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PCR Multiplexes Discriminate Fusarium Symbionts of Invasive Euwallacea Ambrosia Beetles that Inflict Damage on Numerous Tree Species Throughout the United States

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Abstract

Asian Euwallacea ambrosia beetles vector Fusarium mutualists. The ambrosial fusaria are all members of the ambrosia *Fusarium* clade (AFC) within the Fusarium solani species complex (FSSC). Several Euwallacea-Fusarium mutualists have been introduced into nonnative regions and have caused varying degrees of damage to orchard, landscape, and forest trees. Knowledge of symbiont fidelity is limited by current identification methods, which typically requires analysis of DNA sequence data from beetles and the symbionts cultured from their oral mycangia. Here, polymerase chain reaction (PCR)-based diagnostic tools were developed to identify the six Fusarium symbionts of exotic Euwallacea spp. currently known within the United States. Whole-genome sequences were generated for representatives of six AFC species plus F. ambrosium and aligned to the annotated genome of F. euwallaceae. Taxon-specific primer-annealing sites were identified that rapidly distinguish the AFC species currently within the United States. PCR specificity, reliability, and sensitivity were validated using a panel of 72 Fusarium isolates, including 47 reference cultures. Culture-independent multiplex assays accurately identified two AFC fusaria using DNA isolated from heads of their respective beetle partners. The PCR assays were used to show that Euwallacea validus is exclusively associated with AF-4 throughout its sampled range within eastern North America. The rapid assay supports federal and state agency efforts to monitor spread of these invasive pests and mitigate further introductions.

Several destructive fungal plant pathogens vectored by exotic bark and ambrosia beetles (Coleoptera: Curculionidae: Scolytinae) have invaded American landscapes and forests within the past century. For example, the redbay ambrosia beetle Xyleborus glabratus (Eichhoff) and its fungal symbiont Raffaelea lauricola T. C. Harr., Fraedrich & Aghayeva are responsible for the death of hundreds of millions of native Lauraceous plants, including avocado (Persea americana), redbay (P. borbonia), and sassafras (Sassafras albidum) (Fraedrich et al. 2008; Ploetz et al. 2013) throughout the coastal southeastern United States. Another group of scolytine beetles, the exotic Euwallacea ambrosia beetles, pose a threat to landscape trees as well as avocado (Eskalen et al. 2013; Kasson et al. 2013; Mendel et al. 2012).

At least six Euwallacea spp. from Asia have become established within the United States (Cognato et al. 2015; O'Donnell et al.

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2015): Euwallacea interjectus (Blandford), E. validus (Eichhoff), E. denticulus (Motschulsky), and three morphologically cryptic species within the E. fornicatus species complex (Eichhoff) (Atkinson 2016; O'Donnell et al. 2015; Storer et al. 2015). Euwallacea spp. are fungus-farming ambrosia beetles that cultivate mutualistic fungi in the genus Fusarium, although this is presently unconfirmed for E. denticulus. Most of these insect species do not cause noticeable economic or ecological damage and, instead, attack and colonize declining and recently killed trees. However, some of them are able to colonize living trees, sometimes in massive numbers, and the joint action of the wood borer and their cultivated fungi can cause symptoms known as Fusarium dieback or Fusarium canker. The greatest impact has been reported from the polyphagous shot hole borer on avocado and boxelder in California and Israel, which farms Fusarium euwallaceae S. Freeman, Z. Mendel, T. Aoki & O'Donnell (Eskalen et al. 2013; Freeman et al. 2013). However, similar testing is needed to confirm pathogenicity and host range of Fusarium stains associated with other introduced Euwallacea spp. in the United States.

The ambrosial fusaria belong to the F. solani species complex (FSSC), a clade that includes a number of important canker pathogens of deciduous trees (Park and Juzwik 2012; Tisserat 1987). Molecular systematics has revealed that the FSSC itself comprises three main clades (clades 1 to 3) encompassing over 60 species (O'Donnell et al. 2008; Short et al. 2013; Zhang et al. 2006) and recognizable by genealogical concordance phylogenetic species recognition (Taylor et al. 2000). The ambrosia Fusarium clade (AFC) is a newly discovered lineage within clade 3 of the FSSC. In total, 12 putatively clonal species-level lineages of AFC symbionts have been identified (O'Donnell et al. 2015), 10 of which lack Latin binomials. Therefore, an ad hoc nomenclature (i.e., AF followed by 1 to 12) was developed to distinguish the 12 fusaria within this clade. Most members of the AFC are morphologically indistinguishable (Kasson et al. 2013) and produce clavate macroconidia rather than the iconic fusiform conidia characteristic of *Fusarium* spp. Evolution of clavate macroconidia within this clade has been posited as an adaptation for the symbiosis (Gadd and Loos 1947; Kasson et al. 2013). Some *Euwallacea–Fusarium* associations appear to be highly specific but parsimony-based cophylogenetic analyses suggest that there have been multiple symbiont shifts over the evolutionary history of this mutualism (O'Donnell et al. 2015).

Despite the genealogical exclusivity of *Fusarium* symbionts and the successful delimitation of several AFC species using arbitrary-primer polymerase chain reaction (PCR) (Freeman et al. 2013), rapid detection and discrimination of *Fusarium* symbionts remains challenging for several reasons: (i) at least one *Euwallacea* sp. present within the United States and one in Sri Lanka farm two closely related ambrosia fusaria (O'Donnell et al. 2015); (ii) preliminary evidence suggests that interspecific hybridization may occur between members of the AFC and possibly with FSSC members outside the AFC, which presents challenges in resolving species boundaries; (iii) identification of species within the FSSC generally necessitates DNA sequence analysis of phylogenetically informative loci; and (iv) the current reliance on culturing and isolation of the symbionts requires living beetles, which precludes analysis of ethanol-preserved specimens.

Surveillance programs focused on the detection of beetles (genus *Euwallacea*) and disease (Fusarium dieback) have been initiated in several states and on the federal level within the United States Department of Agriculture program Cooperative Agricultural Pest Surveys. However, given the morphological uniformity of many species within the AFC (Kasson et al. 2013; O'Donnell et al. 2015), reliable molecular tools to discriminate exotic *Fusarium* symbionts are needed in order to advance our understanding of their ecology, distribution, and global invasion dynamics.

The present study was initiated to (i) develop and validate diagnostically robust multiplex assays to discriminate *Fusarium* symbionts of exotic *Euwallacea* ambrosia beetles currently present within the United States and (ii) validate multiplex assays that accurately identify AFC fusaria from DNA extracts of freshly macerated heads from living beetles. To bypass the need for fungal culturing, which often requires freshly collected live or recently killed beetles, we tested whether the multiplex PCR assays could be used to amplify DNA extracted directly from beetle heads. Amplification of fusarial DNA directly from beetle heads would facilitate research and detection of these fungi through a rapid culture-independent assessment and through simultaneous detection of different *Fusarium* spp. co-occurring in the same mycangium.

Materials and Methods

AFC comparative genomics. All of the AFC fusaria were originally isolated from live ambrosia beetles or their active galleries (Supplementary Table S1). Cultures were maintained on potato dextrose agar (Beckton, Dickinson and Company, Franklin Lakes, NJ) and transferred to potato dextrose broth to obtain mycelium for DNA extraction. A hexadecyltrimethyl-ammonium bromide (Sigma-Aldrich, St. Louis) protocol was used to obtain total genomic DNA for Sanger sequencing on an ABI3730 and for the multiplex assays (O'Donnell et al. 1998). Total genomic DNA for whole-genome sequencing was obtained using a ZR Fungal/bacterial DNA MiniPrep kit (Zymo Research, Irvine, CA). Whole-genome sequences of the following seven AFC species were generated with the Illumina MiSeq platform using libraries prepared with the NexteraXT kit following the manufacturer's instructions (Illumina, San Diego, CA): NRRL 20438 F. ambrosium (AF-1) from tea (Camelia chinensis) in India; NRRL 62626 F. euwallaceae (AF-2) from avocado in Los Angeles, CA; NRRL 62606 (AF-3) Fusarium sp. from boxelder (Acer negundo) in Gainesville, FL; NRRL 62579 Fusarium sp. (AF-4) from Ailanthus altissima in Pennsylvania; NRRL 62590 Fusarium sp. (AF-6) from avocado in Miami-Dade County, FL; NRRL 62584 Fusarium sp. (AF-8) from avocado in Miami-Dade County, FL; and KOD 792 Fusarium sp. (AF-12) from California sycamore (*Platanus racemosa*) in San Diego County, CA.

Sequence reads of the AFC genomes were trimmed and assembled with applications in CLC Genomics Workbench (Qiagen Bioinformatics, Redwood City, CA). The reads were assembled into 3,362 contigs. The assembly L50 was 68.1 kb, with the longest contig 312 kb in length. Genome annotation was performed using MAKER (Campbell et al. 2014), which takes in ab initio predictions from SNAP (Korf 2004), Augustus (Stanke et al. 2006), and GeneMark-ES (Borodovsky and Lomsadze 2011; Ter-Hovhannisyan et al. 2008). SNAP and Augustus ab initio parameters were initially run with F. graminearum parameters (https://github.com/hyphaltip/ fungi-gene-prediction-params) followed by parameter training on high-quality predictions from MAKER determined by transcript and protein support (AED better than 0.1). Repetitive elements were first identified by de novo repeat element discovery by RepeatModeler Open-1.0.7 (http://www.repeatmasker.org/RepeatModeler.html, Institute for Systems Biology, Seattle) followed by genome masking with RepeatMasker Open 3.0 (http://www.repeatmasker.org/faq. html, Institute for Systems Biology, Seattle, WA). RNAseq data were used in the genome annotation (Elmore et al. 2015), which was first assembled into transcripts with Trinity (Grabherr et al. 2011; Haas et al. 2013) using the Genome-Guided mode. Transcripts were aligned to the genome with BLASTN followed by exonerate polishing of alignments to obtain accurate intron/exon splice sites (Slater and Birney 2005) within MAKER. Proteins from annotated Sordariomycete genomes from the Swissprot database were clustered with cd-hit (Fu et al.2012) at 80% and aligned to the genome with TBLASTN followed by exonerate splice-site polishing in MAKER. Gene calls resulted in 17,438 predicted protein-coding genes. Raw Illumina reads from the remaining genomes were converted to FASTQsanger using Galaxy tools (https://usegalaxy.org/) and mapped to the F. euwallaceae genome using CLC Genomics Workbench (Qiagen Bioinformatics). A track list was created in CLC Genomics containing all seven genomes and predicted gene annotations. Contigs were searched manually for potential primer binding sites that would selectively anneal to DNA from each target species, due to unique nucleotide polymorphisms and insertion/ deletions (Fig. 1), and yielded an amplicon between 200 and 2,000 bp.

Fusarium spp.-specific oligonucleotide primer development and PCR multiplexing. In all, 3 to 15 different primer pairs per Fusarium sp. were tested for specificity against a panel that included all 12 AFC species but only 1 primer pair per species was selected due to nontarget amplifications in the other primer sets (Integrated DNA Technologies, Coralville, Iowa). Potential species-specific primers were tested in PCR using DNA from two representatives each of AF-1 through AF-12, except for AF-5 and AF-10, where only one reference strain was available (Table 1). Primers were also tested using DNA from three AFC strains previously identified as putative interspecific hybrids (Kasson et al. 2013). All PCR were performed in 25.5-µl reactions containing 12.5 µl of Bioline HS 2× PCR Mix (Bioline USA Inc., Taunton, MA), 10 μl of sterile distilled water, 1 µl each of 10 nM oligonucleotide primer, and 1 µl of genomic DNA (gDNA). Optimal annealing temperatures were determined on a gradient from 57 to 67°C across all primer sets (data not shown). All reactions were carried out in an MJ Research thermocycler with the following conditions: 95°C for 2 min, followed by 34 cycles of 95°C for 30 s, the optimal annealing temperature (Table 1) for 30 s, and 72°C for 60 to 90 s, depending on expected target length (Table 1). PCR products (10 µl each) were visualized on 1.5% (wt/vol) agarose gels with 4 μl of 1× SYBR Gold (Invitrogen, Grand Island, NY) and 4 µl of 1× gel loading dye (5Prime, Gaithersburg, MD) in 0.5% Tris-borate-EDTA buffer (Amresco, Solon, OH). Gels were run at 80 to 90 V for 1.5 to 2 h before visualizing on a UV transilluminator (Syngene, Frederick, MD). Amplicons were sized using 100-bp and 1-kb molecular weight ladders (Omega Bio-tek, Nor-

Primer pairs were considered species-specific if amplicons were only produced for a given target species. To confirm that the expected region had been amplified, amplicons were purified using EXOSAP-IT (Affymetrix, Santa Clara, CA) and sequenced using

the same forward and reverse primers used in PCR (Eurofins Operon, Huntsville, AL). After the six primer pairs were designed and tested, three multiplex assays were developed that combined primers specific to the following three pairs of AFC species based on their geographic proximity: *F. euwallaceae* (AF-2) and AF-12 in southern

California, AF-3 and AF-4 in the eastern United States, and AF-6 and AF-8 in Miami-Dade County, FL. To assess whether the assays could coamplify the target loci from each pair of AFC species, a 1:1 mixture of template DNA from each pair was included in each assay.

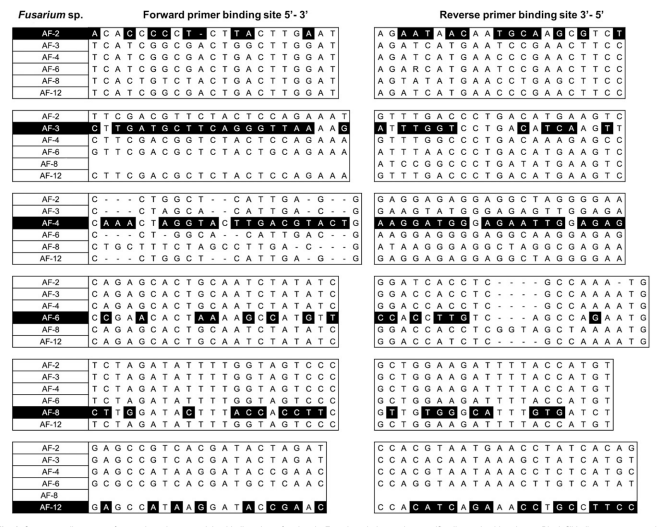


Fig. 1. Sequence alignments of genomic regions containing binding sites of ambrosia Fusarium clade species-specific oligonucleotide primers. Black fill indicates target species and nucleotide polymorphisms within primer binding sites. Blank sequence indicates no genomic coverage available.

Table 1. Primer sequences for taxon-specific marker loci used to distinguish six ambrosia Fusarium clade (AFC) species^a

Target	Forward primer 5'-3'	Reverse primer 5'-3'	Amp (bp)	T (°C)	Ext (min)	GenBank	BLASTp search results
AF-2	ACACCCCCTCTTACTTGAAT	AGACGCTTGCATTGTTATTCT	260	62	1:00	KT835019	Fungal transcription factor regulatory middle homology region
AF-3	CTTGATGCTTCAGGGTTAAAAG	AACTTGATGTCAGGACCAAAT	1,370	64	1:10	KT835020	Rossmann-fold NAD(P)(+)- binding proteins
AF-4	CAAACTAGGTACTTGACGTACTG	CTCTCCAATTCTCCCATCCTT	751	64	1:10	KT835021	Adenylate forming domain, Class I superfamily bifunctional fatty acid transporter/very-long- chain acyl-CoA synthetase
AF-6	CCGAACACTAAAAGCCATGTT	CATTCTGGCTGACAAGGTGG	1,202	65	1:30	KT835022	No putative conserved domains detected
AF-8	CTTGGATACTTTACCACCTTC	AGATCACAAATGCCCACAAC	621	65	1:30	KT835023	Uncharacterized conserved protein (DUF2196)
AF-12	GAGCCATAAGGATACCGAAC	GGAAGGCAGGTTTCTGATGTGG	1,243	62	1:00	KT835024	No putative conserved domains detected

^a Target = target Fusarium spp., Amp = expected amplicon size, T = annealing temperature, Ext = extension time, and GenBank = GenBank accession numbers.

Primer validation and conservation of primer binding sites. In addition to the fusaria mentioned above, DNA from nine clade 3 FSSC species related to the AFC were included in the test of specificity of the six primer pairs (Table 2). This panel contained two representatives of phylospecies FSSC 6, 8, and 10, which are the closest relatives of the AFC (Table 2) (O'Donnell et al. 2008). Of the 18 non-AFC FSSC strains chosen for validation, 10 were fusaria recovered from several bark beetle species (i.e., FSSC 13, 14, 18, 25, and 29).

To further confirm the utility of primer binding sites, 25 additional AF-4 strains were obtained from mature E. validus collected in 2014 to 2015 from multiple locations in the United States (Supplementary Table S2). E. validus was chosen primarily because of its longer residency (Wood 1982) compared with other Euwallacea spp. currently established in the United States and its wide distribution throughout the eastern United States (Cognato et al. 2015). Samples were collected as far south as Clarkesville, GA (Rabun County) and as far north as Huntingdon, PA (Huntingdon County, PA). DNA was extracted from Fusarium cultures derived from single conidia, as previously described (Kasson et al. 2013). Templates were used in PCR containing two primer pairs in the following combinations: primers specific to AF-3/AF-4, AF-6/AF-8, and AF-2/AF-12. In addition to testing whether primer binding sites were conserved in Fusarium symbionts in E. validus throughout the beetle's invaded range in the United States, multiplexing also permitted testing hypotheses regarding symbiont cocultivation and symbiont switching, both of which have been documented in *Euwallacea* spp. (O'Donnell et al. 2015). This was especially relevant given that E. interjectus overlaps with E. validus at the southernmost extent of the latter species' known range, and the two species vector different AFC species.

Sensitivity. The sensitivity of simplex PCR and the three multiplexes was tested using DNA extracted from pure cultures. For simplexes (individual-species PCR), two reference isolates from each of the six target AFC species were used (Table 3). Genomic DNA was diluted to the following concentrations: 100, 50, 25, 10, and 1 ng/ μ l and 100, 10, and 1 pg/ μ l. Concentrations were verified using a NanoDrop spectrophotometer (NanoDrop Products, Wilmington, DE). PCR assays were performed in duplicate using 1 μ l of gDNA template per reaction. To test the sensitivity of multiplexes, the dilutions of 100, 25, and 1 ng/ μ l were used in mixed template reactions, which utilized 1 μ l of gDNA from each isolate of the two different AFC species.

Amplification of fusarial DNA directly from mature beetle heads. *E. validus* heads that were removed from preemergence, mature females in tree-of-heaven (*A. altissima*) stems in West Virginia and mature *E. interjectus* from infested boxelder in Florida were placed individually or in groups of 2, 5, 10, or 20 in 1.5-ml Eppendorf tubes. Whole DNA was assayed using each of the three multiplex PCR. Samples containing two or more heads permitted simultaneous processing of multiple individuals from the same colony to determine what AFC species they vectored. DNA was extracted as previously described (Short et al. 2015).

Results

Primer validation and PCR multiplexing. All six primer pairs proved to be species specific and capable of distinguishing ambrosial fusaria AF-2, AF-3, AF-4, AF-6, AF-8, and AF-12 (Table 2). Amplicons of the expected size were produced for the two target species in each of the three multiplex assays, and not for the six other AFC species or the nine closely related FSSC clade 3 species (Table 2; Fig. 2). Amplicon sizes were as follows: AF-3 and AF-4: 1,370 and 751 bp, AF-6 and AF-8: 1,202 and 621 bp, and AF-2 and AF-12: 260 and 1,243 bp. PCR of the putative triparental interspecific hybrid NRRL 62605 *F. ambrosium* (AF-1) from Sri Lanka produced a positive genotype using the AF-3 primer pair. The sequence from the putative hybrid, which was identified as a NAD(P)-binding Rossmann fold domain-specific protein, was 100% identical to the amplicon obtained from NRRL 62606 (AF-3) *Fusarium* sp. from Gainesville, FL (Table 1). BLASTp search results of protein sequences

revealed that all but two translated sequences (AF-6 and AF-12 species-specific marker loci) contained conserved protein domains (Table 2).

Sensitivity. All six primer pairs were able to amplify their respective target of the gDNA template at or below 1 ng/μl (Table 3). The primer pairs for two AF-4, one AF-6, and two AF-8 isolates showed sensitivity several orders of magnitude below 1 ng/μl, with positive detection at or below 10 pg/μl (Table 3). Multiplex PCR using mixed template gDNA resulted in successful coamplification for seven of nine combinations at 100 ng/μl, 8 of 11 combinations at 25 ng/μl, and 5 of 11 at 1 ng/μl (Fig. 3; Supplementary Table S3). At least one of the two PCR products amplified in all reactions. A majority of the multiplexes that failed involved mixed templates of AF-3 and AF-4. The remaining coamplifications that failed were at the 1 ng/μl concentration and included at least one mixed template of AF-2 and AF-12, AF-6 and AF-8, and AF-3 and AF-4.

Amplification of fusarial DNA directly from beetle heads. The AF-3/AF-4, AF-2/AF-12, and AF-6/AF-8 PCR multiplexes were tested on DNA templates extracted from 1, 2, 5, 10, and 20 heads of freshly caught *E. validus* and *E. interjectus* from the eastern United States. The AF-3/AF-4 multiplex yielded amplicons of the expected size for both species in all batched samples (i.e., 2 to 20 heads) but amplicons were not obtained from individual heads. The AF-3-specific target was amplified from all *E. interjectus* and an AF-4-specific target was amplified from all *E. validus* (Table 2), which demonstrates both the specificity of our assay as well as the specificity of these particular beetle–fungus associations within the United States. As expected, AF-2/AF-12 and AF-6/AF-8 multiplexes yielded no amplicons from any of the *E. validus* or *E. interjectus* DNA extractions.

Discussion

In this study, we validated a rapid diagnostic tool for characterizing fusarial symbionts of *Euwallacea* ambrosia beetles present in the United States. Primer specificity and reliability were confirmed through a screen of a phylogenetically diverse set of AFC strains as well as representatives of nine closely related species within clade 3 of the FSSC (n = 72). PCR were further validated on whole DNA extracted from macerated beetle heads containing conidia. For both fungal and whole-beetle DNA, all primer annealing sites appeared to be species specific and conserved intraspecifically and even added support for a tripartite hybrid origin of NRRL 62605 *F. ambrosium* (Kasson et al. 2013).

Other methods for identifying *Fusarium* symbionts cultivated by *Euwallacea* spp. have relied on analysis of DNA sequence data from phylogenetically informative loci (O'Donnell et al. 2015). A recent method using arbitrarily primed (ap)-PCR was able to discern *F. euwallaceae* from several strains of AF-1 and AF-4 (Freeman et al. 2013). ap-PCR is inexpensive and convenient, in that it only requires three universal primers with no additional genomic data. However, without testing additional isolates representing the remaining nine AFC species and other FSSC, it remains unclear whether ap-PCR could resolve all of the AFC species. Taxon-specific PCR assays such as the one developed here are reliable and easily interpreted, and have been developed in several important pathosystems (Dreaden et al. 2014; Inderbitzin et al. 2013).

The newly designed primer sets amplify their correct targets in concentrations at or below 1 ng/µl for all gDNA templates, and at or below 10 pg/µl for three AFC species, which is comparable with previously reported sensitivity for multiplex assays of phytopathogenic fungi (Dreaden et al. 2012; Rigotti et al. 2002) and bacteria (Balestra et al. 2013). The multiplex assays successfully identified AF-3 and AF-4 from pooled DNA from eight batches of freshly macerated *E. interjectus* and *E. validus* beetle heads, respectively. These results, coupled with the sensitivity assays, may indicate that singleton beetles contained fungal gDNA below detectable limits. However, more work is needed to confirm the concentration of gDNA from the mycangia of *Euwallacea* ambrosia beetles. Follow-up studies comparing whole-DNA extraction methods from freshly collected and ethanol-preserved beetle specimens are needed to

Table 2. Strain and beetle identification (ID) and results of polymerase chain reaction (PCR) and sequencing experiments

		Tested via multiplex PCR						Sequence confirmed		
Fusarium spp.a	ID_p	AF-2	AF-3	AF-4	AF-6	AF-8	AF-12	EF1-α	AFCc	
AF-1 Fusarium ambrosium	NRRL 20438	-	-	-	-	-	-	+		
AF-1 F. ambrosium	NRRL 62942	_	-	-	_	-	-	+		
AF-2 F. euwallaceae	NRRL 54727	_	_	_	_	_	_	+	+	
AF-2 F. euwallaceae	NRRL 62626 ^d	+	_	_	_	_	_	+		
AF-3 Fusarium sp.	NRRL 62629	_	+	_	_	_	_	+	+	
AF-3 Fusarium sp.	NRRL 62606 ^d	_	+	-	_	_	-	+	+	
AF-3 Fusarium sp.	EiF20e	_	+	_	_	_	-		+	
AF-3 Fusarium sp.	EiF10e	-	+	-	-	-	_		+	
AF-3 Fusarium sp.	EiF5 ^e	-	+	-	-	-	_			
AF-3 Fusarium sp.	EiF2 ^e	-	+	_	_	-	_			
AF-3 Fusarium sp.	EiF1 ^e	-	-	-	-	-	-			
AF-4 Fusarium sp.	NRRL 62579 ^d	-	-	+	_	-	_	+	+	
AF-4 Fusarium sp.	NRRL 62578	-	-	+	-	-	_	+		
AF-4 Fusarium sp.	NRRL 62582	-	-	+	_	-	_	+		
AF-4 Fusarium sp.	EvF20e	-	-	+	-	-	_		+	
AF-4 Fusarium sp.	EvF10e	_	_	+	_	_	_		+	
AF-4 Fusarium sp.	EvF5e	_	_	+	_	_	_			
AF-4 Fusarium sp.	EvF2e	_	_	+	_	_	_			
AF-4 Fusarium sp.	EvF1e	_	-	-	_	_	_			
AF-5 Fusarium sp.	NRRL 22231	-	-	-	-	-	_	+		
AF-6 Fusarium sp.	NRRL 62591	_	_	_	+	_	_	+		
AF-6 Fusarium sp.	NRRL 62590 ^d	_	-	_	+	-	_	+	+	
AF-6 Fusarium sp.	KOD 133	_	_	_	+	_	_	+		
AF-6 Fusarium sp.	KOD 134	_	_	_	+	_	_	+		
AF-7 Fusarium sp.	NRRL 62610	_	_	_	_	_	_	+		
AF-7 Fusarium sp.	NRRL 62611	_	_	_	_	_	_	+		
AF-8 Fusarium sp.	NRRL 62585	_	_	_	_	+	_	+	+	
AF-8 Fusarium sp.	NRRL 62584d	_	_	_	_	+	_	+		
AF-9 Fusarium sp.	NRRL 22643	_	_	_	_	_	_	+		
AF-9 Fusarium sp.	NRRL 66088	_	_	_	_	_	_	+		
AF-10 Fusarium sp.	NRRL 62941	_	_	_	_	_	_	+		
AF-11 Fusarium sp.	NRRL 62944	_	_	_	_	_	_	+		
AF-11 Fusarium sp.	NRRL 62943	_	_	_	_	_	_	+		
AF-12 Fusarium sp.	NRRL 62945	_	_	_	_	_	+	+	+	
AF-12 Fusarium sp.	KOD 792d	_	_	_	_	_	+	+		
AF-12 Fusarium sp.	NRRL 62946	_	_	_	_	_	+	+		
AF-1 F. ambrosium (hybrid)	NRRL 46583	_	_	_	_	_	_	+		
AF-1 F. ambrosium (hybrid)	NRRL 22345	_	_	_	_	_	_	+		
AF-1 F. ambrosium (hybrid)	NRRL 62605	_	+	_	_	_	_	+	+ (AF-3)	
FSSC 6-p	KOD 327	_	_	_	_	_	_	+	`	
FSSC 6-s	KOD 330	_	_	_	_	_	_	+		
FSSC 8-a (F. neocosmosporiellum)	NRRL 43467	_	_	_	_	_	_	+		
FSSC 8-b	NRRL 22436	_	_	_	_	_	_	+		
FSSC 10-a	NRRL 22135	_	_	_	_	_	_	+		
FSSC 10-b	NRRL 22098	_	_	_	_	_	_	+		
FSSC 11-1	KOD 313	_	_	_	_	_	_	+		
FSSC 11-o	KOD 320	_	_	_	_	_	_	+		
FSSC 13-g	KOD 398	_	_	_	_	_	_	+		
FSSC 13-k	KOD 267	_	_	_	_	_	_	+		
FSSC 14-a	KOD 450	_	_	_	_	_	_	+		
FSSC 14-b	KOD 450 KOD 452	_	_	_	_	_	_	+		
FSSC 18-f	KOD 432 KOD 288	_	_	_	_	_	_	+	•••	
FSSC 18-m	KOD 288 KOD 312	_	_	_	_	_	_	+		
FSSC 25-f	KOD 312 KOD 384	_	_	_	_	_	_	+	•••	
FSSC 25-k	KOD 389	_		_	_	_	_			
FSSC 23-k FSSC 29-d	KOD 389 KOD 411	_	_	_	_	_	_	+		
FSSC 29-e	KOD 411 KOD 407	_	_	_	_	_	_	+		

^a An informal ad hoc nomenclature was developed to distinguish the 12 fusaria within the ambrosia Fusarium clade (AFC) (i.e., AF followed by 1 to 12). FSSC = Fusarium solani species complex.

b Strain designations are as follows: NRRL = ARS Culture Collection, Peoria, IL and KOD = lab collection of Kerry O'Donnell.

^c AFC-specific targets.

^d Genome was sequenced in the current study.

^e DNA was extracted directly from beetle heads. Ei = Euwallacea interjectus, Ev = E. validus, and F = female beetles. Number following F indicates number of heads included in single DNA extraction.

optimize extraction and amplification, especially at the single-beetle level. This is particularly important given that fungal communities within mycangia are diverse (Kostovcik et al. 2015) and dynamic (Freeman et al. 2016), changing as adult females mature and prepare to emerge from their natal galleries. Results of multiplex PCR using mixed templates confirmed successful coamplification for a majority of combinations at or above 25 ng/µl and several at 1 ng/µl, especially for AF-2/AF-12 mixed templates. Such coamplification is important when testing interactions (symbiont cocultivation or swapping) among Euwallacea ambrosia beetles in areas where their invaded ranges already overlap or where overlap is imminent. The latter includes the southeastern United States and California, where two or more species of exotic Euwallacea beetles might colonize the same woody host (Eskalen et al. 2013; O'Donnell et al. 2015). Given the fact that Euwallacea sp. #1 farming F. euwallaceae, the AF-2-specific PCR assay should prove useful in tracking its spread into new areas. The AF-2/AF-12 multiplex will likely prove most useful in that *Euwallacea* spp. 1 and 5, farming AF-2 and AF-12, respectively, are present in southern California and pose a potential threat to avocado production in the commercial groves there and in Mexico.

Globally, several Fusarium canker diseases have inflicted widespread mortality and economic loss on both native and cultivated crop and timber trees. Canker pathogens in the *F. torreyae* species complex currently threaten endemic Florida torreya (*Torreya taxifolia* Arn.) and cultivated prickly ash (*Zanthoxylum bungeanum* Maxim.) in China (Aoki et al. 2013; Smith et al. 2011; Zhou et al. 2016). Certain members of the FSSC have also been implicated as causal agents in the disease complex thousand cankers disease of walnut (*Juglans* spp.) in western United States and Europe (Montecchio et al. 2015; Tisserat et al. 2009). *F. circinatum*, a member of the *F. fujikuroi* species complex (Nirenberg and O'Donnell 1998) and causal agent of pitch canker,

Table 3. Results of the polymerase chain reaction simplexes targeting six ambrosia *Fusarium* clade (AFC) species

Sample	AFC species	IC (ng/μl) ^b	Approximate concentrations ^a							
			100 ng/μl	50 ng/μl	25 ng/μl	10 ng/μl	1 ng/μl	100 pg/μl	10 pg/μl	1 pg/μl
54727	AF-2	47	n/a	+	+	+	+	_	_	_
62626	AF-2	115	+	+	+	+	+	_	_	_
MB202	AF-3	94	+	+	+	+	+	_	_	_
62605	AF-3c	104	+	+	+	+	+	_	_	_
MB14	AF-4	112	+	+	+	+	+	+	+	+
MB42	AF-4	99	+	+	+	+	+	_	+	+
62591	AF-6	103	+	+	+	+	+	+	_	+
62590	AF-6	99	+	+	+	+	+	+	_	_
62585	AF-8	84	+	+	+	+	+	+	+	+
62584	AF-8	117	+	+	+	+	+	+	+	_
62945	AF12	75	+	+	+	+	+	_	_	_
62946	AF12	98	+	+	+	+	+	_	_	_

a Initial DNA template was approximately 50 ng/μl; therefore, no PCR was conducted for the approximately 100 ng/μl concentration (n/a).

^c Multiparent interspecific hybrid.

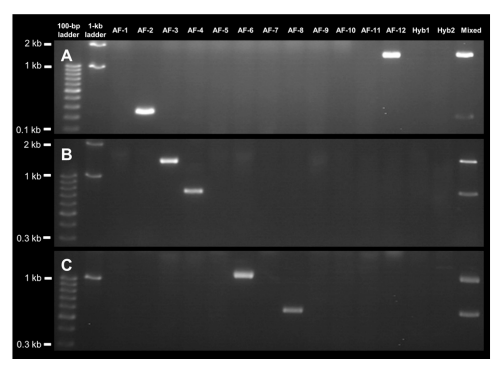


Fig. 2. Gel electrophoresis of amplicons generated using three different species-specific multiplexes. **A**, AF-2/AF-12; **B**, AF-3/AF-4; **C**, AF-6/AF-8. Strains used in multiplexes are as follows: AF-1 = NRRL 20438, AF-2 = NRRL 54727, AF-3 = NRRL 62629, AF-4 = NRRL 62579, AF-5 = NRRL 22231, AF-6 = NRRL 62590, AF-7 = NRRL 62610, AF-8 = NRRL 62585, AF-9 = NRRL 22643, AF-10 = NRRL 62941, AF-11 = NRRL 62944, AF-12 = NRRL 62945, Hyb1 = NRRL 46583, and Hyb2 = NRRL 22345. Mixed templates consisted of DNA from the same strains listed.

^b Initial concentration.

continues to affect numerous pine species throughout the northern and southern hemispheres (Wingfield et al. 2008).

The spread of Fusarium tree diseases, including pitch canker, which has been confirmed on dozens of native and cultivated pine species and associated with numerous insect vectors, is of great concern to countries such as Australia and New Zealand, where extensive plantations of highly susceptible *Pinus radiata* are grown (Dick 1998). In response to these threats, several molecular methods for the rapid and sensitive detection of *F. circinatum* from bark beetles (Fourrier et al. 2015), seed (Dreaden et al. 2012), and spore traps (Schweigkofler et al. 2004) were developed. These molecular tools have likely mitigated additional spread of pitch canker, especially in infected nursery stock, which historically relied on less accurate and less sensitive culture-based screening methods for detection of pathogens, which were prone to false negatives (Dreaden et al. 2012).

Fusarium canker and associated dieback, caused by members of the monophyletic AFC and their *Euwallacea* vectors, also affects numerous tree species within the United States, as well as cultivated crop and timber species planted worldwide. Several species, including boxelder, California sycamore, and native willow species, are

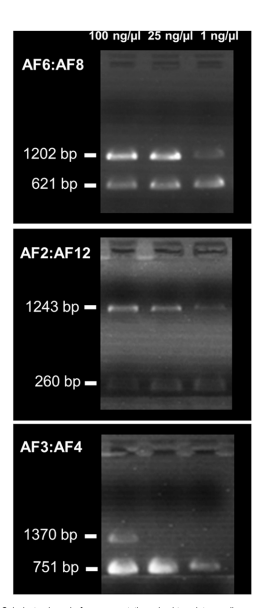


Fig. 3. Gel electrophoresis for representative mixed-template amplicons generated using three different species-specific multiplexes across three different genomic DNA (gDNA) concentrations. Concentrations included 100, 25, and 1 ng/ μ l. Mixed templates consisted of 1 μ l of gDNA from each of two strains. Mixed templates were as follows: AF-6/AF-8 = NRRL 62591/NRRL 62584, AF-2/AF-12 = NRRL 62626/NRRL 62945, and AF-3/AF-4 = NRRL 62605/MB14.

currently at risk for widespread beetle infestations and associated mortality, especially in California. *Euwallacea* spp. within the *E. fornicatus* species complex have been confirmed attacking native vegetation in many new areas throughout San Diego, Los Angeles, Orange, and Riverside Counties (Boland 2016; Lynch et al. 2016). Avocado has also been affected in California, Israel, and partially in Florida, although the severity of damage is highly variable regionally and temporally (Mendel et al. 2012; Freeman et al. 2013). Future introductions of exotic beetles are anticipated and the ranges of established polyphagous *Euwallacea* spp. are expected to expand. As beetle populations grow and disseminate, additional native plant species will likely be affected, including some that may serve as suitable reproductive hosts for the beetles.

The availability of inexpensive whole-genome sequencing coupled with user-friendly sequence analysis software greatly facilitated identification of species-specific primer binding sites for distinguishing the six AFC species currently known within the United States. Whole-genome sequences of closely related fungal species has provided new insight into many other economically important fungal pathogens (Jones et al. 2014; Schmidt et al. 2016), including other fusaria (Maphosa et al. 2016). This diagnostic tool can easily be expanded in the event that other AFC-Euwallacea mutualists are introduced into the United States. Individual (simplex) diagnostic primer pairs can economize screening in regions where only a single AFC species is present, such as AF-2 in Israel. PCR assays provide several obvious advantages over multilocus sequencing for identifying AFC species, such as increased speed and volume of samples that can be screened as well as reduced cost. In summary, the PCR assays to discriminate Fusarium symbionts of exotic Euwallacea ambrosia beetles provide a foundation for rapid and widespread molecular surveillance focused on tracking changes in AFC species' host range and geographic distribution over time.

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