# **Evolutionary Comparisons of the Chloroplast Genome in** Lauraceae and Insights into Loss Events in the Magnoliids

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## Abstract

Available plastomes of the Lauraceae show similar structure and varied size, but there has been no systematic comparison across the family. In order to understand the variation in plastome size and structure in the Lauraceae and related families of magnoliids, we here compare 47 plastomes, 15 newly sequenced, from 27 representative genera. We reveal that the two shortest plastomes are in the parasitic Lauraceae genus *Cassytha*, with lengths of 114,623 (*C. filiformis*) and 114,963 bp (*C. capillaris*), and that they have lost NADH dehydrogenase (*ndh*) genes in the large single-copy region and one entire copy of the inverted repeat (IR) region. The plastomes of the core Lauraceae group, with lengths from 150,749 bp (*Nectandra angustifolia*) to 152,739 bp (*Actinodaphne trichocarpa*), have lost *trnl*-CAU, *rpl23*, *rpl2*, a fragment of *ycf2*, and their intergenic regions in IRb region, whereas the plastomes of the basal Lauraceae group, with lengths from 157,577 bp (*Eusideroxylon zwageri*) to 158,530 bp (*Beilschmiedia tungfangensis*), have lost *rpl2* in IRa region. The plastomes of *Caryodaphnopsis henryi* (Lauraceae) remain intact, as do those of the nonLaurales magnoliid genera *Piper*, *Liriodendron*, and *Magnolia*. On the basis of our phylogenetic analysis and structural comparisons, different loss events occurred in different lineages of the Laurales, and fragment loss events in the IR regions have largely driven the contraction of the plastome in the Lauraceae. These results provide new insights into the evolution of the Lauraceae as well as the magnoliids as a whole. **Key words:** Lauraceae, chloroplast, genome, phylogenetic relationship, loss event.

#### Introduction

In land plants, most chloroplast genomes are single, circular, double-stranded DNA sequences 100–220 kb in size, with a quadripartite structure including one large single-copy (LSC) region, one small single-copy (SSC) region, and a pair of inverted repeat (IR) regions (Bock 2007). Together these regions include >30 structural RNA genes and around 80 protein-coding genes, with the latter including genes related to photosynthesis, transcription or translation, and other functions (Gao et al. 2010). Generally, the ribosomal RNA genes are in the IR region, almost all of the photosynthesis related

genes in the LSC region, and a number of the NADPH dehydrogenase genes in the SSC region. The plastomes of land plants originated once, from a free-living algal ancestor (Turmel et al. 2006), but the gene contents and order vary considerably among species, and significant structural rearrangements and gene losses have been reported in several unrelated lineages, including ferns (Roper et al. 2007; Karol et al. 2010), gnetophytes (McCoy et al. 2008; Wu et al. 2009), and multiple angiosperm families (Goremykin et al. 2003a; Cai et al. 2006), as well as nonphotosynthetic plants (Wicke et al. 2016).

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Comparative analyses of the plastomes of algae and embryophytes show that four genes, tufA, ftsH, odpB, and rpl5, have been lost or transferred to the nucleus and three genes, matK, ycf1, and ycf2, have been gained in charophyte algae and embryophytes (Turmel et al. 2006). For example, the tufA gene, encoding chloroplast protein synthesis elongation factor Tu, is encoded in the plastomes of most algae, but is a pseudogene in *lsoetes*, fragmented in Anthoceros, cycads, and Gingko, and completely lost in the angiosperms (Karol et al. 2010). Within the angiosperms, three genes, ycf1, ycf2, and accD, have been lost in the Poaceae (Guisinger et al. 2010), whereas rpl22, infA, and accD were lost in the legumes, Lemnoideae, and Acoraceae, respectively (Wang and Messing 2011; Goremykin et al. 2005; Doyle et al. 1995). In plants with a heterotrophic lifestyle, pseudogenization and entire loss events of ndh-genes were detected (Wickett et al. 2008; Barrett et al. 2014; Wicke et al. 2016). However, the ndh-gene loss events have also occurred in autotrophic orchids, gnetophytes, and Pinaceae (Braukmann et al. 2009; Kim et al. 2015; Wakasugi et al. 1994).

In addition to gene losses, large inversions, and other structural rearrangements have been also reported. In ferns and seed plants, a 30-kb fragment flanked by the complete matK and rpoC2 has been identified as an inversion, with gene organization different from that in liverworts, mosses, hornworts, lycophytes, and Chaetosphaeridium (Wickett et al. 2011). In rice, maize, Calamus, and orchids, two identical trnH-rps19 gene clusters were detected as a duplication event before the diversification of extant monocot lineages (Chang et al. 2006; Wang et al. 2008; Luo et al. 2016). In Tetracentron and Trochodendron, a 4-kb extra region containing the five genes rpl22, rps3, rpl16, rpl14, and rps8 was found as evidence for unstable boundaries of the IR region across early-diverging eudicots (Sun et al. 2013, 2016). Interestingly, most of the rearrangements were detected in the boundary regions of IR, suggesting that the IR regions represent hotspots for structural rearrangements within the plastome (Wicke et al. 2011; Zhu et al. 2016).

The IR regions in the plastome of angiosperms have been used as evolutionary markers for elucidating relationships among some taxa, because they are frequently subject to contraction, expansion, or even complete loss (Lavin et al. 1990; Kim and Jansen 1994; Plunkett and Downie 2000; Luo et al. 2016; Sun et al. 2016; Zhu et al. 2016). In the early-diverging eudicots, the IR regions range from 24.3 to 36.4 kb in length and contain from 18 to 33 genes (Sun et al. 2016). In early-diverging monocots, the IR regions range from 25.2 to 33.3 kb in length and contain from 16 to 20 genes (Luo et al. 2016). As extreme examples, loss of one or two IR regions has been detected in Cephalotaxaceae (Yi et al. 2013), Pinaceae (Wu et al. 2011b), Taxodiaceae (Hirao et al. 2008), Leguminosae (Palmer et al. 1987; Lavin et al. 1990), Geraniaceae (Guisinger et al. 2011), and Cactaceae (Sanderson et al. 2015).

After the eudicots and monocots, the magnoliids is the third-largest group of Mesangiospermae, and includes four orders, 19 families, and over 9,000 woody species from all over the world (www.theplantlist.org). However, <30 species have assembled chloroplast genome sequences, and there has not been a systemic structural comparison of these plastomes. To improve understanding of the dynamics and evolution of plastome structure in magnoliids, we therefore focused on the plastomes of the important family Lauraceae and the related families Calycanthaceae (Laurales), Chloranthaceae (Chloranthales), Magnoliaceae (Magnoliales), Piperaceae (Piperales), and Winteraceae (Canellales). We included 15 newly sequenced and 33 previously reported plastomes in our study, representing 25 genera from all four orders of magnoliids. The main objectives of this study were 1) to reconstruct the phylogenetic relationships using the sequenced magnoliid plastomes, 2) to reveal plastome structural variations in Lauraceae, 3) to trace the evolutionary pattern of plastome contraction.

## **Materials and Methods**

#### Plant Material and Plastome Sequencing

Fresh leaves and silica-gel dried materials were sampled from 15 species representing 10 genera of Lauraceae. The voucher specimens for the 15 sampled plants collected from China and Indonesia were deposited at the Herbarium of Xishuangbanna Tropical Botanical Garden (HITBC), Chinese Academy of Sciences (CAS; table 1). Genomic DNA was extracted from 2 g leaves using the CTAB method (Doyle and Dickson 1987), in which 4% CTAB was used, and we added  $\sim 1\%$  polyvinyl polypyrrolidone (PVP) and 0.2% <sub>DL-</sub> dithiothreitol (DTT). From each purified sample of total DNA,  $0.5 \mu g$  was fragmented to construct short-insert (500 bp) libraries following the manufacturer's manual (Illumina) and then used for sequencing. The DNA samples were indexed by tags and pooled together in one lane of a Genome Analyzer (Illumina HiSeg 2000) for sequencing at BGI-Shenzhen, and >4.0 Gb of reads for each sample were obtained.

#### Genome Annotation and Comparison

The paired-end reads were filtered using GetOrganelle pipeline (https://github.com/Kinggerm/GetOrganelle) to get plastid-like reads, then the filtered reads were assembled using SPAdes version 3.10 (Bankevich et al. 2012). To retain pure chloroplast contigs, the final "fastg" files were filtered using the "slim" script of GetOrganelle. The filtered De Brujin graphs were viewed and edited using Bandage (Wick et al. 2015), then a circular chloroplast genome was generated. The genome was automatically annotated using CpGAVAS (Liu et al. 2012), then adjusted using Geneious version 9.1.7 (Kearse et al. 2012). The annotated chloroplast genomes

#### Table 1

Sampled Species of Lauraceae and Their Voucher Specimens Sequenced in This Study

No	Species	Herbarium	Taxon	Voucher	Geographic Origin	Accession Number in GenBank
1	Eusideroxylon zwageri	HITBC-BRG	Eusideroxylon zwageri Teijsm. & Binn.	SY34806	Sulawesi, Indonesia	MF939351
2	Cryptocarya chinensis	HITBC-BRG	Cryptocarya chinensis (Hance) Hemsl.	SY34239	Jianfenglin, Hainan	MF939349
3	Cryptocarya hainanensis	HITBC-BRG	Cryptocarya hainanensis Merr.	SY01426	Menghai, Yunnan	MF939350
4	Beilschmiedia tungfangensis	HITBC-BRG	Beilschmiedia tungfangensis S.K. Lee & L.F. Lau	SY34805	Wenshan, Yunnan	MF939348
5	Beilschmiedia pauciflora	HITBC-BRG	Beilschmiedia pauciflora H.W. Li	SY01364	Mengla, Yunnan	MF939347
6	Cassytha filiformis	HITBC-BRG	Cassytha filiformis Linnaeus	SY34802	Menghai, Yunnan	MF939337
7	Cassytha capillaris	HITBC-BRG	Cassytha capillaris Meisn.	SY34803	Sulawesi, Indonesia	MF939338
8	Neocinnamomum caudatum	HITBC-BRG	Neocinnamomum caudatum (Nees) Merr.	SY01561	Puer, Yunnan	MF939344
9	Neocinnamomum lecomtei	HITBC-BRG	Neocinnamomum lecomtei H. Liu	SY33249	Wenshan, Yunnan	MF939345
10	Caryodaphnopsis henryi	HITBC-BRG	Caryodaphnopsis henryi Airy Shaw	SY01542	Honghe, Yunnan	MF939346
11	Caryodaphnopsis malipoensis	HITBC-BRG	Caryodaphnopsis malipoensis Bing Liu & Y. Yang	SY32618	Wenshan, Yunnan	MF939343
12	Actinodaphne trichocarpa	HITBC-BRG	Actinodaphne trichocarpa C.K. Allen	SY32938	Emei, Sichuan	MF939342
13	Neolitsea sericea	HITBC-BRG	Neolitsea sericea (Blume) koidzumi	SY33307	Linan, Zhejiang	MF939341
14	Nectandra angustifolia	HITBC-BRG	Nectandra angustifolia (Schrad.) Nees & Mart.	SY34804	Sulawesi, Indonesia	MF939340
15	Sassafras tzumu	HITBC-BRG	Sassafras tzumu (Hemsl.) Hemsl.	SY34790	Anqing, Anhui	MF939339

have been submitted to GenBank (accession number: MF939337 to MF939351). The genome maps of all the 15 plastomes were drawn by OrganellarGenomeDRAW tool (OGDRAW; Lohse et al. 2013) and the gene organization maps were drawn by Gene Structure Display Server (GSDS) version 2.0 (Hu et al. 2015). Mauve version 2.4.0 software was used for alignment and determining the plastome rearrangements among the Magnoliids (Darling et al. 2004).

#### Phylogenetic Analysis

To estimate phylogenetic relationships within the magnoliids, 47 taxa with available complete plastomes were compared, including one taxon each from Canellales and Chloranthales, four from Piperales, six from Magnoliales, and 35 from Laurales. The 35 taxa included the 15 new plastomes and 20 complete plastomes which have been published elsewhere or adopted from NCBI (Song et al. 2015, 2016; Wu et al. 2017). Amborella trichopoda (AJ506156) was treated as the outgroup. For the species tree, maximum likelihood (ML) analyses were performed on data sets of 48 plastome sequences with single IR, SSC, and LSC regions. The whole genome matrix was aligned using MAFFT version 3.73 (Katoh and Standley 2013), then manually edited using Geneious version 9.1.7 (Kearse et al. 2012). ML analysis was conducted using RAxML version 7.2.6 with the GTR + G model to search the best-scoring ML tree (Tamura et al. 2011). One thousand bootstrap replicates were performed to obtain the confidence support. Bayesian inference (BI) was performed using MrBayes version 3.2.6 (Ronquist and Huelsenbeck 2003). The best-fit DNA substitution model of the Bayesian information criterion (BIC) was evaluated by using jModeltest version 2.1.10 (Darriba et al. 2012; Guindon et al. 2003). Markov Chain Monte Carlo (MCMC) analyses were run in MrBayes for 10,000,000 generations. The BI analysis started with a random tree and sampled every 1,000 generations. The first 25% of the trees was discarded as burn-in, and the remaining trees were used to generate a majority-rule consensus tree (supplementary fig. S1, Supplementary Material online). The trees were viewed and edited with the Fig tree version 1.4.0 software (http://tree.bio.ed.ac.uk/software/figtree/).

## **Results**

### Overall Structure and Gene Pool

Thirteen of the 15 newly sequenced Lauraceae plastomes displayed the typical quadripartite structure of angiosperms, including LSC, SSC, and a pair of IR regions, whereas the two plastomes from *Cassytha*, a genus of parasitic vines, have lost one copy of the IR (fig. 1). The complete plastome of *Cassytha filiformis* is 114,623 bp in length, 340 bp shorter than that of *Cassytha capillaris* (114,963 bp), and 42,954 bp shorter than that of *Eusideroxylon zwageri* (157,577 bp; table 2). Among the other 13 plastomes, genome size ranged from 150,749 bp (*Nectandra angustifolia*) to 158,530 bp (*Beilschmiedia tungfangensis*). In the LSC region, the length varied from 86,035 (*Caryodaphnopsis henryi*) to 93,803 bp (*Neolitsea sericea*), in the SSC region from 15,751 bp (*Caryodaphnopsis malipoensis*) to 19,222 bp (*Cryptocarya chinensis*), and in the IR region from 19,292 (*N. angustifolia*) to



Fig. 1.—Gene maps of the plastomes of Cassytha, Eusideroxylon, Cryptocarya, Beilschmiedia, Caryodaphnopsis, Neocinnamomum, Nectandra, Sassafras, Neolitsea, and Actinodaphne in the Lauraceae. The pink asterisks indicate the structural differences of IR loss.

25,601 bp (C. henryi). The plastomes of Eusideroxylon, Cryptocarya, Beilschmiedia, and C. henryi shared identical complements of coding genes; a total of 130 genes, including 8 rRNA genes, 37 tRNA, and 85 protein-coding genes, of which 17 are duplicated in IR regions. A total of 128 genes were detected on the plastomes of Neocinnamomum, Nectandra, Sassafras, Actinodaphne, Neolitsea, and Carvodaphnopsis malipoensis, 113 of which are single copy, while 15 are duplicated in IR regions. The different gene numbers reflect the duplication of rpl23 and trnl-CAU in the first group. The plastomes of Cassytha have not only lost the duplicated genes in the IR region, but also six NADH dehydrogenase (ndh) genes, ndhA, ndhC, ndhG, ndhI, ndhJ, and ndhK, and their five ndh genes are pseudogenes.

## Phylogenomic Analysis

The matrix of complete plastomes was used to reconstruct a phylogenetic tree of magnoliids (fig. 2). Magnoliids are divided into five main clades (ML-BS = 100%) corresponding to five orders: Canellales, Chloranthales, Laurales, Magnoliales, and Piperales. Sisterhood of Laurales and Magnoliales, with Piperales and Canellales being the next sister groups, was highly supported. Two major clades, including

Calycanthaceae and Lauraceae, were recognized within the Laurales. There was 100% support for the monophyly of Lauraceae family. Five well-supported groups were recovered within the Lauraceae (ML-BS = 100%). The basal group (ML-BS = 100%), including the genera Eusideroxylon, Cryptocarya, Beilschmiedia, and Endiandra, the Cassytha group (ML-BS = 100%), the Neocinnamomum group (ML-BS = 100%), the Carvodaphnopsis group (ML-BS = 100%), and the core group (ML-BS = 100%), including Alseodaphne, Persea, Phoebe, Machilus, Lindera, Laurus, Actinodaphne, Neolitsea, Litsea, Nectandra, Sassafras, and Cinnamomum.

#### Plastome Comparisons

Synteny and rearrangements were detected in ten plastomes of Lauraceae. A significant degree of synteny was found within the basal group, including *E. zwageri* and *B. tungfangensis*, and the core group, including *N. angustifolia*, *Laurus nobilis*, *Lindera communis*, *Machilus balansae*, *Alseodaphne semecarpifolia*, *Neocinnamomum caudatum*, and C. *capillaris*. However, the two groups differ in the orientation of a 13.7-kb fragment flanked by *rps7* and *rpl2* (fig. 3). In the basal group, the *rps7-ndhB-trnL-ycf2-trnl-rpl23-rpl2* segment has been

#### Table 2

Summary of 15 Complete Plastomes of Lauraceae

	Eusideroxylon zwageri	Cryptocarya chinensis	Cryptocarya hainanensis	Beilschmiedia tungfangensis	Beilschmiedia pauciflora	Cassytha filiformis	Cassytha capillaris
Total cpDNA size (bp)	157,577	157,675	157,145	158,530	157,901	114,623	114,963
Length of LSC region (bp)	89,231	89,199	89,002	89,351	88,673	-	-
Length of IR region (bp)	24,717	24,627	24,621	25,473	25,496	-	-
Length of SSC region (bp)	18,912	19,222	18,901	18,233	18,236	-	-
Total GC content	39.10%	39.10%	39.10%	39.00%	39.00%	36.90%	36.90%
Total number of genes (unique)	130 (113)	130 (113)	130 (113)	130 (113)	130 (113)	107 (107)	107 (107)
protein encoding	85	85	85	85	85	73	73
tRNA	37	37	37	37	37	30	30
rRNA	8	8	8	8	8	4	4
Length of <i>ycf1</i> (bp)	5,493	5,460	5,436	5,436	5,460	5,211	5,211
Length of truncated <i>ycf1</i> (bp)	971	977	974	1,863	1,863	-	-
Length of <i>ycf2</i> (bp)	6,882	6,885	6,885	6,843	6,849	5,583	5,583
Length of complete or truncated ycf2(bp)	6,882	6,885	6,885	6,843	6,849	_	-

Neocinnamomum	Neocinnamomum	Caryodaphnopsis	Caryodaphnopsis	Actinodaphne	Neolitsea	Nectandra	Sassafras
caudatum	lecomtei	henryi	malipoensis	trichocarpa	sericea	angustifolia	tzumu
150,842	150,838	154,938	149,239	152,739	152,442	150,749	151,798
91,881	91,912	86,035	91,901	93,783	93,803	93,783	92,752
20,257	20,257	25,601	20,036	20,078	20,067	19,292	20,096
18,447	18,412	17,701	17,266	18,800	18,505	18,382	18,854
38.80%	38.80%	39.00%	39.00%	39.20%	39.20%	39.20%	39.20%
128 (113)	128 (113)	131 (113)	128 (113)	128 (113)	128 (113)	128 (113)	128 (113)
84	84	86	84	84	84	84	84
36	36	37	36	36	36	36	36
8	8	8	8	8	8	8	8
5,517	5,517	5,526	5,526	5,574	5,568	5,535	5,586
928	928	1,473	1,473	1,378	1,372	1,372	1,419
6,831	6,831	6,894	6,894	6,876	6,846	6,909	6,294
3,110	3,110	6,894	3,186	3,168	3,162	2,478	3,168

combined with *tmH*-GUG, whereas the segment of the core group species has been combined with *rps19* (fig. 4), indicating that a rearrangement event occurred in Lauraceae plastome evolution. In the plastomes of *C. henryi* and in the basal group species, two unbroken protein-coding copies of *ycf2* were detected, suggesting that fragmentation of *ycf2* has occurred in other species of Lauraceae. Moreover, upstream of *rps19* adjoining the IR region, we detected one copy of a protein-coding gene *rpl23* and a tRNA gene *trnM*-CAU in the plastome of *C. henryi* and the basal group species, but not in the plastomes of other species, indicating that significant IR boundary changes occurred in Lauraceae plastome evolution.

# IR Expansion and Contraction

In the sequenced plastomes of Lauraceae, two complete or fragmented copies of *ycf1* and *ycf2* were located at the

boundaries between the IR regions and the LSC or SSC regions. The full lengths of ycf2 and ycf1 ranged from 5,583 bp in Cassytha filiformis to 6,894 bp in Carvodaphnopsis malipoensis and from 5,211 bp in Cassytha filiformis to 5,586 bp in Sassafras tzumu, respectively (table 2). Double complete copies of the ycf2 genes were detected in the seven sequenced Lauraceae plastomes of the basal group species, but only one complete copy and one fragment in the 24 plastomes of C. malipoensis, Neocinnamomum, and the core group species, except those of C. henryi and both Cassytha species. The length of the fragment of ycf2 ranged from 2,478 bp in N. angustifolia to 3,168 bp in Actinodaphne trichocarpa. In contrast, all 32 sequenced Lauraceae plastomes, except the two species of Cassytha, had one complete copy and a fragment of ycf1. The length of the fragment of ycf1 ranged from 971 bp in E. zwageri to 1,863 bp in Beilschmiedia pauciflora. Neither Cassytha plastome had fragments of ycf1 and ycf2, but only one complete copy of each due to the IR loss.



Fig. 2.—Molecular phylogenetic tree of 47 taxa of Magnoliids based on complete plastome sequences using unpartitioned ML. Numbers at each node are bootstrap support value.

## Discussion

#### Relationships in Lauraceae

This study included 47 complete chloroplast genomes for plants from all five orders (Canellales, Chloranthales, Laurales, Magnoliales, and Piperales) of the magnoliids. All of these complete plastome sequences of Lauraceae and related families yielded a fully resolved tree, consistent with the Angiosperm Phylogeny Group's most recent phylogeny, APG IV (Byng et al. 2016). Relationships among the five orders of the magnoliids are clarified as sisterhood of Laurales and Magnoliales, with Piperales and Canellales being the next sister groups, and Chloranthales the most basal group. Calycanthaceae and Lauraceae were recognized within the Laurales. All of these clades were recognized by Renner (Renner 1999).

The deep relationships of 34 Lauraceae taxa are separated into the following groups in our study. *Eusideroxylon*, *Cryptocarya*, *Beilschmiedia*, and *Endiandra* form the first

	LSC	*1	*2 IRa	SSC	IRb
5000 10000 15000 20000 25000 30000 35000 40000 4	45000 50000 55000 60000 65000 7000	0 75000 80000 85000 90000		5000 120000 125000 130000 135000 1	40000 145000 1500
Nectandra_angustifolia	···· <sup>·</sup> ○ <sup>□</sup> œ <b>d</b> ⊂⊂⊂ <sup>©</sup> œ <b>v</b> <sup>1</sup> !!	ber the transmission			
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Lindera_communis					
5000 10000 15000 20000 25000 30000 35000 40000	45000 50000 55000 60000 65000 7000			5000 120000 125000 130000 135000 1	40000 145000 150000
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Alseodaphne_semecarpifolia 5000 10000 15000 20000 25000 30000 35000 40000	45000 50000 55000 80000 85000 7000	0 75000 80000 85000 90000	95000 100000 105000 110000 11	5000 120000 125000 130000 135000 1	
Caryodaphnopsis_henryi					
5000 10000 15000 20000 25000 30000 35000 40000 4	45000 50000 55000 60000 65000 7000		disaal a state	5000 120000 125000 130000 135000 1	40000 145000 50000
5000 10000 15000 20000 25000 30000 35000 40000 4	45000 50000 55000 60000 65000 7000	0 75000 80000 85000 90000	95000 106000 105000 110000	-	
Cassytha_capillaris	45000 50000 55000 80000 85000 7000	0 75000 80000 85000 90000	95000 100000 105000 110000 11	5000 120000 125000 130000 135000 1	40000 145000 15000 155000
	45000 50000 55000 60000 65000 7000		95000 100000 105000 110000 11	5000 120000 125000 130000 135000 1	40000 145000 150000 155000
	ישים מים בים מישים שישים איי				

Eusideroxylon\_zwageri

Fig. 3.—Synteny and rearrangements detected in Lauraceae plastomes using the Mauve multiple-genome alignment program. The yellow asterisks 1 indicate the varied gene locus of *rpl2*, the blue asterisks 2 indicate a rearrangement of the fragment flanked by *rpl2* and *rps7*, and the green asterisks 3 indicate an insert of intergenic region *rpl2–rpl23*.



Fig. 4.—Comparisons of gene loci in the fragments flanked by rps19 and trnL in IRa and trnL and trnH in IRb among 15 plastomes of Magnoliids.

group in the phylogeny. *Cassytha*, *Neocinnamomum*, and *Caryodaphnopsis* form the second, third, and fourth groups, respectively. The fifth group includes *Alseodaphne*, *Persea*, *Phoebe*, and *Machilus*. The sixth group includes *Nectandra*, *Sassafras*, and *Cinnamomum*. And the last group includes

*Lindera, Laurus, Litsea, Actinodaphne, and Neolitsea.* The phylogenetic placements of the first, fourth, fifth, and sixth groups are consistent with previously published phylogenetic relationships (Chanderbali et al. 2001; Rohwer and Rudolph 2005). The position of *Cassytha*, considered as a 'jumping

genus' by Rohwer and Rudolph (2005), was settled here in the way predicted from morphology (Chanderbali et al. 2001). The seventh group, equivalent to the tribe Laureae (Chanderbali et al. 2001), was confirmed as sister to the sixth group, tribe Cinnamomeae (including *Sassafras*), which has always been assumed based on morphological characters, although previous molecular analyses failed to prove it convincingly (Chanderbali et al. 2001; Rohwer and Rudolph 2005).

#### Unusual Structure of the Cassytha Plastomes

The sizes of the fifteen newly sequenced Lauraceae plastomes differed greatly, from 114,623 bp in the hemiparasitic vine, C. capillaris, to 158,530 bp in B. tungfangensis, as a result of the loss of one IR copy and six ndh genes in Cassytha. Cassytha is the only stem hemiparasitic genus with reduced leaves and roots in the magnoliids, and the only nonwoody member of the Lauraceae. We show that it is also unique in the Lauraceae the loss of one IR copy in its plastome, although similar losses have occurred independently in the Leguminosae (Cai et al. 2008). Pinaceae (Raubeson and Jansen 1992). Cephalotaxaceae (Yi et al. 2013), and cupressophytes (Wu et al. 2011a). In addition, six ndh genes, ndhA, ndhC, ndhG, ndhI, ndhJ, and ndhK, have been lost, and the other five, ndhB, ndhD, ndhE, ndhF, and ndhH, are clearly pseudogenes in both Cassytha taxa sequenced in this study. All eleven *ndh* genes encode independent subunits of a plastid NADPH-dehydrogenase complex (Ndh 1-complex) which carries out one of the recycled electron pathways around Photosystem I (Casano et al. 2000). Cyclic electron flow is vital for maintenance of efficient photosynthesis and enablement of photoprotection under environmental stresses in higher plants (Wang et al. 2006). The ndh genes are frequently pseudogenized or lost in plant groups with a degree of heterotrophy, such as Aneura, Cuscuta, Epifagus, Hydnora, and nonphotosynthetic orchid species, and in some autotrophic gymnosperms and ferns (dePamphilis and Palmer 1990; Wicke et al. 2011; Wickett et al. 2008; McNeal et al. 2007; Kim et al. 2015; Naumann et al. 2016), but this is first report for Cassytha, the only hemiparasitic genus in the Laurales. This adds to the evidence that the Ndh1-complex is not essential for plant survival, while the ndh-independent antimycin-Asensitive pathway, which functions in cyclic electron flow as another choice, could be more important under most conditions (Shikanai 2014).

#### Loss Events in the Laurales

Comparative genomic analysis indicated that missing segments of DNA in Lauraceae plastids mainly drive the genome contraction events. A fragment flanked by *rps7* and *rpl2* was detected as a rearrangement event between the basal group species and the other species except *C. henryi*. However, it looks more like two or more independent loss events when we choose the plastomes of C. henryi or nonLaurales species as reference. Double IR fragments with the gene order of trnLycf2-trnl-rpl23-rpl2 are highly conserved in the plastomes of C. henryi (fig. 4) and nonLaurales genera such as Drimys, Piper, Liriodendron, and Magnolia (Cai et al. 2006; Zhu et al. 2016; Yang et al. 2014), indicating the plastome of C. *henryi* is evolutionarily conserved. In *Calycanthus* (Laurales) plastome (Goremykin et al. 2003a), one copy of rpl2 with the length of 1,480 bp disappeared from the trnL-rpl2 fragment in IRb, but all of the sequenced Lauraceae plastomes of the basal group, including Endiandra, Beilschmiedia, Cryptocarya, and Eusideroxylon, lost another copy of rpl2 from the *trnL-rpl2* fragment in the IRa region (fig. 4). More interesting are the sequenced Lauraceae plastomes of the core group, including Alseodaphne, Persea (Song et al. 2016), Phoebe, Machilus (Song et al. 2015), Lindera, Laurus, Litsea, Nectandra, Sassafras, Cinnamomum (Wu et al. 2017), Actinodaphne, and Neolitsea, which have further lost a segment of at least 4,500 bp which contains a fragment of ycf2 and one copy of rpl23 and trnl-CAU in IRb of Calycanthus. This segment was also lost in the plastomes of Neocinnamomum species and C. malipoensis. Taken together, these independent loss events show that in the Lauraceae the plastomes of Neocinnamomum, Cassytha, the core group, and the basal group could share a common ancestral genome structure like that of C. henryi, but have subsequently evolved independently with different loss patterns.

#### Evolutionary Pattern in Angiosperms

To put these results in a wider phylogenetic context, we traced the fragments flanked by trnL-CAA and rps19 in the IRa region and by trnL-CAA and trnH-GUG in the IRb region in the six major groups of the angiosperms and found that the gene backbone and order are conserved (fig. 5). In the earlydiverging angiosperm species, A. trichopoda and Nymphaea alba, of the ANITA group (Qiu et al. 1999), the gene orders of the fragments are rps19-rpl2-rpl23-trnl-ycf2-trnL and trnLycf2-trnl-rpl23-rpl2-trnH (Goremykin et al. 2003b, 2004). These orders are retained in the early diverging monocot Tofieldia thibetica (Luo et al. 2016) and Ceratophyllum demersum in the Ceratophyllaceae (Moore et al. 2007). In the early diverging eudicot Euptelea pleiosperma (Sun et al. 2016), the only change in the gene order is a new insertion of a fragment of rps19. In the magnoliids, the same gene order for both fragments is retained in the sequenced species of Choranthaceae (Hansen et al. 2007), Piperales (Cai et al. 2006), and Magnoliales (Zhu et al. 2016), but a new copy of trnH has been inserted between rps19 and rpl2 in the IRa fragment of Drimys granadensis in the Canellales (Cai et al. 2006) and the copy of *rpl2* has been lost between *rps19* and rps23 in the IRa region of Endiandra, Beilschmiedia, Cryptocarya, and Eusideroxylon species in Lauraceae, and in



Fig. 5.—Model of the origin and variation of the fragments flanked by *rps19* and *trnL* in IRa and *trnL* and *trnH* in IRb among plastomes of angiosperms. The pink asterisks indicate the varied gene loci.

IRb of *Calycanthus* in Calycanthaceae (Goremykin et al. 2003a). Nevertheless, our comparative genomic analysis concluded that the regions encompassing the *ycf2* and the adjoined *trnH*-GUG or *trnL*-CAA gene in the plastomes of *C. henryi* and other early-diverging angiosperms are the retained IRs, corresponding to either IRa or IRb in the basal and core groups of Lauraceae.

## **Supplementary Material**

Supplementary data are available at *Genome Biology and Evolution* online.

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