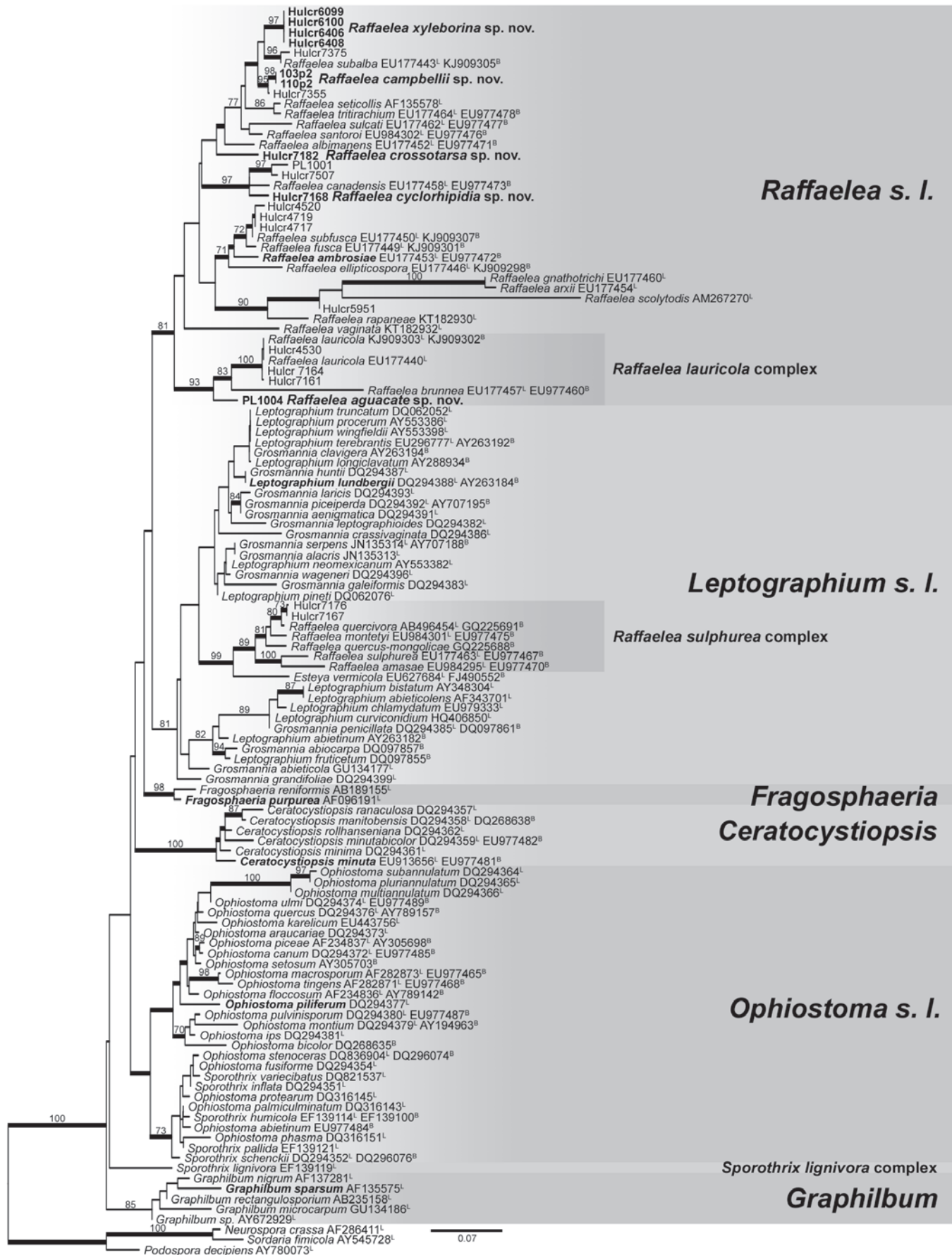


Fig. 1. Best ML tree from GARLI analysis of 28S rDNA and β T data matrix of *Ophiostomatales* genera with *Sordariales* as outgroup (Musvuugwa et al. 2015). Values at nodes represent ML bootstrap percentages ≥ 70 % from a summary of 500 replicates, and branches in bold represent BPP ≥ 95 %. ^L denotes GenBank accession number of 28S rDNA sequence for taxon; ^B denotes GenBank accession number of β T sequence for taxon. Types of genera and new species of *Raffaelea* in bold.

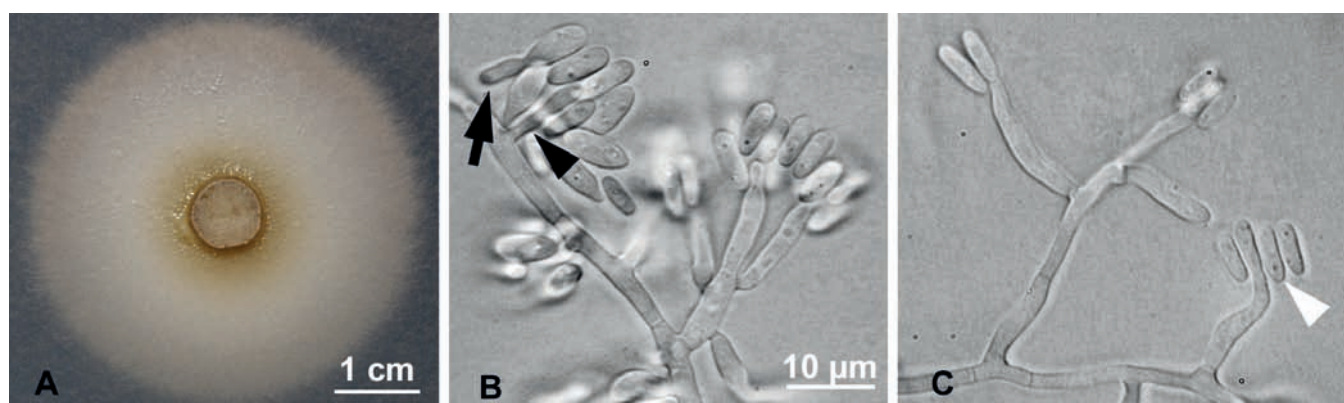


Fig. 2. *Raffaelea aguacate* (PL1004) morphological features in pure culture on MEA. **A.** Colony growth after 9 d at 25 °C. **B.** Hyphae bearing long, tapering conidiogenous cells with conidia at apex, and occasional sessile lateral conidia (black arrowhead); elongated conidia may bud yeast-like daughter cell (black arrow). **C.** Hyphae with long and slightly irregular conidiogenous cells, with conidia truncated at base (white arrowhead). Bar in B applies also to C.

phylogenetic analyses of the 28S rDNA and β T data matrices (Fig. 1). The *R. lauricola* complex was sister to the *Raffaelea* s. str. clade with 81 % ML bootstrap and 100 % BPP support, and the well-supported *R. sulphurea* clade resolved within *Leptographium* s. lat. The ITS and 18S rDNA sequences were not included in the phylogenetic analyses, but these sequences were used for molecular identification (Table 1). The data matrix for the 28S rDNA and β T regions has been deposited in TreeBASE as submission 19323.

The new species in *Raffaelea* s. str. and the *R. lauricola* complex (Table 1) possessed all β T introns (3/4/5). Isolates Hulcr7167 and Hulcr7176 possessed two introns (3/4/-). These patterns of intron presence were the expected conditions for the majority of species in each clade (de Beer & Wingfield 2013). Although they were not isolated from *Platypus quercivorus*, isolates Hulcr7167 and Hulcr7176 resolved in the *R. sulphurea* complex with *R. quercivora*, and were 99 % (396/400 bp) and 98 % (392/400 bp) similar, respectively, to the β T sequence (including introns) of *R. quercivora*. The 28S rDNA sequences for isolates Hulcr7167 and Hulcr7176 were 98 % (492–493/499 bp) similar to *R. quercivora*, but the representative *R. quercivora* sequence (GenBank accession AB496454) had six ambiguous bases that increased the level of dissimilarity with our isolates.

TAXONOMY

Raffaelea aguacate D.R. Simmons, Dreaden & Ploetz, sp. nov.

MycoBank MB817170
(Fig. 2)

Etymology: The epithet “*aguacate*” refers to the Spanish for avocado (*Persea americana*), from which this isolate was cultured.

Diagnosis: Conidiogenous cells $13 (\pm 2) \times 2.7 (\pm 0.3) \mu\text{m}$, hyaline, sometimes irregular; conidia at conidiogenous cell apex or sessile and lateral; conidia $7.2 (\pm 0.6) \times 2.6 (\pm 0.5) \mu\text{m}$, elongate, truncated at base, hyaline, rarely with yeast-like budding.

Type: **USA:** Florida: Miami-Dade Co., Homestead, from bioassay of *Persea americana*, 2009, C. L. Harmon (BPI 910154 – holotype; 272 = PL1004 = CMW38067 – ex-type cultures).

Description: Colonies initially cream, turning light green to olivaceous, aging to dark green on MEA; reverse subhyaline. Optimal colony diameter after 9 d at 25 °C in the dark was $70.2 (\pm 3.9) \text{ mm}$; $46.0 (\pm 2.6) \text{ mm}$ at 10 °C; no growth at 35 °C. Conidiogenous cells hyaline, sometimes irregular, tapering at ends, $13 (\pm 2) \times 2.7 (\pm 0.3) \mu\text{m}$. Conidia forming from apex of conidiogenous cells, hyaline, occasionally sessile and lateral. Conidia produced singly, aseptate, elongate, and occasionally truncated at the base, $7.2 (\pm 0.6) \times 2.6 (\pm 0.5) \mu\text{m}$. Conidia rarely budding. Sexual morph unknown.

Raffaelea campbellii D.R. Simmons, A. Campbell & Ploetz, sp. nov.

MycoBank MB817171
(Fig. 3)

Etymology: The epithet “*campbellii*” is in honor of Donald and Princessa Campbell, parents of Alina S. Campbell, collector of the specimen, for their guidance and support.

Diagnosis: Conidiogenous cells $13.7 (\pm 1.6) \times 3.7 (\pm 0.3) \mu\text{m}$, hyaline, flask-shaped; conidia at conidiogenous cell apex; conidia $6.7 (\pm 1.2) \times 3.6 (\pm 0.5) \mu\text{m}$, ovoid to elliptical, truncated at base, hyaline.

Type: **USA:** Florida: Miami-Dade Co., cultured from *Xyleborus glabratus* that infected *Persea palustris*, Jun. 2013, A. S. Campbell (BPI 910156 – holotype; 103p2 = CMW44800 – ex-type culture).

Additional specimen examined: Loc. cit (110p2 = CMW44801).

Description: Colonies initially cream, turning olivaceous to blackish on MEA, surface tough and wrinkled; reverse subhyaline. Optimal colony diameter after 9 d at 25 °C in the dark $25.7 (\pm 1.3) \text{ mm}$; no growth at 10 °C or 35 °C.

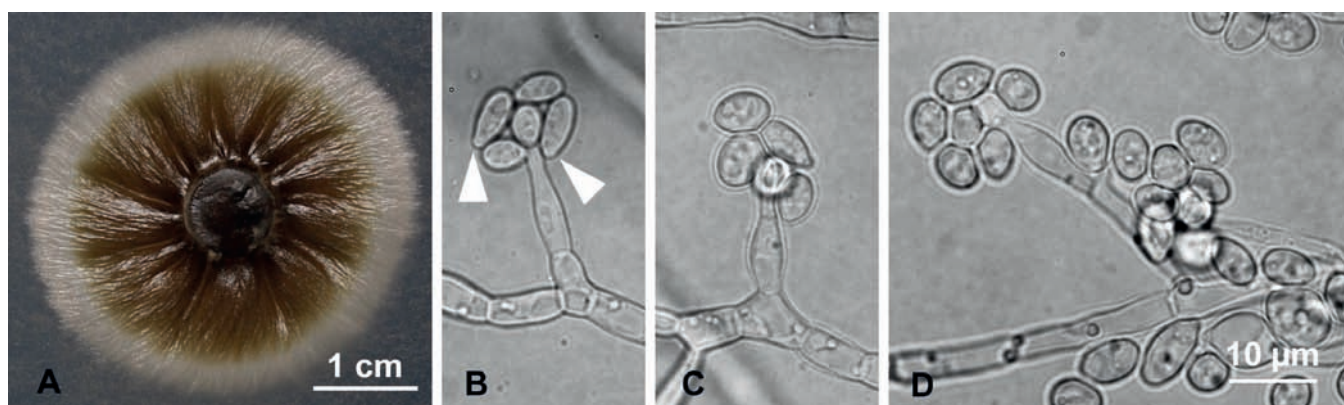


Fig. 3. *Raffaelea campbellii* (103p2) morphological features in pure culture on MEA. **A.** Colony growth after 9 d at 25 °C. **B–D.** Hyphae bearing flask-shaped conidiogenous cells with ovoid to elliptical conidia, often truncated at the base (white arrowheads). Bar in D applies also to B–C.

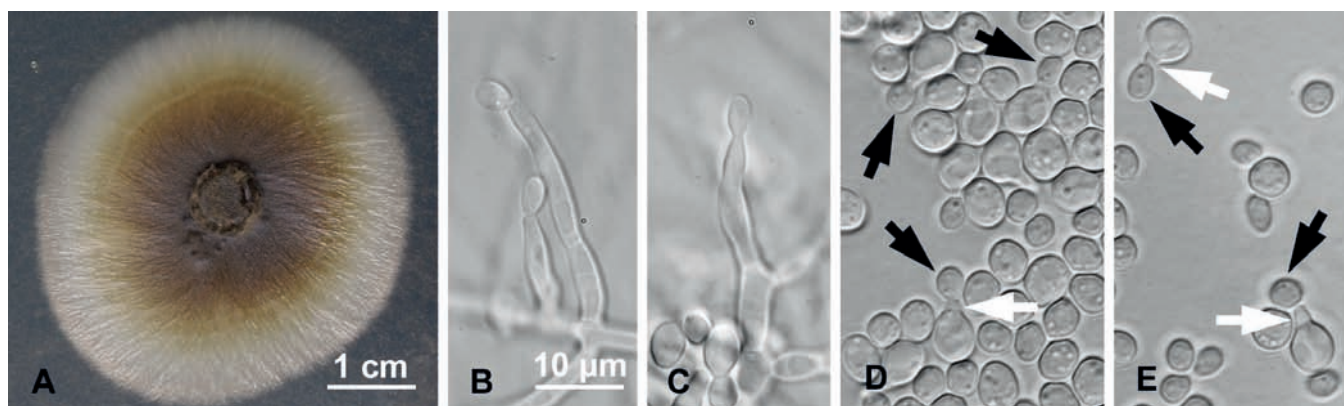


Fig. 4. *Raffaelea crossotarsa* (Hulcr7182) morphological features in pure culture on MEA. **A.** Colony growth after 9 d at 25 °C. **B–C.** Hyphae bearing long, tapering conidiogenous cells with conidia. **D–E.** Globose to ovoid conidia budding yeast-like daughter cells (black arrows), which protrude from prominent isthmuses (white arrows). Bar in B applies also to C–E.

Conidiogenous cells hyaline, flask-shaped, tapering towards the apex, $13.7 (\pm 1.6) \times 3.7 (\pm 0.3) \mu\text{m}$. *Conidia* forming from apex of conidiogenous cells, hyaline. Conidia produced singly, accumulating at tip of conidiogenous cells, aseptate, ovoid to elliptical, sometimes fusiform, and often truncate at the base, $6.7 (\pm 1.2) \times 3.6 (\pm 0.5) \mu\text{m}$. *Sexual morph* unknown.

***Raffaelea crossotarsa* D.R. Simmons & Y.T. Huang, sp. nov.**

MycoBank MB817172
(Fig. 4)

Etymology: The epithet “*crossotarsa*” refers to the genus of the host beetle (*Crossotarsus emancipatus*), the mycangium of which yielded this fungus.

Diagnosis: Conidiogenous cells $15.2 (\pm 2.1) \times 3 (\pm 0.3) \mu\text{m}$, hyaline, slender; conidia at conidiogenous cell apex; conidia $6 (\pm 0.4) \times 4.9 (\pm 0.3) \mu\text{m}$, globose to ovoid, hyaline, yeast-like budding from prominent isthmus.

Type: **Taiwan:** Fushan, cultured from *Crossotarsus emancipatus* collected from *Lithocarpus* sp., Mar. 2015, J. Hulcr, A. Black & D. R. Simmons (BPI 910157 – holotype; Hulcr7182 = CMW44793 – ex-type culture).

Description: Colonies initially cream, aging from golden olivaceous to dark green or dark ruddy brown on MEA, surface tough; reverse subhyaline. Optimal colony diameter after 9 d at 25 °C in the dark was $39.2 (\pm 1.2) \text{ mm}$; $9.0 (\pm 0.5) \text{ mm}$ at 10 °C; no growth at 35 °C. *Conidiogenous cells* hyaline, slender, tapering at ends, $15.2 (\pm 2.1) \times 3 (\pm 0.3) \mu\text{m}$. *Conidia* forming from apex of conidiogenous cells, hyaline. Conidia produced singly, aseptate, globose to ovoid, $6 (\pm 0.4) \times 4.9 (\pm 0.3) \mu\text{m}$. Conidia producing budding cells from prominent isthmus, 1–2 μm long. *Sexual morph* unknown.

***Raffaelea cyclorhipidia* D.R. Simmons & Y.T. Huang, sp. nov.**

MycoBank MB817173
(Fig. 5)

Etymology: The epithet “*cyclorhipidia*” refers to the genus of the host beetle (*Cyclorhipidion ohnoi*), the mycangium of which yielded this fungus.

Diagnosis: Conidiogenous cells $12 (\pm 1.7) \times 3.6 (\pm 0.3) \mu\text{m}$, hyaline, flask-shaped; conidia at conidiogenous cell apex or sessile and lateral; conidia $7.3 (\pm 1.0) \times 3.5 (\pm 0.7) \mu\text{m}$, elliptical to elongate, hyaline, yeast-like budding.

Type: **Taiwan:** Fushan, cultured from *Cyclorhipidion ohnoi*

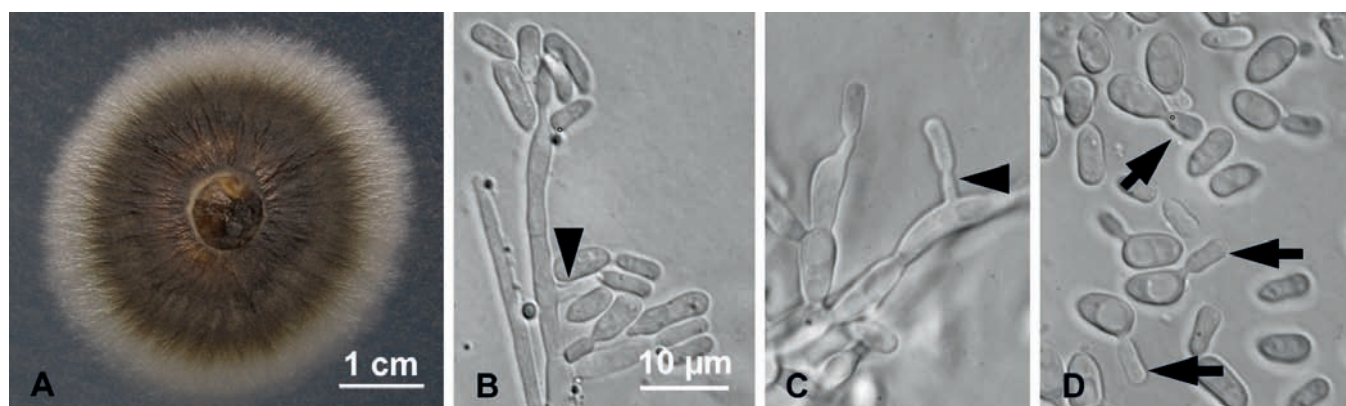


Fig. 5. *Raffaelea cyclorhipidia* (Hulcr7168) morphological features in pure culture on MEA. **A.** Colony growth after 9 d at 25 °C. **B–C.** Hyphae bearing typical flask-shaped conidiogenous cells with conidia at apex, and occasional lateral sessile conidia (black arrowheads). **D.** Elliptical to elongate conidia budding yeast-like daughter cells (black arrows). Bar in B applies also to C–D.

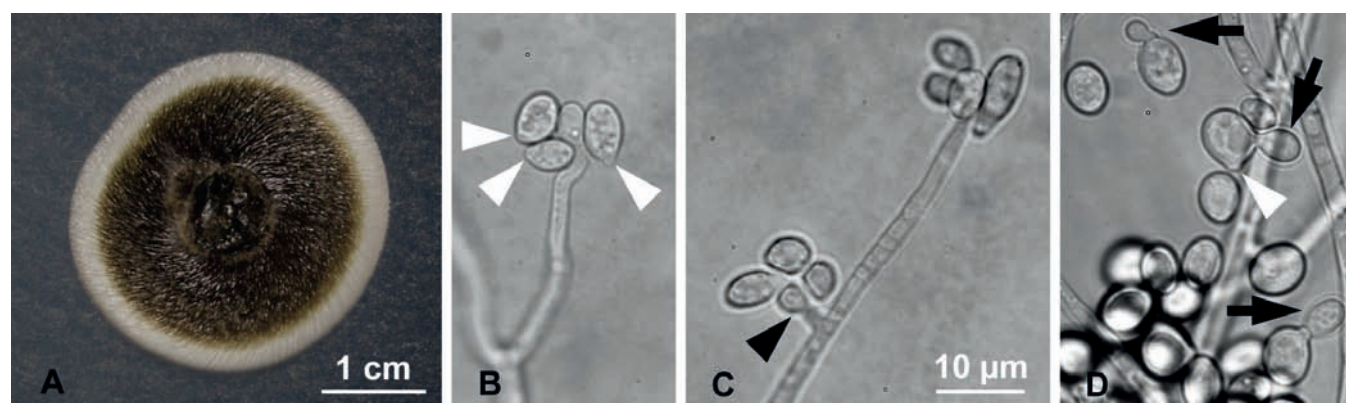


Fig. 6. *Raffaelea xyleborina* (Hulcr6099) morphological features in pure culture on MEA. **A.** Colony growth after 9 d at 25 °C. **B.** Micronematous conidiogenous cells with ovoid conidia truncated at base (white arrowhead). **C.** Micronematous conidiophore with short conidiogenous cell sessile at side (black arrowhead) and at apex. **D.** Globose to ovoid conidia truncated at the base (white arrowhead) and budding yeast-like daughter cells (black arrows). Bar in C applies also to B–D.

collected infesting *Lithocarpus* sp., Mar. 2015, J. Hulcr, A. Black & D. R. Simmons (BPI 910158 – holotype; Hulcr7168 = CMW44790 – ex-type culture).

Description: Colonies initially cream, aging from olivaceous to golden brown or blackish on MEA, surface tough and wrinkled; reverse subhyaline. Optimal colony diameter after 9 d at 25 °C in the dark was 47.5 (±1.9) mm; 20.6 (±1.8) mm at 10 °C; no growth at 35 °C. *Conidiogenous cells* hyaline, flask-shaped, tapering towards the apex, 12 (±1.7) × 3.6 (±0.3) µm. *Conidia* forming at apex of conidiogenous cells, occasionally sessile and lateral, hyaline. Conidia produced singly, aseptate, elliptical to elongate, occasionally truncate at base, 7.3 (±1.0) × 3.5 (±0.7) µm. Conidia produce budding cells. *Sexual morph* unknown.

Raffaelea xyleborina D.R. Simmons & C. Bateman, sp. nov.
MycoBank MB817174
(Fig. 6)

Etymology: The epithet “*xyleborina*” refers to the genus of the host beetle (*Xyleborinus andrewesii*), the mycangium of which yielded this fungus.

Diagnosis: Conidiophores micronematous, hyaline; conidia at conidiogenous cell apex or lateral and sessile; conidia 6.5 (±0.7) × 4.9 (±0.8) µm, globose to ovoid, truncated at base, hyaline, yeast-like budding.

Type: **USA:** Florida: Highlands Co., Venus, cultured from *Xyleborinus andrewesii* collected from bait trap, 3 Jan. 2013, C. Bateman, C. Gibbard & L. L. Stelinski (BPI 910159 – holotype; Hulcr6099 = CMW45859 – ex-type culture;

Additional specimens examined: Loc. cit (Hulcr6100, Hulcr6406, Hulcr6408).

Description: Colonies initially cream, varying with age from cream to dark green to blackish on MEA, surface tough and spiral in appearance; reverse subhyaline. Optimal colony diameter after 9 d at 35 °C in the dark was 26.8 (±3.0) mm; 14.9 (±1.3) mm at 25 °C; no growth at 10 °C. *Conidiogenous cells* hyaline, micronematous, with conidia forming at apex, occasionally sessile and lateral. *Conidia* produced singly, aseptate, hyaline, globose to ovoid, sometimes elongate, and often truncated at base, 6.5 (±0.7) × 4.9 (±0.8) µm. Conidia produce budding cells. *Sexual morph* unknown.

DISCUSSION

Considering the damage that ambrosia fungi and their vectors cause (Ploetz et al. 2013), there is an urgent need to determine not only the diversity of these fungi globally but also to gain an enhanced knowledge of the host vector range for these potentially devastating species. Comparison of fungal isolates in this study with known species of *Raffaelea* revealed that two isolates from Taiwan, Hulcr7167 and Hulcr7176, grouped with *R. quercivora*. *Raffaelea quercivora* has been isolated from *Platypus quercivorus* in Japan, where it is responsible for ongoing epidemics of Japanese oak wilt (Kubono & Ito 2002), as well as in Taiwan (Kusumoto et al. 2014). Our isolates of *R. quercivora* were not isolated from the mycangia or fungal galleries of *P. quercivorus*, however, but rather from the mycangia of *Cyclorhipidion ohnoi* and the fungal galleries of *Crossotarsus emancipatus* from Taiwan. Though the latter two beetle species have not been implicated in oak wilt, these symbioses suggest that other vectors of *R. quercivora* exist. These isolates were collected from the same beetle host populations in Taiwan from which two of the species newly described in this study, *R. cyclorhipidia* and *R. crossotarsa*, were recovered. Therefore, these beetle-associated species display a degree of promiscuity with fungi within and exterior to *Raffaelea s. str.*

Raffaelea lauricola, the causative agent of laurel wilt, was found in Taiwan, from the documented host *Xyleborus glabratus*, and in Florida, from the previously unrecorded host *Ambrosiodmus lecontei*. Carrillo et al. (2014) reported that *R. lauricola* was laterally transferred to additional ambrosia beetle hosts, other than *X. glabratus*, when these species co-inhabit trees infected with this fungal pathogen. This finding demonstrates that the pathogen is a relatively promiscuous symbiont of ambrosia beetles, raising its importance from the biosecurity perspective. Despite Carrillo et al. (2014) having examined 41 adult *A. lecontei* females emerging from laurel wilt-affected swamp bay bolts, they failed to isolate *R. lauricola* from this host species. However, we recovered *R. lauricola* from *A. lecontei* infesting *Persea borbonia* near Lake Kissimmee (FL). The presence of *Raffaelea* with *Ambrosiodmus* may be phoretic or facultative, because *Ambrosiodmus* species examined to date carry a highly specific ambrosial basidiomycetous species (Li et al. 2015).

Besides information on known ambrosia fungi, results of this study suggest that under-explored regions of the world contain a large diversity of undescribed ambrosia fungi. Phylogenetic analyses of DNA sequence data for 22 isolates of *Raffaelea*-like fungi led to the discovery of the five new species described here. Some additional isolates resolved in lineages that would generally support their description as novel taxa (i.e. Hulcr5951; Hulcr7355; Hulcr7507 and PL1001), but these cultures could not be revived for morphological characterization after cryopreservation. Four of the new species described in this study were isolated from mycangia of ambrosia beetle hosts. Although sampling efforts that provided the foundation for this study included many different parts of the world, three of the novel taxa were from the eastern US. Whether this is a true reflection of an unexamined area of *Raffaelea* species diversity, or due to sampling bias, is unknown but deserves future consideration.

Results from this study indicate that *Raffaelea s. str.* and the *R. lauricola* complex are monophyletic (Fig. 1; *Raffaelea s. lat.*). This is consistent with previous analyses using rDNA and β T sequences (Dreaden et al. 2014). Analyses of 28S rDNA across *Ophiostomatales* have shown the same association with some support (Musvuugwa et al. 2015) or that these clades are disparate (de Beer & Wingfield 2013). Until a more accurate determination of their relationship is conducted with additional genetic loci, we conclude that these two clades are distinct.

Fungal symbioses with ambrosia beetles have become especially fertile topics for research, and further study will likely identify an increasingly large diversity of fungal associates. Indeed, Bateman et al. (2016) described a new genus in *Ophiostomatales* from *Premnobius cavipennis* (Scolytinae; Ipini), an independently evolved ambrosia beetle lineage largely confined to Africa. Furthermore, ambrosia beetles' mycosymbionts are not limited to the ascomycetous *Ophiostomatales*. Li et al. (2015) found a new basidiomycetous *Polyporales* fungus, *Flavodon ambrosius* (Simmons et al. 2016), in symbiosis with *Ambrosiodmus* species, and Kasson et al. (2016) found the same mycosymbiont associated with another genus, *Ambrosiophilus*, which is sister to *Ambrosiodmus* (Hulcr & Cognato 2010). Thus, as investigations into these insects increase in number, additional fungal genera in unexpected lineages may be found in symbioses with ambrosia beetles.

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REFERENCES

- Arx JA von, Hennebert GL (1965) Deux champignons ambrosia. *Mycopathologia et Mycologia Applicata* **25**: 309–315.
- Bateman CC, Huang Y-T, Simmons DR, Kasson MT, Stanley EL, Hulcr J (2016) Ambrosia beetle *Premnobius cavipennis* (Scolytinae: Ipini) carries highly divergent ascomycotan ambrosia fungus *Afroraffaelea ambrosiae* gen. nov. sp. nov. (*Ophiostomatales*). *Fungal Ecology*: in press.
- Beaver RA (1989) Insect-fungus relationship in the bark and ambrosia beetles. In: *Insect-fungus Interactions* (Wilding N, Collins NM, Hammond PM, Webber JF, eds): 121–143. San Diego: Elsevier.
- Campbell AS, Ploetz RC, Dreaden TJ, Kendra PE, Montgomery WS (2016) Geographic variation in mycangial communities of *Xyleborus glabratus*. *Mycologia* **108**: 657–667.

- Carrillo D, Duncan RE, Ploetz JN, Campbell AF, Ploetz RC, Peña JE (2014) Lateral transfer of a phytopathogenic symbiont among native and exotic ambrosia beetles. *Plant Pathology* **63**: 54–62.
- de Beer ZW, Seifert KA, Wingfield MJ (2013) A nomenclator for ophiostomatoid genera and species in the *Ophiostomatales* and *Microascales*. In: *Ophiostomatoid Fungi: expanding frontiers* (Seifert KA, de Beer ZW, Wingfield MJ, eds): 245–322. [CBS Biodiversity Series 12.] Utrecht: CBS-KNAW Fungal Biodiversity Centre.
- de Beer ZW, Wingfield MJ (2013) Emerging lineages in the *Ophiostomatales*. In: *Ophiostomatoid Fungi: expanding frontiers* (Seifert KA, de Beer ZW, Wingfield MJ, eds): 21–46. [CBS Biodiversity Series 12.] Utrecht: CBS-KNAW Fungal Biodiversity Centre.
- Dreaden TJ, Davis JM, de Beer ZW, Ploetz RC, Soltis PS, *et al.* (2014) Phylogeny of ambrosia beetle symbionts in the genus *Raffaelea*. *Fungal Biology* **118**: 970–978.
- Gardes M, Bruns TD (1993) ITS primers with enhanced specificity for basidiomycetes: application to the identification of mycorrhizae and rusts. *Molecular Ecology* **2**: 113–118.
- Glass NL, Donaldson GC (1995) Development of primer sets designed for use with the PCR to amplify conserved genes from filamentous ascomycetes. *Applied and Environmental Microbiology* **61**: 1323–1330.
- Guindon S, Gascuel O (2003) A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. *Systematic Biology* **52**: 696–704.
- Harrington TC, Aghayeva DN, Fraedrich SW (2010) New combinations in *Raffaelea*, *Ambrosiella*, and *Hyalorhinocladia*, and four new species from the Redbay ambrosia beetle, *Xyleborus glabratus*. *Mycotaxon* **111**: 337–361.
- Harrington TC, Fraedrich SW, Aghayeva DN (2008) *Raffaelea lauricola*, a new ambrosia beetle symbiont and pathogen on the *Lauraceae*. *Mycotaxon* **104**: 399–404.
- Hawksworth DL (2011) A new dawn for the naming of fungi: impacts of decisions made in Melbourne in July 2011 on the future publication and regulation of fungal names. *MycKeys* **1**: 7–20.
- Hubbard HG (1897) The ambrosia beetles of the United States. *USDA, Division of Entomology, Bulletin, n.s.* **7**: 9–30.
- Hulcr J, Cognato AI (2010) Repeated evolution of crop theft in fungus-farming ambrosia beetles. *Evolution* **64**: 3205–3212.
- Kasson MT, Wickert KL, Stauder CM, Macias AM, Berger MC, *et al.* (2016) Mutualism with aggressive wood-degrading *Flavodon ambrosius* (*Polyporales*) facilitates niche expansion and sub-sociality in *Ambrosiophilus* ambrosia beetles. *Fungal Ecology* **23**: 86–96.
- Kubono T, Ito S (2002) *Raffaelea quercivora* sp. nov. associated with mass mortality of Japanese oak, and the ambrosia beetle (*Platypus quercivorus*). *Mycoscience* **43**: 255–260.
- Kusumoto D, Masuya H, Hirao T, Goto H, Hamaguchi K, *et al.* (2014) Discoloration induced by *Raffaelea quercivora* isolates in *Quercus serrata* logs and its relation to phylogeny: a comparison among isolates with and without the Japanese oak wilt incidence including outside of Japan. *Journal of Forest Research* **19**: 404–410.
- Li Y, Simmons DR, Bateman CC, Short DPG, Kasson MT, Rabaglia RJ, Hulcr J (2015) New fungus-insect symbiosis: culturing, molecular, and histological methods determine saprophytic *Polyporales* mutualists of *Ambrosiodmus* ambrosia beetles. *PLoS ONE* **10**: e0137689.
- Musvuugwa T, de Beer ZW, Duong TA, Dreyer LL, Oberlander KC, Roets F (2015) New species of *Ophiostomatales* from *Scolytinae* and *Platypodinae* beetles in the Cape Floristic Region, including the discovery of the sexual state of *Raffaelea*. *Antonie Van Leeuwenhoek* **108**: 933–950.
- O'Donnell KL, Cigelnik E (1997) Two divergent intragenomic rDNA ITS2 types within a monophyletic lineage of the fungus *Fusarium* are nonorthologous. *Molecular Phylogenetics and Evolution* **7**: 103–116.
- Ploetz RC, Hulcr J, Wingfield MJ, de Beer ZW (2013) Destructive tree diseases associated with ambrosia and bark beetles: black swan events in tree pathology? *Plant Disease* **97**: 856–872.
- Posada D (2008) jModeltest: phylogenetic model averaging. *Molecular Biology and Evolution* **25**: 1253–1256.
- Rehner SA, Samuels GJ (1994) Taxonomy and phylogeny of *GlIOClaidium* analysed from nuclear large subunit ribosomal DNA sequences. *Mycological Research* **98**: 625–634.
- Ronquist F, Huelsenbeck JP (2003) MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* **19**: 1572–1574.
- Simmons DR, Li Y, Bateman CC, Hulcr J (2016) *Flavodon ambrosius* sp. nov., a basidiomycetous mycosymbiont of *Ambrosiodmus* ambrosia beetles. *Mycotaxon* **131**: 277–285.
- Sukumaran J, Holder MT (2010) DendroPy: a python library for phylogenetic computing. *Bioinformatics* **26**: 1569–1571.
- Vilgalys R, Hester M (1990) Rapid genetic identification and mapping of enzymatically amplified ribosomal DNA from several *Cryptococcus* species. *Journal of Bacteriology* **172**: 4238–4246.
- White TJ, Bruns TD, Lee SB, Taylor JW (1990) Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In: *PCR Protocols: a guide to methods and applications* (Innis MA, Gelfand DH, Sninsky JJ, White TJ, eds): 315–322. San Diego: Academic Press.
- Yin M, Duong TA, Wingfield MJ, Zhou X, de Beer ZW (2015) Taxonomy and phylogeny of the *Leptographium procerum* complex, including *Leptographium sinense* sp. nov. and *Leptographium longiconidiophorum* sp. nov. *Antonie Van Leeuwenhoek* **107**: 547–563.
- Zwickl DJ (2006) *Genetic algorithm approaches for the phylogenetic analysis of large biological sequence datasets under the maximum likelihood criterion*. PhD dissertation, University of Texas at Austin.