

## PRIMER NOTE

## AN EXPANDED NUCLEAR PHYLOGENOMIC PCR TOOLKIT FOR SAPINDALES<sup>1</sup>

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- *Premise of the study:* We tested PCR amplification of 91 low-copy nuclear gene loci in taxa from Sapindales using primers developed for *Bursera simaruba* (Burseraceae).
- *Methods and Results:* Cross-amplification of these markers among 10 taxa tested was related to their phylogenetic distance from *B. simaruba*. On average, each Sapindalean taxon yielded product for 53 gene regions (range: 16–90). *Arabidopsis thaliana* (Brassicaceae), by contrast, yielded product for two. Single representatives of Anacardiaceae and Rutaceae yielded 34 and 26 products, respectively. Twenty-six primer pairs worked for all Burseraceae species tested if highly divergent *Aucoumea klaineana* is excluded, and eight of these amplified product in every Sapindalean taxon.
- *Conclusions:* Our study demonstrates that customized primers for *Bursera* can amplify product in a range of Sapindalean taxa. This collection of primer pairs, therefore, is a valuable addition to the toolkit for nuclear phylogenomic analyses of Sapindales and warrants further investigation.

**Key words:** Anacardiaceae; Burseraceae; low-copy nuclear genes; microfluidic PCR; Rutaceae.

Low-copy nuclear gene regions offer increased phylogenetic utility for species- and population-level studies of plants as compared to chloroplast and nuclear ribosomal markers (Zimmer and Wen, 2012), yet sampling these regions remains challenging due to the dearth of universal primers and barriers to sequencing whole or partial nuclear genomes from multiple individuals. Consequently, assessing the phylogenetic limits of custom-designed target sequences or primers for low-copy nuclear gene regions is critical to fully realizing their broader impacts for advancing plant systematics. We report the results of a cross-amplification study incorporating primers for 91 low-copy nuclear gene loci created by Gostel et al. (2015) for species-level phylogenetics of Malagasy *Commiphora* Jacq. (Burseraceae). Primers for these markers were developed using genomic resources from two rosid orders by mapping sequence data from a transcriptome of *Bursera simaruba* (L.) Sarg. (Burseraceae; Sapindales) (Matasci et al., 2014) to 950 putative low- or single-copy nuclear gene loci of *Arabidopsis thaliana* (L.) Heynh. (Brassicaceae; Brassicales) (Duarte et al., 2010). Gostel et al. (2015) further optimized the primers for microfluidic

PCR-based target enrichment, a method that allows simultaneous and cost-effective amplification of multiple loci (Blow, 2009; Uribe-Convers et al., 2016).

We tested cross-amplification of these primers using 10 taxa that have varying phylogenetic distances from *B. simaruba* within Sapindales and included *A. thaliana* as the outermost limit of the survey. Sapindales is a widespread group that includes ca. 6700 species within nine families (Angiosperm Phylogeny Group, 2016) (Fig. 1). Molecular phylogenies of this order often lack sufficient phylogenetic support along their backbone as well as at the species level (e.g., Fine et al., 2014; Grudinski et al., 2014), thus our understanding of Sapindalean systematics could benefit from an expanded phylogenetic toolkit such as that provided by the Gostel et al. (2015) primers.

### METHODS AND RESULTS

**Taxonomic sampling and molecular methods**—Appendix 1 contains accession information for the 11 taxa sampled; Fig. 1 displays their phylogenetic relationships. *Bursera simaruba* (*Bursera* Jacq. ex L. subgenus *Bursera*) and *C. grandifolia* Engl. were included as positive controls; prior work has shown that all or most of the custom-designed primers amplify PCR product in these two species (Gostel et al., 2015). For experimental taxa, we included *B. tonkinensis* Guillaumin, which is sister to *Commiphora* (Weeks and Simpson, 2007), as well as *Aucoumea* Pierre, the monotypic genus sister to *Bursera* and *Commiphora* (Weeks et al., 2014). One species from each of *Boswellia* Roxb. ex Colebr., *Canarium* L., and *Protium* Burm. f. were included, as well as *Beiselia* Forman, the monotypic genus sister to all other Burseraceae (Weeks et al., 2014). We included one species of Anacardiaceae, the family that is sister to Burseraceae (Weeks et al., 2014), and one species of Rutaceae, which represents the Sapindalean clade sister to Burseraceae–Anacardiaceae–Kirkiaeae (Muellner-Riehl et al., 2016). *Arabidopsis thaliana* (Brassicaceae) was included because its genomic resources were used in primer design and can test the applicability of these primers to other closely related rosid lineages (Wang et al., 2009).

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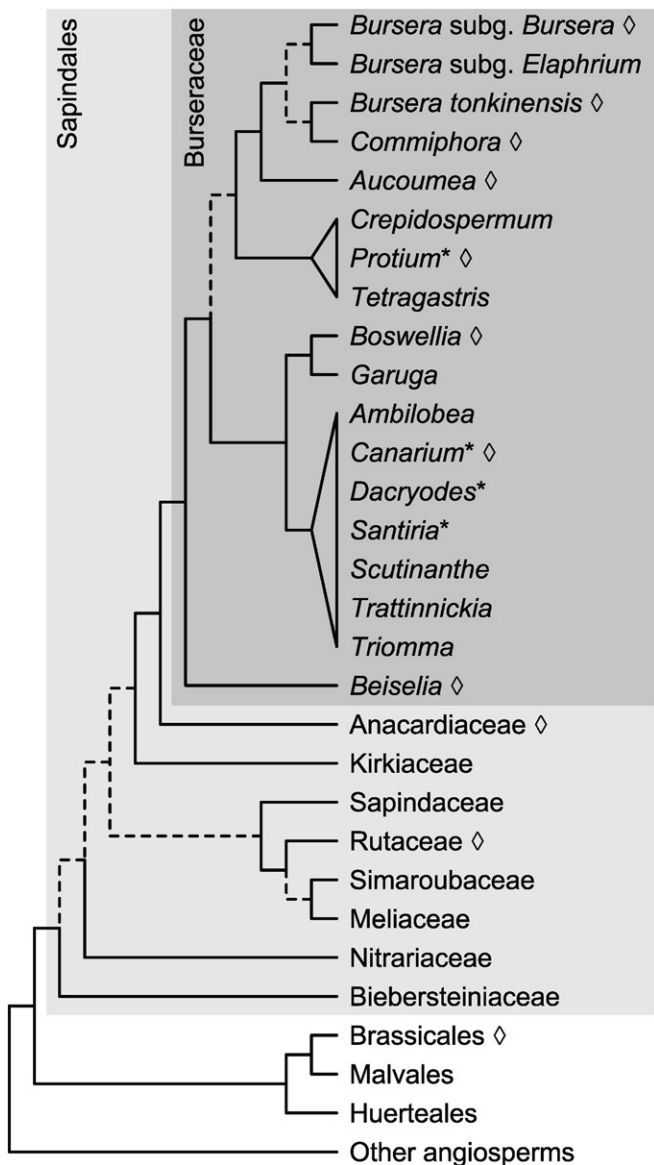


Fig. 1. Phylogeny of Sapindalean lineages condensed from Wang et al. (2009), Weeks et al. (2014), and Muellner-Riehl et al. (2016); nodes having low or conflicting support are indicated by dashed branches. Lineages sampled by the current study are noted by open diamonds. Generalized generic phylogeny of Burseraceae does not depict *Rosselia* or *Pseudodacryodes*, which have not been included in any molecular phylogenetic analysis; paraphyletic genera are indicated by asterisks.

Whole genomic DNA was extracted from taxa using the FastPrep FastDNA Spin Kit (Bio101 Systems, La Jolla, California, USA) or the cetyltrimethylammonium bromide (CTAB) method (Weeks et al., 2005). Primer development for the 91 markers is detailed by Gostel et al. (2015); primer sequences are listed in Table 1. Markers were amplified via PCR in 15- $\mu$ L reactions including: 0.15  $\mu$ L of forward and reverse primers (50  $\mu$ M), 0.75  $\mu$ L spermidine (4 mM), 7.5  $\mu$ L GoTaq Green Master Mix (Promega Corporation, Madison, Wisconsin, USA), 5.6  $\mu$ L nuclease-free water, and 1  $\mu$ L genomic DNA (0.1–25.8 ng/ $\mu$ L). Markers that failed to amplify for *B. simaruba* and *C. grandifolia* were then trialed using reaction chemistry based on that recommended for microfluidic PCR-based target enrichment including: 0.15  $\mu$ L of forward and reverse primers (50  $\mu$ M); FastStart High Fidelity PCR System reagents (Roche Diagnostics, Mannheim, Germany), composed of 1.5  $\mu$ L FastStart High Fidelity Reaction Buffer without MgCl<sub>2</sub> (10x concentration), 2.7  $\mu$ L MgCl<sub>2</sub> (25 mM), 0.75  $\mu$ L DMSO, 1.2  $\mu$ L Nucleotide Mix (10 mM), 0.15  $\mu$ L FastStart High Fidelity Enzyme Blend

(5 U/ $\mu$ L); 0.75  $\mu$ L Loading Reagent (Fluidigm Corporation, San Francisco, California, USA); 6.8  $\mu$ L nuclease-free water; and 1  $\mu$ L genomic DNA.

The PCR thermocycler protocol followed that of Gostel et al. (2015) and included three alternating standard and C<sub>o</sub>t cycles (Mathieu-Daude et al., 1996), beginning with 2 min at 50°C, 20 min at 70°C, and 10 min at 95°C. The first set of 10 standard cycles included a denaturation step at 95°C for 15 s, annealing at 60°C for 30 s, and extension at 72°C for 1 min. Two C<sub>o</sub>t cycles followed, including four steps consisting of 95°C for 15 s, 80°C for 30 s, 60°C for 30 s, and 72°C for 1 min. Standard and C<sub>o</sub>t cycles alternated two more times with eight, two, eight, and five cycles, respectively. After 35 cycles, samples were held at 4°C prior to being visually verified via agarose gel electrophoresis (1% agarose; 94 V for 40 min). Low DNA mass ladder (Invitrogen, Carlsbad, California, USA) was included in the first and last wells of each gel to guide length estimation of PCR products.

**Marker amplification results**—Table 1 contains amplification results for the low-copy nuclear loci, including the range of amplicon lengths for all taxa and GenBank numbers for markers sequenced by Gostel et al. (2015) for *B. simaruba* and *C. grandifolia* that had  $\geq 15$  sequence reads mapped. Table 2 summarizes marker amplification success for each taxon. Ninety primer pairs amplified product in *B. simaruba* and, on average, 54 primer pairs worked for other Burseraceae taxa. The low number of markers amplified in *Aucoumea* (16) was unexpected given its close relationship to *Bursera*. This result may have been caused by primer mismatch due to increased genetic change within this monotypic genus, as evidenced by its long branch within Burseraceae phylogeny (Weeks et al., 2014). In total, nine primer pairs worked for every Burseraceae taxon tested, and if *Aucoumea* is excluded as an outlier, the panel of family-universal primer pairs increases to 26. Thirty-four and 26 primer pairs generated product in Anacardiaceae and Rutaceae, respectively, while only two primer pairs worked in *Arabidopsis*. Comparing the Burseraceae panel to that of Anacardiaceae and Rutaceae reveals 16 and 12 successfully amplified regions in common, respectively, with eight shared among the three families. PCR chemistry may have suppressed amplification of markers, as high-fidelity PCR reagents were not used due to their high cost. Among the positive controls, high fidelity as compared to standard PCR reagents increased amplification success by 8% (*Bursera*, 83 to 90 primer pairs) and 85% (*Commiphora*, 39 to 72 primer pairs). Thus, our experimental results report a conservative baseline for the cross-amplification success of these primer pairs.

## CONCLUSIONS

Our study demonstrates that 90 of 91 primer pairs for novel low-copy nuclear loci developed by Gostel et al. (2015) for *B. simaruba* successfully amplify product in a broad range of Sapindalean taxa and effectively expand the phylogenomic toolkit for this order. Twenty-six markers amplify all Burseraceae taxa (excluding *Aucoumea*) and eight amplify all Sapindalean groups tested. Our results present a new source for universal targets or primers for phylogenetic reconstruction of taxa within Sapindales. Future efforts will include sequencing amplicons to determine the number of phylogenetically informative characters for each locus.

## LITERATURE CITED

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TABLE 1. Primer pair sequences and validation results by taxon.

Locus ID <sup>a</sup>	Primer sequences (5'-3') <sup>a</sup>	GenBank accession no. <sup>b</sup>	B. simaruba	C. grandifolia	Amplicon length range among all taxa
AT3G54460 <sup>c</sup>	F: GGACACACCCTGGCTCTAG R: CTCCCATGACTTTGGTTCTGTC	KX767982	KX767983	270–290	X X X X X X X X X X
AT2G04620	F: TCCACCATATTGAGTGAAGCAA R: AATGGGAGTGGAAATGAGATGTTG	KX76792	KX767929	420–520	X X X X X X X X X X
AT4G37510 <sup>c</sup>	F: TTCAATTGAGACCTCCATAGATGAC R: GCTTAGCCGGATTAACTGTCTGC	KX76800	KX76800	280	X X X X X X X X X X
AT3G22660 <sup>c</sup>	F: AGATGAGATGAAATTGGTGAAC R: TTTCCTGCTTAAGCTCTCTTCTATCT	KX76794	KX76795	450	X X X X X X X X X X
AT1G21840 <sup>c</sup>	F: TGTTGGAGAACCTGAAGAGAGG R: CACCAATTATCCAAACCTCTGAA	KX767930	KX767931	630–640	X X X X X X X X X X
AT2G04740 <sup>d</sup>	F: CAAACTCCAAAACCTAAACCGG R: TCAAAAGCCCTCAAAAGCTCTCCTC	KX767986	KX767987	460–590	X X X X X X X X X X
AT4G14605 <sup>d</sup>	F: CTCTCACTCATAGCAGGAGAAG R: CTCTCTCACAGCTTATAAAGTCA	KX767990	KX767991	510–580	X X X X X X X X X X
AT4G19900 <sup>c</sup>	F: GTTCTCTGAGCGGATTGAGCTTGA R: CTGTAGAGAGCAAGTGG	KX767994, KX767995, KX767996, KX768005	KX767995, KX767997, KX768006	350–420	X X X X X X X X X X
AT4G29590	F: GAGCAATTCCCCTCAAGAGCA R: GTGCTTGTAACTCTTTGGTATGG	KX767994, KX767995, KX767996, KX768005	KX767995, KX767997, KX768006	490	X X X X X X X X X X
AT5G04910	F: TAAGAGTCCACAGCAGCATGAGT R: TAAAGAAATGATGCACTAGCTTCG	KX767994, KX768005	KX767995, KX767997, KX768006	260	X X X X X X X X X X
AT3G15110 <sup>e</sup>	F: CTCACTGGTGCCATATGCTGCTC R: ATTCTCTGTAACCTTGGCTCTGGA	KX767902	KX767903	1560	X X X X X X X X X X
AT1G18060 <sup>d</sup>	F: AACAAAGAAAAGTTGCAGTAAAGGA R: GCTGGGCTCTGTCACTTTTTG	KX767926	KX767927	740–930	X X X X X X X X X X
AT2G03667 <sup>c</sup>	F: CTAGTTGGCTGGTGTGATG R: CACAAAGGAAATAAACGAAAGTCC	KX767926	KX767927	590	X X X X X X X X X X
AT2G40760	F: GGTTATCATGGAAAGGG R: CGCTCTGCCCTCTCTTTC	KX768007	KX768008	400	X X X X X X X X X X
AT2G20790 <sup>c,g</sup>	F: CCATTGTCATGGTCTCAAGAGATG R: COATGGTGCAATTTAACATTAACGTGTC	KX767940	KX767941	320–350	X X X X X X X X X X
AT2G36740	F: AGTCACAAAGCACTGAGTAT R: CATCCCTTGAGAAATAACCGPATCTGT	KX767958, KX767959, KX767960, KX767961	KX767958, KX767959, KX767960, KX767961	640–810	X X X X X X X X X X
AT3G01380 <sup>d</sup>	F: AATCATCATATAAGGGCAGCG R: COAAGAAATAAGAAGTTAGTGGTGGAC	KX767962	KX767962	530–930	X X X X X X X X X X
AT3G10400	F: CGGTCTTGGTACGCTTAAATAAGC R: GCTACTTGGTACGCTTAAATAAGC	KX767967	KX767967	340	X X X X X X X X X X
AT1G59990 <sup>d</sup>	F: TGACACCACGAAATAAACGACAGC R: AACCCACATGGFACTGTTAACATG	KX767908, KX767909, KX767910, KX767944	KX767908, KX767909, KX767910, KX767944	450–510	X X X X X X X X X X
AT2G22370B <sup>c</sup>	F: CATCAGACATAAGATGAGCAGCAG R: CTGTGCTTGGTACTTGTGATCCA	KX767904, KX767905, KX767906	KX767904, KX767905, KX767907	610–780	X X X X X X X X X X
AT1G31780 <sup>c,g</sup>	F: CTGTGCTTGGTACTTGTGATCCA R: GGACCCAAAAGTGTACTACAGAG	KX767907	KX767907	380–832	X X X X X X X X X X
AT1G31780 (INT) <sup>c,g</sup>	R:				

TABLE 1. Continued.

Locus ID <sup>a</sup>		Primer sequences (5'-3') <sup>a</sup>	GenBank accession no. <sup>b</sup>		Amplicon length range among all taxa
			<i>B. simaruba</i>	<i>C. grandifolia</i>	
AT2G27760	F:	GAACCTTAACCCCTAACAACTGGAGAA			930
	R:	GGCGGTTCCTGGACCATAT			
	F:	GAACCTTAAACCCCTAACAACTGGAGAA			160–470
	R:	CGAAATTCCCTAGCAGTGAACTCC			
	F:	GACGCTGTATCTAGGCTCCAG			220–640
AT1G63160 (INT)	R:	AAAATGTTGCACTGTGAAGTTGGC			
AT1G63160	F:	GACGCTGTATCTAGGCTCCAG			
	R:	CAACATGAGAACAGTAT	KX767912	KX767913	1070–1490
AT1G65030	F:	CGGTTTCTGAAACTCGGGTACAG			
	R:	CGGGGAAAAGAGGGTTTGG			
	F:	CTTCGAGAAATTGGTGGAAATGT	KX768003	KX768004	340
AT5G52180	R:	CATACAGAAAGCCGCTCGATA			
	F:	CAGCATGGAAATACGTGCTAGTA	KX767954	KX767955	
	R:	TATCAACTGGAAACCCCTGGATAAG			
	F:	TGCCAAGTAAACAGATGGTTAAAGG	KX767934	KX767935	
AT2G44760 <sup>d</sup>	R:	TCTCCA AACACTCTGGTTAAAGGA			
	F:	GGGGTGAGAAATGAGAAATGACATG	KX767998	KX767999	580–780
AT4G31770 <sup>c</sup>	R:	ACAAGTTCTCCAATTCCAAA			
	F:	TCATTGAAGGTTGGGATTCAGC	KX767938	KX767939	
	R:	ACGACTTGGCGATCTCTGATAAA			
AT2G20330 <sup>e,g</sup>	F:	CCTCTCTCATAGTGTGCT			
	R:	CCACAAAAGCACTGCATAAAGTT			
AT1G66080	F:	AGTACTTCCCGAGAAATTGAA			
	R:	GCACAGTACATAAACCCATTGGT			
AT2G05170B <sup>c</sup>	F:	TGGCTTGTGGNCATATGAGAACTT			
	R:	CCTAATACTGGAGGAAAATGCT	KX767914	KX767915	
AT5G67220	R:	CAGTACTTCCCGAGAAATTGAA			
	F:	CGGTAAAAAGCTCTCAGATCC			
	R:	CTCTGGCAAGTGAACCCFT	KX767936	KX767937	
AT2G17265 <sup>c,d,g</sup>	F:	TTATGGGAGGTTCTGTTTACAG			
	R:	CTAGACCAACTCTATCCACCTC			
	F:	TCTTTGCTGCTACCTCTCAG			
	R:	CGATGTCGTTCTCTGATATAGCCT			
AT2G31890B <sup>c,g</sup>	F:	CTCTCCAGTGTCACTTTAACAG	KX767946	KX767947	
	R:	CTTGAGAAATCTGTGGTCATCA			
	F:	TITAAGGACTTCGGCTTCAAA	KX767956	KX767957	
	R:	GGCAGAAAAGATAAGCCCTCAG			
AT3G26580 <sup>c,g</sup>	F:	AGGTGAACGGTGGATTATGAT	KX767976,	KX767977,	
	R:	GTGACGGTATTGCTCTGFAAG	KX767978	KX767979	
AT2G44660B	F:	GTTTTGCGAGAAGGGATGATT	KX767952	KX767953	
AT2G44660B (INT) <sup>c</sup>	R:	TGAAGGTTGGCTGGAGTATCT			
	F:	GTTTTGCGAGAAGGGATGATT	KX767952	KX767953	
AT3G49730	R:	COAAAATGGAGATGGCTT			
	F:	AATCAACTCAGGCCTTCTCTC			
	R:	X			

TABLE 1. Continued.

Locus ID <sup>a</sup>	Primer sequences (5'-3') <sup>a</sup>	GenBank accession no. <sup>b</sup>		Amplifon length range among all taxa
		<i>B. simaruba</i>	<i>C. grandifolia</i>	
AT2G44660A <sup>f</sup>	F: ATCGTATCAAGCACAGACATTGAR: R: GAAAAAACAAACACCCATCAA	KX767950	KX767951	790
AT2G21710 <sup>c</sup>	F: TTTCCTCCCTTAACATAAGCCT R: CTTGTCCTGCACCTCTGTGATGAA	KX767942 (5' only)	KX767943 (5' only)	1040-1360
AT2G21710 (INT) <sup>d</sup>	F: TTTCCTCCCTTAACATAAGCCT R: GCTGCCATCCCAGAGCTCTGG			750-860
AT2G22370A <sup>c</sup>	F: ATGGTGAGGCCCTTGAGATCTTC R: TAGTGCTGTACTCAAACAGATT			980-1320
AT1G77930A <sup>d</sup>	F: ACCCTAATTCTGTTCTGCATTG R: GAGCAGTTCAAAAGCAGCTTGAAT	KX767924	KX767925	580-740
AT1G77930A (INT)	F: ACCCTAATTCTGTTCTGCATTG R: GCATCCCTCATGTTACTCTGAATT			410-460
AT5G02250 <sup>d</sup>	F: CACTTATCCCTATGTTCCAAGAAC R: GGATCTGCCCTGTTTCAAATAT			1240-1680
AT2G31440	F: GTATGGAGGGTTCTTCCTTTG R: ATTCTCTGCAAGAGATGAACATCA			1000-1350
AT1G77550A <sup>d</sup>	F: TGTGAGCTTCTCTATATTGTGCC R: TGATGCTTCAGAACAGAACAGA	KX767920	KX767921	740-860
AT3G15290 <sup>e</sup>	F: GATGTTGATGGCTATTGTT R: ATCTGCAAGTCTAAAGGCCAT			1090
AT5G111980	F: TTCAACCATTGATCCCCAAATTAC R: GACAGAGATCCCCCTCAAGTATC		N/A	
AT5G14580 <sup>e</sup>	F: TATACTGTTATGCCAGAATTCCGG R: TCCTGTGCAAACTTATCTAAGGCCT			1030-1750
AT2G31840 <sup>d</sup>	F: AGTGTATTGATGGTGTCCCTGATGT R: CATCTTGGTAGGTGCTACAG			480-1220
AT5G57655	F: TTGGTTATGCUCTGTAATTGCA R: CTACAGTGCACATTGGAAACCAT			340-1340
AT2G47760	F: CAGCATGGAAATACGTTGCTAGTA R: TATCAACTGGACCCCTGGATAAG			620-1480
AT3G29130 <sup>d</sup>	F: TTGCCGAGGTCTGGTGAATT R: AAGTACTTCTCTGTTGATTTCCG	KX767980, KX767981		980-1720
AT3G13200 <sup>e</sup>	F: AACTCATCGCTTTCCTCTCT R: GAATCATCAGAATCTACATCGT			1970
AT4G33030 <sup>d</sup>	F: GATGGTGTCTTGTACTGCTTGT R: COAGAAAACAGTGGCATATTCTG			770-1340
AT1G73180 <sup>d</sup>	F: AACTCTGCCAGTGTCCAATATA R: AGAATGCCATTACCCAGTAGT	KX767948	KX767949	810-1000
AT2G31890A	F: AGATTGGAGGGAGCTACTTTATT R: CCTCCCTATACGTGCTGAAATCC			450-620
AT3G46220 <sup>d</sup>	F: CAATTGAGGAGTGAATGGTGCCT R: TCCATTCTGCTGAAAGCTTGT			330-570
AT2G05120 <sup>e,g</sup>	F: TGCTCAAAGCTCTGGTCTCATCAA R: CGAGGAAGAACACTGAAGCACTAG	KX767932	KX767933	370-570
AT1G73740 <sup>d</sup>	F: TTGATATTGGAGGTCTTGGG R: CACAGCTCTGAAACAAACAGAG			870-1230

TABLE 1. Continued.

Locus ID <sup>a</sup>		GenBank accession no. <sup>b</sup>	Amplicon length range among all taxa		
			<i>B. simaruba</i>	<i>C. grandifolia</i>	
AT4G31790 <sup>d</sup>	F: ATTTGGTTGTTGAGCCAAGAAA R: GTCCAAAATGACCATCTGGAGTT		1620–2180		X X X X X X
AT5G10460	F: TGGTCATCAATTAGCAATTCTCAGC R: GCTCTTCAAAATCTCCAACT		1320–1800	X X X X X X	X X X X
AT4G26980 <sup>d</sup>	F: CTGCTAGTGGGTTCTGAATTGG R: ACTTCTCAAGCATTGACAACCTCAT		940–1170	X X X X X X	X X X X
AT5G48790 <sup>d</sup>	F: GAGGATTGGTTACTGAAGAAAGG R: TCGGCACCTTAAATTGTGAATATG		680–1250	X X X X X X	X X X X
AT5G15680A	F: TTCTCATCAAAACATCTGGGCC R: GAGGAATTGCAATCAGATTCTGGTC	KX768001	560	X X X X X X	X X X X
AT3G04650	F: CAAATCGCTTCTGGATGTTCATCA R: CTGGGGCAGTGGATGTTCATTC	KX767962, KX767964	490–660	X X X X X X	X X X X
AT2G25570 <sup>d</sup>	F: GACAACACTCAAACATCACGCCAG R: GTCCCTCTCATGAGCCATAG	KX767965	660–1160	X X X X X X	X X X X
AT2G31040 <sup>d</sup>	F: AAGTACTGGGGAGAAAGAG R: CCAAGTGTGAGGATTGCACTTC		1230–1690	X X X X X X	X X X X
AT4G04955	F: GAACAGATACTGAGAACAGCCAG R: TGAGCTTTAGTCCCTGAAGG		440–1170	X X X X X X	X X X X
AT3G21540 <sup>c</sup>	F: GTTGCTATTAGCTGATGCCAAA R: AATGGTTCTTGTAGTACGATCCAA	KX767970, KX767972	730–1020	X X X X X X	X X X X
AT2G05170A <sup>c</sup>	F: GAAGGAAATGTTACAGGTGAGGA R: TGAGAAGAATGGGGAGCTTCTT	KX767973	650–880	X X X X X X	X X X X
AT2G28450 <sup>f</sup>	F: TTCTGAGATAATGCTTATGTCAGG R: CGCCAATTGTCAGTACCA		1330	X X X X X X	X X X X
AT3G07750 <sup>d</sup>	F: GCTATATTGTTGATTGAGCCCT R: TGGTTGCTCACGTTTAATGATC		940–1330	X X X X X X	X X X X
AT1G76450 <sup>d</sup>	F: CGTCGGCACAAAATTACAAGAATGG R: TCAAATTCCCCAAGTCCCAATC	KX767916, KX767918	1330–1570	X X X X X X	X X X X
AT3G10530 <sup>c,d</sup>	F: ATTCCCCATCAAATTCCACTCG R: ACTCATGGCATCCAGTACTAA	KX767919	700–1960	X X X X X X	X X X X
AT3G61620 <sup>c</sup>	F: TTAGTATTTGAGGCAAGACGG R: CCTAAAAGGTATGGTCAAGGTT		1330	X X X X X X	X X X X
AT4G21170 <sup>d</sup>	F: TGGAGCTGTTATTATGCCCTTGT R: TAGTCCTAGTAAACAAACAGC	KX767992	730–1130	X X X X X X	X X X X
AT3G22990 <sup>d</sup>	F: TCTCCCTTCACTGAACTGAGA R: ACAGTTCAAGGGCACATGATC		900–1130	X X X X X X	X X X X
AT4G18810B	F: TTATGATATTGAGGCAAGACGG R: TCTAGGTCCATCTCAAGTGC	KX767988	540–640	X X X X X X	X X X X
AT1G77550B <sup>d</sup>	F: TGGTGTGTTATGTGATTGAGTC R: CTGGTCCCTGAGAACATTGTC	KX767922	760–820	X X X X X X	X X X X
AT5G16690 <sup>d</sup>	F: TGCTCCCTGAGAACATTGTC R: TCAAAGAACGCGTAAACAGTA		760–820	X X X X X X	X X X X
AT4G00560	F: CTGCTATGTCATAAACGCTTCCC R: GTCCACCAACATCAACAGTA	KX767984 (5' only)	900–1180	X X X X X X	X X X X

TABLE 1. Continued.

Locus ID <sup>a</sup>		GenBank accession no. <sup>b</sup>			Amplicon length range among all taxa
		<i>B. simaruba</i>	<i>C. grandifolia</i>		
AT3G17170 <sup>d</sup>	F: GATGATGAACATTATTTCTTGAGGC R: TCTTGAACCTTCTCATTCACACTGC	KX767968	KX767969	360–840	630–900
AT3G14910 <sup>e</sup>	F: GGAGCTATTTATCAAAGTTGTGCC R: AAAGCAATAATACGACCAAAGAACTG			2	16
Total no. of primers amplified/taxon				47	68
				90	90
				53	71
				72	72
				26	54
				34	34

Note: INT = reverse primer is an internal primer for the locus.

<sup>a</sup>Primer originally developed by Gostel et al. (2015).

<sup>b</sup>GenBank accession numbers from loci used in phylogenetic analysis in Gostel et al. (2015). Some loci were only created for loci of *Bursera simaruba* and *Commiphora grandifolia* that were used in the phylogenetic analysis in Gostel et al. (2015). Some loci have two GenBank numbers for a species because sequence reads did not cover the full length of the locus. The first GenBank number corresponds to the read from the 5' end of the locus; the second GenBank number corresponds to the read from the 3' end of the locus.

<sup>c</sup>Universal Burseraceae primer (excluding *Aucoumea*).

<sup>d</sup>Primer for which high-fidelity TAQ increased amplification success for *Commiphora grandifolia*.

<sup>e</sup>Primer for which high-fidelity TAQ increased amplification success for *Bursera simaruba*.

<sup>f</sup>Primer for which high-fidelity TAQ increased amplification success for *Bursera simaruba* and *Commiphora grandifolia*.

<sup>g</sup>Universal Sapindales primer (excluding *Aucoumea*).  
<sup>h</sup>Faint double band observed.

TABLE 2. Number of primer pairs amplified of the 91 primer pairs tested for each of the 11 taxa.

Species tested (Order; Family)	Primer pairs amplified/tested (%)
<i>Arabidopsis thaliana</i> (Brassicaceae)	2/91 (0.02)
<i>Aucoumea klaineana</i> (Sapindales; Burseraceae)	16/91 (17)
<i>Beiselia mexicana</i> (Sapindales; Burseraceae)	47/91 (52)
<i>Boswellia neglecta</i> (Sapindales; Burseraceae)	68/91 (75)
<i>Bursera simaruba</i> (Sapindales; Burseraceae)	90/91 (99)
<i>Bursera tonkinensis</i> (Sapindales; Burseraceae)	53/91 (58)
<i>Canarium pilosum</i> (Sapindales; Burseraceae)	71/91 (78)
<i>Commiphora grandifolia</i> (Sapindales; Burseraceae)	72/91 (79)
<i>Phellodendron amurense</i> (Sapindales; Rutaceae)	26/91 (28)
<i>Protium guianense</i> (Sapindales; Burseraceae)	54/91 (59)
<i>Schinus fasciculatus</i> (Sapindales; Anacardiaceae)	34/91 (37)

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APPENDIX 1. Accession information for taxa used in this study, including voucher information, country of origin, and latitude and longitude coordinate data, if available, and DNA extraction method.

Species	Voucher (Herbarium)	Country of origin	Geographic coordinates	DNA extraction method <sup>a</sup>
<b>Sapindales</b>				
Burseraceae				
<i>Aucoumea klaineana</i> Pierre	<i>Walters et al.</i> 466 (MO) <i>McPherson</i> 16293 (MO)	Gabon	00°07'12"S, 11°42'57"E 00°27'S, 11°45'E	1 1
<i>Beiselia mexicana</i> Forman	<i>Pell s.n.</i> (TEX)	Mexico	NA	1, 2
<i>Boswellia neglecta</i> S. Moore	<i>Weeks</i> 00-VII-29-1 (TEX)	Ethiopia	NA	2
<i>Bursera simaruba</i> (L.) Sarg.	<i>Weeks</i> 16-VI-16-01 (GMUF) <i>Goldman s.n.</i> (BH)	USA	NA NA	1 2
<i>Bursera tonkinensis</i> Guillamin	<i>Daly et al.</i> 13929 (NY)	Vietnam	20°15'12.6"N, 105°43'2.5"E	1
<i>Canarium pilosum</i> A. W. Benn.	<i>Bogler s.n.</i> (TEX)	Malaysia	NA	2
<i>Commiphora grandifolia</i> Engl.	<i>Gostel</i> 121 (GMUF)	Madagascar	23°39'19.64"S, 44°37'44.36"E	1
<i>Protium guianense</i> (Aubl.) Marchand	<i>Weeks</i> 10-I-09-10 (GMUF) <i>Miller and Hauk</i> 9391 (MO)	Madagascar Suriname	12°14'16.14"S, 49°22'12.906"E 04°45'22"N, 056°52'30"W	1 1
Anacardiaceae				
<i>Schinus fasciculatus</i> (Griseb.) I. M. Johnst.	<i>Silva-Luz</i> 287 (NY)	Argentina	24°52'05.4"S, 65°32'41.4"W	1
Rutaceae				
<i>Phellodendron amurense</i> Rupr.	<i>Weeks</i> 15-VII-13-01 (GMUF)	USA	38°49'53.76"N, 77°18'32.04"W	1
<b>Brassicales</b>				
Brassicaceae				
<i>Arabidopsis thaliana</i> (L.) Heynh.	<i>Gostel s.n.</i> (GMUF)	USA	NA	1

Note: NA = not available.

<sup>a</sup>1 = FastDNA, 2 = CTAB.