

Phylotranscriptomics of Theaceae: generic-level relationships, reticulation and whole-genome duplication

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• **Background and Aims** Theaceae, with three tribes, nine genera and more than 200 species, are of great economic and ecological importance. Recent phylogenetic analyses based on plastomic data resolved the relationships among the three tribes and the intergeneric relationships within two of those tribes. However, generic-level relationships within the largest tribe, Theeae, were not fully resolved. The role of putative whole-genome duplication (WGD) events in the family and possible hybridization events among genera within Theeae also remain to be tested further.

• **Methods** Transcriptomes or low-depth whole-genome sequencing of 57 species of Theaceae, as well as additional plastome sequence data, were generated. Using a dataset of low-copy nuclear genes, we reconstructed phylogenetic relationships using concatenated, species tree and phylogenetic network approaches. We further conducted molecular dating analyses and inferred possible WGD events by examining the distribution of the number of synonymous substitutions per synonymous site (*Ks*) for paralogues in each species. For plastid protein-coding sequences, phylogenies were reconstructed for comparison with the results obtained from analysis of the nuclear dataset.

• **Results** Based on the 610 low-copy nuclear genes (858 606 bp in length) investigated, Stewartieae was resolved as sister to the other two tribes. Within Theeae, the *Apterosperma–Laplacea* clade grouped with *Pyrenaria*, leaving *Camellia* and *Polyspora* as sister. The estimated ages within Theaceae were largely consistent with previous studies based mainly on plastome data. Two reticulation events within *Camellia* and one between the common ancestor of *Gordonia* and *Schima* were found. All members of the tea family shared two WGD events, an older At- γ and a recent Ad- β ; both events were also shared with the outgroups (Diapensiaceae, Pentaphylacaceae, Styracaceae and Symplocaceae).

• **Conclusions** Our analyses using low-copy nuclear genes improved understanding of phylogenetic relationships at the tribal and generic levels previously proposed based on plastome data, but the phylogenetic position of the *Apterosperma–Laplacea* clade needs more attention. There is no evidence for extensive intergeneric hybridization within Theeae or for a Theaceae-specific WGD event. Land bridges (e.g. the Bering land bridge) during the Late Oligocene may have permitted the intercontinental plant movements that facilitated the putative ancient introgression between the common ancestor of *Gordonia* and *Schima*.

Key words: Theaceae, phylogeny, transcriptome, low-copy nuclear genes, molecular dating, phylogenetic network, whole-genome duplication.

INTRODUCTION

Theaceae, the tea family, comprise nine genera in three tribes and contain 372 accepted species (WFO, 2021) of evergreen and deciduous trees and shrubs. Members of Theaceae have great economic and ecological importance; the family contains familiar plants such as tea [e.g. *Camellia sinensis* (L.) Kuntze], oil plants (e.g. *C. oleifera* Abel) and a number of woody ornamentals (e.g. *C. japonica* L., *C. reticulata* Lindl.). Some large tree representatives (e.g. *Schima*) and small tree lineages (e.g. *Camellia, Stewartia*) are dominant or common species of the subtropical evergreen broadleaved forests in East Asia (Tang, 2015). Due to excessive collection and habitat destruction, several species (mainly species of *Camellia*) have been listed as (critically) endangered, including *C. fangchengensis* S. Ye Liang & Y. C. Zhong, *C. hekouensis* C. J. Wang & G. S.

Fan and *C. piquetiana* (Pierre) Sealy (IUCN, 2020). In addition, many new species (Orel, 2006; Orel and Wilson, 2010*b*, 2012; Orel *et al.*, 2013; Orel and Curry, 2015, 2019; Lee and Yang, 2019; Liu *et al.*, 2019, 2020*a*, *b*; Yu *et al.*, 2021) and subgeneric taxa (Orel and Wilson, 2010*a*; Orel *et al.*, 2014) have been described and published for *Camellia* in the past decade.

Since the establishment of Theaceae (Mirbel, 1813), the systematic boundaries of genera in the family as defined by morphology have changed significantly (Bentham and Hooker, 1862; Melchior, 1925; Takhtajan, 1997), from two genera to as many as six tribes and 32 genera. Molecular systematic studies recognized three tribes and nine genera (Stevens, 2001 onwards; APG IV, 2016). Since then, a number of systematic studies using DNA markers, morphology, anatomy and cytology have been conducted to explore the relationships among tribes and genera (Ye, 1990; Tsou, 1998; Prince and Parks, 2001; S. X. Yang *et al.*, 2004; Wang *et al.*, 2006; J. B. Yang *et al.*, 2006; M. M. Li *et al.*, 2013; W. Zhang *et al.*, 2014; Yu *et al.*, 2017b). However, many phylogenetic relationships remain unresolved in this family.

First, phylogenetic relationships among the three tribes in Theaceae remain to be confirmed using nuclear genes. Evidence based on floral development indicated a sister relationship between tribes Gordonieae and Stewartieae (Tsou, 1998; treated as subtribes in this study). The same topology was recovered based on the small single-copy region (SSC) of the plastome, but with conflicting support among maximum parsimony (MP), maximum likelihood (ML) and Bayesian inference (BI) analyses (M. M. Li et al., 2013). However, phylogenetic analysis of 46 morphological characters supported a sister relationship between Gordonieae (=Schimeae) and Theeae (Wang et al., 2006), which was also supported by phylogenetics of plastid rbcL, matK and trnL-F, mitochondrial matR and nuclear ribosomal internal transcribed spacer (nrITS) (Prince and Parks, 2001; S. X. Yang et al., 2004; J. B. Yang et al., 2006) and whole plastome sequences (Yu et al., 2017b). In a recent study of Ericales based on 25 loci from the plastid, nuclear and mitochondrial genomes, Theeae and Gordonieae were also recovered as sisters, with Stewartieae as their sister (Rose et al., 2018). However, the sampling of nuclear genes in previous studies was very limited, and further analyses that incorporate more regions of the nuclear genome are needed.

Intergeneric relationships within Theeae have been challenging to resolve, especially the phylogenetic position of Apterosperma and Laplacea. Since the time of its original description (Chang, 1976), Apterosperma has been formally placed in tribe Gordonieae (=Schimeae) and considered close to Schima and Franklinia based on similar morphological characters (Ye, 1990; Chang and Ren, 1998; Tsou, 1998). A combined molecular phylogenetic analysis based on nrITS, plastid trnL-F and mitochondrial DNA (mtDNA) matR sequence data placed Apterosperma as sister to all other genera in Theeae (S. X. Yang et al., 2004). However, based on analyses of 46 morphological characters. Apterosperma was placed as sister to other Theeae (Wang et al., 2006). Using five genomic regions (plastid: *atpI-H*, *matK*, *psbA5'R-ALS-11F*, *rbcL*; nuclear: LEAFY) and 30 species representing four of the five genera within Theeae, W. Zhang et al. (2014) found that Apterosperma formed a sister relationship with *Polyspora* in the chloroplast DNA (cpDNA) tree, but was placed within a clade comprising

Tutcheria (=*Pyrenaria*) and *Parapyrenaria* (=*Pyrenaria*) in the *LEAFY* tree. In that study, *Camellia* and *Pyrenaria* were not monophyletic, and inconsistent phylogenetic placement of some species between the nuclear and plastid trees was proposed to be the result of widespread hybridization among genera in Theeae.

In contrast to *Apterosperma*, very few studies have included *Laplacea*. Prince and Parks (2001) recovered *Laplacea* within a clade comprising *Camellia*, *Tutcheria* (=*Pyrenaria*) and *Glyptocarpa* (=*Camellia*). In Yu *et al.* (2017b), *Apterosperma* and *Laplacea* were sisters with strong support [ML bootstrap support (MLBS) = 96 %, BI posterior probability (PP) = 1.0]. However, the *Apterosperma–Laplacea* clade grouped either with *Camellia–Polyspora* or with *Pyrenaria* with moderate support based on different partitions of the plastome, grouped with *Polyspora* with weak support in the nrITS dataset, or even resolved as an early-branching clade from the combined plastome and nrITS dataset.

Species in the tea family are disjunctly distributed in temperate, subtropical and tropical areas of eastern to south-eastern Asia. and eastern North America to Central and South America, i.e. an Amphi-Pacific disjunction (Kobuski, 1949, 1950; Prince, 1993; Stevens, 2001 onwards; Ming and Bartholomew, 2007). Several studies have been conducted to investigate the biogeographical history of Theaceae, and some focused on the Eastern North American-East Asian disjunct genus Stewartia (Prince, 2002; H. Y. Lin et al., 2019). Others attempted to uncover the spatial-temporal history of the whole family. Based on two calibration points, the biogeographical reconstruction of M. M. Li et al. (2013) indicated that Theaceae originated in the late Cretaceous [~86 million years ago (Mya)] and started to diversify in the early Eocene (~49 Mya); species interchange of Gordonieae and Stewartieae between North America and eastern Asia may have been facilitated by the Bering land bridge. Our previous study also suggested a late Cretaceous origin (~91.6 Mya) of the family, but in contrast to M. M. Li et al. (2013), the crown of Theaceae was inferred as late Palaeocene (~57.3 Mya) (Yu et al., 2017b). Theaceae were proposed to be of Indo-Malaysian origin as part of a broader phylogenetic study of Ericales (Rose et al., 2018), although with relatively sparse sampling. However, a more recent study suggested a broad mid- to high-latitude Northern Hemispheric (e.g. Eurasia, Nearctic) origin of Theaceae (Yan et al., 2021). Biogeographical analyses based on a large number of nuclear genes and strong taxon sampling are needed to resolve this issue.

Ancient whole-genome duplication (WGD), an important evolutionary force in plants, has been reported in the common ancestor of extant seed plants, extant angiosperms and core eudicots (At- γ) (Jiao *et al.*, 2011; Vekemans *et al.*, 2012), as well as at many other critical nodes across the green plant tree of life (Leebens-Mack *et al.*, 2019). WGD events were also found in the early history of numerous families, such as Asteraceae, Brassicaceae, Fabaceae, Poaceae and Rosaceae (Barker *et al.*, 2008; Schranz *et al.*, 2012; C. H. Huang *et al.*, 2016; Xiang *et al.*, 2017; Qiao *et al.*, 2019). Based on the kiwifruit (*Actinidia chinensis*) genome, researchers proposed a WGD event called Ad- β , which was shared by *Actinidia* and *Camellia* (T. Shi *et al.*, 2010; S. X. Huang *et al.*, 2013). However, using genome collinearity and also the MAPS pipeline, Ad- β was recently mapped to the core Ericales (Leebens-Mack *et al.*, 2019), and more specifically to the clade comprising core Ericales, primuloids, polemonioids and Lecythidaceae (C. Zhang *et al.*, 2020). In Theaceae, whole-genome sequence data were only available for *Camellia*, and results for the placement of the WGD event from different studies were not in agreement with each other (Xia *et al.*, 2017, 2020; Wei *et al.*, 2018); more data and broader sampling are needed.

Phylotranscriptomics has become an important approach for plant phylogenetics (Soltis et al., 2013; Yang and Smith, 2013; Wickett et al., 2014; C. H. Huang et al., 2016; Yu et al., 2018; C. Zhang et al., 2020) and has been widely used to explore diverse evolutionary questions, including the origin and early diversification of land plants (Wickett et al., 2014), deep-level (among eight clades) angiosperm phylogeny (Zeng et al., 2014), phylogenetic relationships within eudicots, asterids and rosids, respectively (L. Zhao et al., 2016; Zeng et al., 2017; C. Zhang et al., 2020), and intergeneric relationships for several species-rich orders or families (Caryophyllales, Cupressaceae, Asteraceae, Brassicaceae, Fabaceae, Rosaceae and Pinaceae) (C. H. Huang et al., 2015, 2016; Y. Yang et al., 2015; Xiang et al., 2017; Ran et al., 2018; Mao et al., 2019; Y. Y. Zhao et al., 2021). Additionally, the 1000 Plant Transcriptomes Project (1KP), which sequenced transcriptomes from 1124 species representing the diversity of green plants, provided resolution across the green tree of life (Matasci et al., 2014; Leebens-Mack et al., 2019). Furthermore, comparing evidence from the plastid and nuclear genomes allows the detection of cytoplasmic introgression and other forms of hybridization (Calvo et al., 2013; Folk et al., 2017; Guo et al., 2018; Morales-Briones et al., 2018; Stubbs et al., 2020).

Here, transcriptomes of 55 species of Theaceae were sequenced, and low-depth whole-genome sequencing was conducted for another two species because no fresh tissue was available; both nuclear and plastid genes were extracted. Plastid genes were integrated with plastome data from our previous study (Yu *et al.*, 2017*a*, *b*). We aim to reconstruct the nuclear phylogenetic framework and temporal history of the tea family, focusing on the relationships among tribes and genera and the phylogenetic position of *Apterosperma* and *Laplacea*. Furthermore, we also test the intergeneric hybridization hypothesis proposed previously and investigate whether a previously detected WGD event in *Camellia* is shared by other genera of Theaceae.

MATERIALS AND METHODS

Taxon sampling, transcriptomics and low-depth whole-genome sequencing

We collected 58 samples representing 57 species from all three tribes and all nine genera of Theaceae (Table 1). Fresh and healthy leaves of 27 samples were collected in the field and then frozen immediately in liquid nitrogen. For 29 samples, we used leaf tissue collected in the field and stored in a -80 °C freezer. Additionally, silica-dried leaves of *Stewartia malacodendron* and *Laplacea fruticosa* were used for low-depth whole-genome sequencing (at least 30× coverage).

Total RNA was extracted from the 56 samples of flash frozen leaves using the Spectrum Plant Total RNA Extraction Kit (Sigma-Aldrich, Burlington, MA, USA). Total genomic DNA was isolated from silica-dried leaves of *Stewartia malacodendron* and *Laplacea fruticosa* using the modified CTAB method (Doyle and Doyle, 1987). RNA sequencing and low-depth whole-genome sequencing library construction, Illumina HiseqXten sequencing, raw data cleaning and quality control were performed at Novogene (China). Additionally, plastid genome data were obtained from our previous studies (Yu *et al.*, 2017*a*, *b*). Six species from Diapensiaceae, Pentaphylacaceae, Styracaceae and Symplocaceae were used as outgroups (Table 1). The transcriptome data of these six species were downloaded from GenBank.

Based on the results from previous phylogenetic studies of Theacaee, *Pyrenaria* and *Stewartia* were treated in the broad sense according to the treatment in the Flora of China (Ming and Bartholomew, 2007). In our previous study (Yu *et al.*, 2017*b*), we found that *Laplacea grandis* grouped with *Gordonia lasianthus* and should therefore be moved into *Gordonia*; thus, here we use the name *Gordonia brenesii* H. Keng (*=Laplacea grandis*) following Grayum and Madrigal (2011).

Sequence assembly and orthologue identification for nuclear genes

Quality control for all the raw sequencing reads was performed using Fastp v0.20.1 (S. F. Chen *et al.*, 2018), including removal of adapters, reads containing N and reads with low quality scores (percentage of base \leq Q20). Trinity v2.8.4 was used to conduct *de novo* assembly of cleaned Illumina RNA sequencing reads of each species (Grabherr *et al.*, 2011; Haas *et al.*, 2013). After transcripts had been filtered with Transrate v1.0.3 (Smith-Unna *et al.*, 2016) and clustered using Corset v1.07 (Davidson and Oshlack, 2014), TransDecoder v5.3.0 (Haas *et al.*, 2013) with blastp was utilized to choose open reading frames. CD-HIT v4.6 (Li and Godzik, 2006) was then used to remove redundant contigs with a threshold of 0.99. Orthology inference was conducted following Yang and Smith's (2014) pipeline, setting min_taxon as 48 in each orthologue group.

For the data obtained from low-depth whole-genome sequencing of two species (*Stewartia malacodendron* and *Laplacea fruticosa*), we first *de novo* assembled each genome using Platanus v1.2.4 with default parameters (Kajitani *et al.*, 2014). Then RepeatMasker v4.1.1 (Tarailo-Graovac and Chen, 2009) and RepeatModeler (http://www.repeatmasker. org/) were used to identify tandem repeats and transposable element (TEs). We carried out gene annotation using *de novo* gene prediction in AUGUSTUS (Haas *et al.*, 2008) and homologue prediction using Exonerate (Slater and Birney, 2005) and then generated an integrated gene set using EVidenceModeler (Haas *et al.*, 2008). Preliminary gene trees were reconstructed using RAxML v8.2.12 (Stamatakis, 2014).

To reduce potentially misidentified orthologues, we carefully examined the individual gene trees obtained for the 631 putative orthologues and found the three tribes within Theaceae were not monophyletic in 21 of these gene trees. Because all three tribes within Theaceae were consistently monophyletic in

Pyrenaria spectabilis var. spectabilis

YXQ155

Taxon Voucher Sources No. of reads SRA number Plastid specimen (trimmed) genome Stewartieae Stewartia calcicola YXQ090 Yunnan, China 76 314 960 SRR14596892 KY406783 KY406775 YXQ144 Guangxi, China 101 158 084 SRR14596891 Stewartia cordifolia Stewartia crassifolia YXQ171 Hunan, China 88 878 696 SRR14596880 KY406766 FLAS 260361 1 798 360 616 SRR14596869 KY406773 Stewartia malacodendron Alabama, USA SRR14596858 Stewartia ovata 18847*A The Arnold Arboretum 100 544 128 KY406782 Stewartia pseudocamellia MO-6587810 Missouri Botanical 83 044 916 SRR14596847 KY406786 Garden Stewartia pteropetiolata YXQ038 Yunnan, China 124 092 532 SRR14596838 KY406770 YX015072003 91 741 812 SRR14596837 KY406789 Stewartia rostrata Jiangxi, China 91 059 484 Stewartia rubiginosa YXQ189 Hunan, China SRR14596836 KY406777 YXQ15072001 91 325 600 SRR14596835 KY406748 Stewartia sinensis Jiangxi, China Gordonieae Franklinia alatamaha MO-6587811 Missouri Botanical 101 387 448 SRR14596890 KY406774 Garden N. Zamora 7196 <u>Gordonia brenesii</u> Guanacaste, Costa Rica 104 462 244 SRR14596889 KY406761 82 908 172 SRR14596888 Gordonia lasianthus JCRA 110687 JC Raulston Arboretum, KY406790 Raleigh, NC, USA Schima argentea YXQ226 Yunnan, China 86 065 752 SRR14596887 * YXQ072 Yunnan, China 89 409 700 SRR14596886 KY406784 Schima brevipedicellata YXQ034 Schima noronhae Yunnan, China 88 824 452 SRR14596885 KY406787 YX0053 97 108 488 SRR14596884 KY406779 Schima sericans Yunnan, China SRR14596883 YXQ142 91 704 880 KY406788 Schima superba Guangxi, China Schima wallichii YXQ001 Yunnan, China 89 876 764 SRR14596882 KY406795 Theeae Apterosperma oblata YangSX 5978 Guangdong, China 86 466 584 SRR14596881 YangSX 5010 Hainan, China 98 672 716 SRR14596879 Camellia amplexifolia _ YangSX 5540 98 123 144 SRR14596878 Camellia assimiloides Guangdong, China _ Camellia cordifolia YangSX 5551 Guangdong, China 96 616 940 SRR14596877 _ Camellia cuspidata YangSX 5118 Hubei, China 78 369 336 SRR14596876 _ Camellia flavida YangSX 5865 Guangxi, China 92 242 904 SRR14596875 _ Camellia fluviatilis YangSX 4033 Guangxi, China 68 780 488 SRR14596874 _ YangSX 6064 SRR14596873 Camellia grijsii Guizhou, China 120 959 328 _ Camellia gymnogyna YangSX 5953 Guangxi, China 94 768 804 SRR14596872 _ YangSX 5653 111 316 580 Guizhou, China SRR14596871 Camellia huana _ Camellia ilicifolia YangSX 5287 Guizhou, China 95 559 812 SRR14596870 YangSX 5926 SRR14596868 Camellia longipedicellata Guangxi, China 90 347 792 _ Camellia longissima YangSX 5079 Guangxi, China 99 050 172 SRR14596867 _ Camellia luteoflora YangSX 6063 Guizhou, China 123 788 308 SRR14596866 _ SRR14596865 Camellia pilosperma YangSX 4714 Guangxi, China 92 105 712 _ Camellia pitardii var. compressa YangSX 4576 Hunan, China 76 531 640 SRR14596864 _ YangSX 5555 Camellia semiserrata Guangdong, China 86 751 660 SRR14596863 _ Camellia sinensisvar. pubilimba YangSX 5927 Guangxi, China 85 785 132 SRR14596862 YangSX 5064 Sichuan, China 88 726 296 SRR14596861 Camellia szechuanensis _ YangSX 5798 Camellia tsingpienensis Guangxi, China 90 514 420 SRR14596860 YangSX 5202 Chongqing, China Camellia tuberculata 90 076 900 SRR14596859 NZ10477 Puntarenas, Costa Rica SRR14596857 Laplacea fruticosa 1 532 313 620 YXQ099 KY406760 Polyspora axillaris Hainan, China 92 917 500 SRR14596856 Yunnan, China 84 011 964 SRR14596855 **YXO221** Polyspora chrysandra Polyspora dalgleishiana BROWP 501 Royal Botanic Garden 83 838 300 SRR14596854 KY406769 Edinburgh, UK YXQ097 SRR14596853 KY406776 Polyspora hainanensis Hainan, China 97 170 208 Polyspora longicarpa YangSX 4779 Yunnan, China 89 251 900 SRR14596852 KY406768 Polyspora speciosa SRR14596851 KY406754 **YXQ145** Guangxi, China 96 508 612 Pyrenaria hirta var. cordatula YXQ169 Guangxi, China 92 426 012 SRR14596850 KY406785 YangSX 4067 Pyrenaria hirta yar, hirta Guangxi, China 100 342 800 SRR14596849 KY406772 Pyrenaria jonquieriana subsp. multisepala YXQ106 Hainan, China 87 583 392 SRR14596848 YangSX 5046 KY406756 Myanmar, Kachin 88 528 812 SRR14596846 Pyrenaria khasiana Pyrenaria menglaensis YXQ211 Yunnan, China 87 085 244 SRR14596845 KY406747 YXQ101 Hainan, China 81 080 788 SRR14596844 KY406764 Pyrenaria microcarpa var. microcarpa YXQ216 Yunnan, China 98 658 408 SRR14596843 KY406781 Pyrenaria oblongicarpa Pyrenaria pingpienensis **YXQ210** Yunnan, China 96 601 500 SRR14596842 YangSX 5038 Taiwan, China 101 120 720 SRR14596841 Pyrenaria shinkoensis SRR14596840 KY406753 Pyrenaria spectabilis var. greeniae YXQ172 Hunan, China 93 451 072

Guangxi, China

TABLE 1. List of taxa sampled in this study, with voucher, Illumina reads and GenBank accessions; species names that are underlined represent those species selected for PhyloNet analysis

SRR14596839

KY406765

88 012 020

Taxon	Voucher specimen	Sources	No. of reads (trimmed)	SRA number	Plastid genome
Outgroups					
Galax urceolata	Diapensiaceae	-	9 647 946	ERX2099546	_
Eurya acuminatissima	Pentaphylacaceae	-	25 009 135	SRX2786652	_
Ternstroemia gymnanthera	Pentaphylacaceae	-	14 421 549	ERX2099558	_
Sinojackia xylocarpa	Styracaceae	_	10 611 525	ERX2099565	_
Symplocos tinctoria	Symplocaceae	_	10 196 542	ERX2099566	_
Symplocos paniculata	Symplocaceae	-	28 374 406	SRX1601992	-

TABLE I. Continued

*A dash represents the plastid genes were extracted from the transcriptome data or no data were available.

previous studies (Prince and Parks, 2001; S. X. Yang *et al.*, 2004; M. M. Li *et al.*, 2013; Yu *et al.*, 2017*b*), these 21 orthologues probably contain hidden paralogues and could be problematic for downstream analysis. Therefore, we excluded these 21 orthologues to yield a final gene dataset with 610 orthologues (hereafter referred to as the reduced 610 orthologues dataset). Functional annotations for the 610 and 21 orthologues were performed using eggnog-mapper-2.1.4 (Huerta-Cepas *et al.*, 2017) with the command diamond; the best hit was chosen as the final annotation. Annotation results were visualized in WEGO2.0 (https://wego.genomics.cn/).

Sequence assembly for plastid genomes

To construct the plastid matrix, plastid protein-coding genes from 27 samples were extracted from the transcriptome or whole-genome sequencing data generated here, while the plastid genomes from 31 samples (Table 1) were obtained from complete plastid genomes already available from our previous study (Yu *et al.*, 2017*b*). For those species without plastid genome data, we first assembled the plastid genome using GetOrganelle v1.6.2e (Jin *et al.*, 2020), and the protein-coding sequences (CDS) were extracted from the transcriptome and whole-genome sequencing data. For those species with previously completed plastid genomes, protein-coding sequences were extracted following parallel methods to yield a combined matrix of all 58 samples (57 species) for which nuclear gene sequences were obtained.

Phylogenetic analyses based on nuclear genes and plastid genome

For nuclear genes, the obtained nucleotide sequences were aligned with MAFFT v7.407 (Katoh and Standley, 2013). Alignment statistics were calculated by AMAS (Borowiec, 2016). To better assess evolutionary history, both concatenation and coalescent approaches were used to reconstruct intergeneric relationships of Theaceae. For concatenation, partitioned ML and BI analyses were performed using RAxML v8.2.12 (Stamatakis, 2014) and MrBayes v3.2.6 (Ronquist *et al.*, 2012), respectively. Partitioning schemes and models were selected using PartitionFinder v2.1.1 for 610 orthologues (Lanfear *et al.*, 2016), and 188 subsets were obtained for the dataset. In the ML analysis, BS values were calculated using 1000 replicates.

In the BI analysis, four chains were run for $2\,000\,000$ generations with random initial trees; every 100 generations, trees were sampled, and the first 25 % of the trees were discarded as burn-in.

For the coalescent analysis, an ML gene tree was reconstructed for each orthologue with RAxML using the same parameter settings as above. The best ML gene tree and 100 bootstrap replicate trees generated for each orthologue set were used to estimate the species tree and supporting values in ASTRAL-III v5.6.3 (C. Zhang et al., 2018); nodes with bootstrap support below 10 % in all gene trees were removed using Newick utilities (Junier and Zdobnov, 2010), in order to improve accuracy in the ASTRAL analysis (C. Zhang et al., 2018). Support of the species tree was quantified using the local posterior probability (LPP) of a branch as a function of its normalized quartet support (Sayyari and Mirarab, 2016). The co-phylogenetic plot was created using the phytools package (Revell, 2012) in R to visualize the differences between trees. We used PhyParts (Smith et al., 2015) to examine patterns of gene tree concordance and conflict within the nuclear genome. All 610 gene trees were rooted using Phyx (Brown et al., 2017), and outgroups were removed. The results were visualized using the program phypartspiecharts (https://github.com/ mossmatters/phyloscripts/).

For the plastid genome sequence data, we obtained an 80-CDS dataset for the same taxa represented in the nuclear dataset. The nucleotide sequences were aligned with MAFFT v7.407 (Katoh and Standley, 2013). Phylogenetic reconstructions followed the approaches employed in our previous study (Yu *et al.*, 2017*b*).

Evolutionary network analysis

To test the previously proposed hypothesis of intergeneric hybridization in Theaceae (W. Zhang *et al.*, 2014), we used PhyloNet v3.8.2 (Than *et al.*, 2008) to infer an evolutionary network for Theaceae, using the command 'InferNetwork_MPL' under a maximum pseudo-likelihood framework (Yu and Nakhleh, 2015) and 505 individual gene trees with one outgroup species. Given that the computational time needed by network methods scales very rapidly with taxon number (Folk *et al.*, 2018), we reduced the sampling to 22 species, which is a computationally tractable size (i.e. <30 species; Than *et al.*, 2008; Wen *et al.*, 2018). Given that we mainly focused on hybridization events between genera, representative species from

all nine genera were used. Specifically, we selected one species for each monotypic genus or small genus (i.e. Apterosperma, Franklinia and Laplacea) and selected two (e.g. Schima) to six (e.g. Camellia) species representing the main subclades of each species-rich genus. This level of taxon sampling might impact our ability to detect reticulations within genera but would not necessarily impact our primary goal of investigating deep inter-generic introgression events. Only one outgroup species was used (different outgroup species were selected for each gene tree because of missing data) (Table 1). The maximum pseudo-likelihood algorithm requires a priori specification of the number of reticulating branches; the number of reticulations was set as one, two, three and four in repeated analyses, as per developer recommendations. Gene trees with branches with <70 % BS were collapsed, and five optimal networks were returned for each analysis. The command CalGTProb was used to compute the likelihood scores and select the best network.

Molecular dating based on 610 nuclear genes

We extracted the variable sites (187 255 bp in length) of the 610 orthologues dataset using MEGA v7 (Kumar et al., 2016) for our molecular dating analysis. Three species from Symplocaceae and Styracaceae, the most closely related families to Theaceae, were used as outgroups. Bayesian estimations of divergence times were conducted in BEAST v2.6.4 (Bouckaert et al., 2014), using the GTR + I + G nucleotide substitution model, lognormal uncorrelated relaxed clock model and birthdeath tree prior. We selected two fossil calibration points within Theaceae following our previous study (Table 2; Yu et al., 2017b) and conservatively set the two calibration constraints to the stems of those clades. Each fossil age was used to constrain the 'offset' in the lognormal distribution, with the mean 'M' set as 20 % (e.g. C4 in Table 2: offset = 23, M = 4.6) of the fossil age (checked 'Mean in Real Space') and the standard deviation 'S' set to 1.0. Additionally, three secondary calibration points were also used following our previous study (Yu et al., 2017b) and other studies which also used the Bayesian dating method such as BEAST or MCMCTREE (Magallón et al., 2015; Foster et al., 2017). First, the root of the tree, i.e. the stem age of Theaceae, was constrained under a uniform prior as 79.8-102.5 Mya; 79.8 and 102.5 Mya represent the minimum and maximum age of the 95 % HPD (highest posterior density) of the Theaceae stem in previous studies (Supplementary

Data Table S1; Magallón *et al.*, 2015; Foster *et al.*, 2017; Yu *et al.*, 2017b). Second, the crown of Theaceae and crown of Gordonieae and Theeae were also constrained under a uniform prior using the ages obtained from our previous study (Table 2; Yu *et al.*, 2017b). For each analysis, we ran one billion generations with sampling every 10 000 generations. Convergence was attained within 500–600 million generations, and the effective sample size (ESS) values for all parameters were >100. We removed the first 600 million generations as burn-in and used the remaining 40 000 trees to generate the maximum clade credibility (MCC) tree by TREEANNOTATOR v2.6.4 (with a PP limit of 0.5 and median node heights).

Inference of whole-genome duplication

To investigate the putative ancient WGD in Theaceae, we applied the Python package 'wgd' (Zwaenepoel and Van de Peer, 2019) to construct synonymous substitution (K_s) distributions (ranging from 0.05 to 3) among paralogues from 56 Theaceae transcriptomes and the six outgroup transcriptomes noted in Table 1. Using the command 'mcl' to blast and cluster sequences with each CDS, the commands 'ksd' and 'mix' were used to construct the *Ks* distribution and mixture modelling of *Ks* distributions, respectively. For analysis of the mixture model, we used the BGMM method in the wgd package.

RESULTS

Characteristics of transcriptomes and datasets

We sequenced transcriptomes (ranging from 5.85 to 10.6 Gb) of 56 individuals from 55 species and obtained 138.7 Gb and 165.7 Gb (~89.5× coverage, based on an estimated genome size of 1.85 Gb for *Stewartia pteropetiolata* from flow cytometry) of genomic data for *Laplacea fruticosa* and *Stewartia malacodendron*, respectively. From trimmed reads, assembly of the 64 transcriptomes/genomes (56 transcriptomes and two genomes generated in this study and six transcriptomes from GenBank) provided an average length of unigenes from 382 to 1578 bp. The N50 length ranged from 377 to 1959 bp, with an average of 1491 bp (Supplementary Data Table S1). In total, 610 orthologues were obtained from 56 transcriptomes and two whole-genome sequencing datasets, with the aligned length of the orthologue sets ranging from 309 to 8854 bp and the

TABLE 2. Fossils used for calibrations in this study, selected from Yu et al. (2017)

Calibration nodes	Fossils	Calibration types	Ages (epoch)	Fossil assignment	Constrained age	References
C1	NA	Secondary calibration	79.8–102.5	Root of the tree (stem of Theaceae)	79.8–102.5	(Foster <i>et al.</i> , 2017; Yu <i>et al.</i> , 2017b)
C2	NA	Secondary calibration	39.6–74.7	Crown of Theaceae	39.6–74.7	(Yu <i>et al.</i> , 2017 <i>b</i>)
C3	NA	Secondary calibration	33.8-66.9	Crown of Gordonieae and Theeae	33.8-66.9	(Yu et al., 2017b)
C4	Schima kwangsiensis X. G. Shi, C. Quan et J. H. Jin	Fruits, seeds fossil	Late Oligocene	Stem of Schima	23.0	(M. M. Li <i>et al.</i> , 2013; Quan <i>et al.</i> , 2016; X. G. Shi <i>et al.</i> , 2017)
C5	Hartia quinqueangularis(Menzel)	Fruits, seeds fossil	Late Miocene	Stem of <i>Hartia</i> (now a clade within <i>Stewartia</i>)	5.3	(Mai, 1975)

proportion of missing data ranging from 0 to 29.65 % (Table S2). The aligned length of the concatenated 610 orthologues was 858 606 bp, with 227 708 (26.5 %) variable sites and 21.67 % missing data. The alignment length of the concatenated 80 plastid coding genes was 69 225 bp, with 7236 (10.5 %) variable sites and 11.88 % missing data. Gene function of the 610 and 21 orthologues did not show a significant difference, with genes from both sets mainly related to the cellular component, molecular function and general biological process (Table S3, Figs S1 and S2). All transcriptomic raw reads have been deposited in GenBank (Table 1), and all of the alignments and trees in this study have been submitted to TreeBASE (see Supplementary Data statement at the end of the paper).

Phylogenetic relationships and networks in Theaceae

The topology recovered from RAxML analyses based on the concatenated 610 low-copy nuclear genes strongly supported a sister relationship between Theeae and Gordonieae (MLBS = 100 %, PP = 1.00; Fig. 1). Additionally, PhyParts analysis indicated 435 out of 610 gene trees (71.3 %) supported this topology (Fig. 2). Our coalescent-based species tree inferred from ASTRAL revealed the same relationship among the three tribes as obtained with the concatenation analysis (LPP = 1.00; Fig. 1; Supplementary Data Fig. S3). RAxML analyses based on the 80 protein-coding genes of the plastid genome also supported a sister relationship between Theeae and Gordonieae (MLBS = 91 %, PP = 1.00; Fig. S4). The co-phylogenetic plot between the concatenated and coalescent tree topology of the 610 nuclear genes (Fig. 1), and between the nuclear coalescent tree and plastome topology (Fig. S4) did not show any incongruence of relationships among the three tribes. Hence, support at the tribal level of Theaceae was strong and uniform across the data partitions and analytical methods employed here.

Generic-level relationships recovered within Gordonieae and Stewartieae were largely consistent with previous phylogenetic studies of Theaceae. Within Theeae, the sister relationships between Camellia and Polyspora and between Apterosperma and Laplacea fruticosa were both maximumly supported in all of the analyses using transcriptomic data (Fig. 1; Supplementary Data Fig. S3). Nevertheless, the position of the Apterosperma-Laplacea clade varied among analyses. Using the 610 low-copy nuclear genes, the concatenation analysis placed the Apterosperma-Laplacea clade grouped with Pyrenaria with moderate support (MLBS = 72 %, PP = 1.00; Fig. 1). The ASTRAL topology (Fig. S3), while largely congruent overall with the results from the concatenated analyses (Fig. 1), instead placed Apterosperma-Laplacea as sister to the Camellia-Polyspora clade with very weak support (LPP = 0.35; Fig. S3). At lower taxonomic levels, PhyParts recovered strong discordance across many parts of the tree, with only 112 out of 610 gene trees supporting the sister relationship between the Apterosperma-Laplacea clade and the Camellia-Polyspora clade, and 488 gene trees supporting conflicting/alternative resolutions (Fig. 2). Phylogenetic trees based on the dataset of 80 plastid protein-coding genes were highly consistent with our previous study (Yu et al., 2017b) with the exception that Apterosperma was sister to Camellia (MLBS = 86 %, PP = 1.00; Fig. S4).

For the PhyloNet analyses, the inferred network with the highest log pseudo-likelihood (-8792.9795) included three reticulation events (reticulation numbers were set as 4; Fig. 3). This analysis suggested one reticulation between *Camellia tsingpienensis* and *C. amplexifolia*, and the descendant putative hybrid further backcrossed with *C. tsingpienensis* and formed the clade comprising *C. huana* and *C. szechuanensis*. We also recovered evidence of a reticulation event suggesting that *Franklinia alatamaha* descended from a putative ancient hybridization event between the common ancestor of *Gordonia* and the common ancestor of *Schima*. All other analyses (reticulation numbers = 1, 2, 3), for which likelihood was suboptimal, only detected reticulation within *Camellia* (Fig. 3).

Divergence time estimation

The stem and crown ages of Theaceae were estimated to be 99.7 Mya (95 % HPD: 92.0–102.5) and 64.9 Mya (56.4–73.3), respectively (Fig. 4; Supplementary Data Table S4). The divergence between Gordonieae and Theeae was ~55.7 Mya (48.1–62.8). The crown ages of Stewartieae, Gordonieae and Theeae were estimated to be 18.5 Mya (15.6–21.7), 25.1 Mya (24.1–26.6) and 22.1 Mya (19.2–25.2), respectively. The estimated ages were all within the 95 % HPD of our previous study (Yu *et al.*, 2017*b*) and Yan *et al.* (2021).

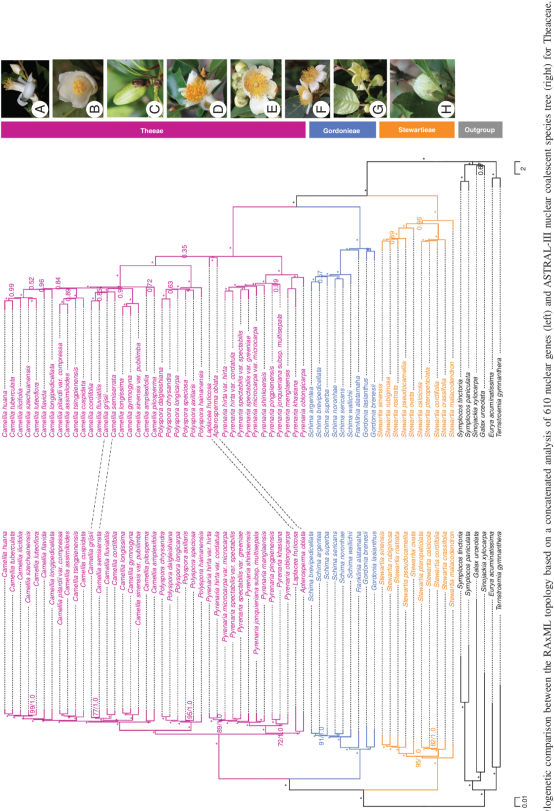
Whole-genome duplication

Ks (synonymous substitution rate) analyses using all 57 ingroup taxa suggested that all species showed a peak at around Ks = 1.5, and all except one species from Theaceae presented a peak at around Ks = 0.4 (Fig. 4; Supplementary Data Fig. S5). The two peaks are consistent with the two WGD events (At- γ : Ks = 1.16; Ad- β : Ks = 0.36) reported in the tea (*Camellia*) sinensis var. sinensis) genome (Xia et al., 2017). No Ks peak around 0.4 was found in Stewartia ovata; given the nested position of this species, the absence may be an artefact due to stochastic error or may be because the transcriptome data only represent the expressed genes in those tissues (e.g. roots, leaves, flowers) sampled. For the six outgroup species, consistent Ks peaks were also found at around 1.5 and 0.4 (Fig. S5), indicating that Symplocaceae, Styracaceae, Pentaphylacaceae and Diapensiaceae shared the two WGD events (i.e. At- γ , Ad- β) with Theaceae.

DISCUSSION

Consistent relationships among tribes between plastome and transcriptome trees

Numerous phylogenetic studies have not been able to elucidate with strong support the relationships among the three tribes within Theaceae, due in part to incomplete taxon sampling and few loci, either a few plastid and nuclear loci or half of the SSC region of the plastid genome (Prince and Parks, 2001; S. X. Yang *et al.*, 2004; J. B. Yang *et al.*, 2006; M. M. Li *et al.*, 2013). Using 25 loci from the plastid, nuclear and mitochondrial





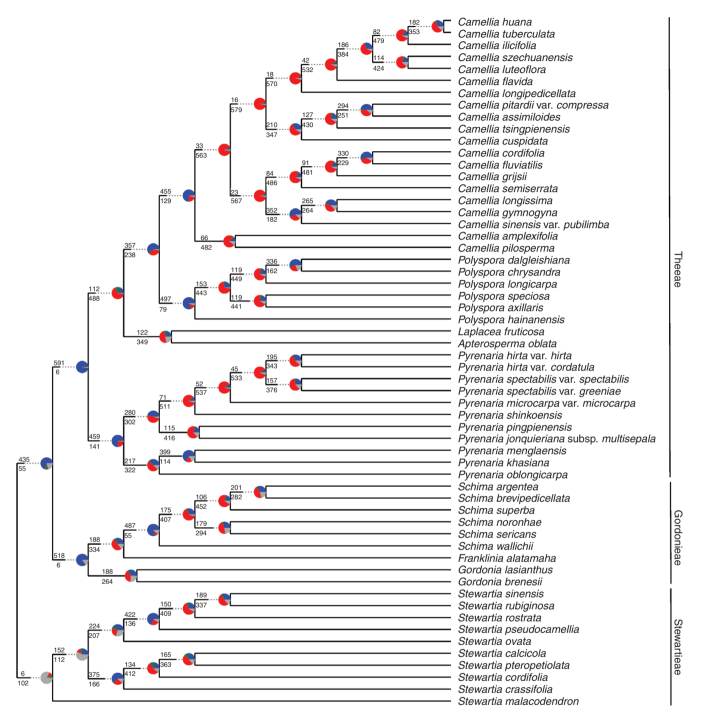


FIG. 2. Patterns of gene-tree concordance and conflict within Theaceae based on the PhyParts analysis. The tree topology used is that inferred by ASTRAL-III. The pie charts at each node show the proportion of genes in concordance (blue), conflict (green = a single dominant alternative; red = all other conflicting trees) and without enough information (grey). The numbers above and below each branch are the numbers of concordant and conflicting genes at each bipartition, respectively.

genomes from 4531 species from Ericales, a recent study found that Theeae and Gordonieae grouped together (MLBS > 70 %, PP > 0.95) with Stewartieae as their sister, but only ten species of Theaceae were included (Rose *et al.*, 2018). Our previous study likewise strongly supported a sister relationship between Theeae and Gordonieae (MLBS = 91 %, PP = 1.00) based on

a combined plastome and nuclear ribosomal DNA dataset (Yu *et al.*, 2017*b*). Here we present a phylogenetic framework of Theaceae using 610 orthologous low-copy nuclear genes. The topology obtained from both the concatenation and the coalescence analyses of the 610 low-copy nuclear genes consistently and strongly supported a sister relationship between Theeae

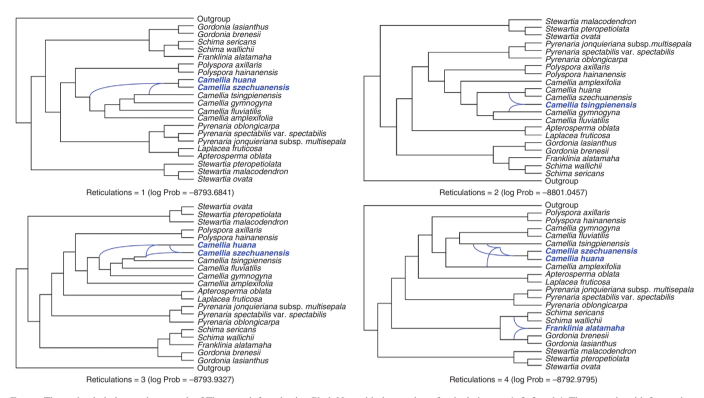


FIG. 3. The optimal phylogenetic network of Theaceae inferred using PhyloNet, with the number of reticulations as 1, 2, 3 and 4. The scenario with four reticulations had the best log pseudo-likelihood.

and Gordonieae (MLBS = 100 %, PP = 1.00, LPP = 1.00; Fig. 1; Supplementary Data Fig. S3), consistent with the results from the plastid genome (MLBS = 91 %, PP = 1.00; Fig. S4). In addition, 435 of the 610 low-copy nuclear genes supported this topology (Fig. 2). The ((Gordonieae, Theeae) Stewartieae) relationship is consistent with the evolutionary pattern of the endosperm in Theaceae, as discussed in Yu *et al.* (2017*b*).

Improved intergeneric relationships based on transcriptome data

Intergeneric relationships within Gordonieae and Stewartieae have been fully resolved in previous studies (Prince and Parks, 2001; S. X. Yang et al., 2004; J. B. Yang et al., 2006; M. M. Li et al., 2013; Yu et al., 2017b). However, the relationships among the five genera in Theeae have been controversial. Laplacea was placed in a clade comprising *Camellia*, *Tutcheria* (=*Pyrenaria*) and Glyptocarpa (=Camellia) (Prince and Parks, 2001). W. Zhang et al. (2014) revealed that Apterosperma formed a sister relationship with Polyspora (MLBS = 73 %, PP = 1.00) in the plastid DNA tree, but these two genera were placed in a clade comprising Tutcheria (=Pyrenaria) and Parapyrenaria (=Pyrenaria) (MLBS = 68 %, PP = 0.72) in the *LEAFY* tree. Based on the 610 low-copy nuclear genes, the resolution of the relationships among the five genera in Theeae has been improved. The Apterosperma-Laplacea clade received maximal support in the concatenation analyses using the 610 low-copy nuclear genes and grouped with Pyrenaria with moderate support (MLBS = 72%, PP = 1.00; Fig. 1). However, the ASTRAL topology suggested that the Apterosperma-Laplacea clade was weakly supported as sister to the Camellia-Polyspora clade

(LPP = 0.35; Fig. 1; Supplementary Data Fig. S3); 112 out of 610 nuclear genes supported this topology (Fig. 2). The strongly supported *Apterosperma–Laplacea* clade also grouped with *Pyrenaria* with moderate support based on the whole plastid genome dataset (MLBS = 67 %), the SSC (MLBS = 80 %) dataset and the protein-coding gene dataset (MLBS = 75 %) from our previous study (Yu *et al.*, 2017*b*). Taken as a whole, based on evidence from both plastid genomes and transcriptome data, we suggest (((*Apterosperma–Laplacea*), *Pyrenaria*), (*Camellia–Polyspora*)) as the most likely topology.

Phylogenetic network inference suggests three reticulation events in Theaceae

In W. Zhang et al. (2014), Camellia and Pyrenaria were not recovered as monophyletic, and widespread hybridization among genera in Theeae was proposed. Phylogenetic conflict found in Stewartia was also suggested to be caused by ancient introgressive hybridization following species diversification, leading to discordant histories in the nuclear and plastid genomes (H. Y. Lin et al., 2019). However, while our PhyloNet analyses supported the presence of hybridization in the history of Theaceae (Fig. 3), they do not support the specific reticulation scenario suggested by W. Zhang et al. (2014); Camellia and Pyrenaria were supported as monophyletic based on the 610 low-copy nuclear genes (Fig. 1; Supplementary Data Fig. S3). Two of the three reticulation events detected in the best-fit network were within Camellia, and another intergeneric reticulation was in Gordonieae, but not Theeae (Fig. 3).

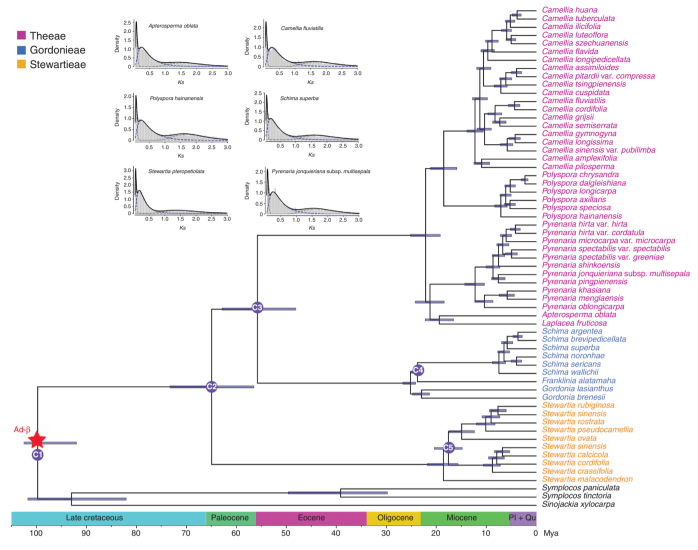


FIG. 4. Chronogram of Theaceae derived from BEAST analysis using 610 nuclear genes with *Ks* distribution plots for paralogues showing inferred WGDs (blue dashed line) in the upper left corner; the six species included represent all three tribes of Theaceae. The ages of stratigraphic boundaries were obtained from the International Chronostratigraphic Chart (Cohen *et al.*, 2013) (Pl, Pliocene; Qu, Quaternary), with a scale as millions of years ago (Mya). Blue bars at each node represent the 95 % highest posterior density (HPD) with posterior probability above 0.5. C1–C5 represent the five calibration points used, which correspond to those nodes listed in Table 2. The red star represents the WGD event shared by all members of the clade including core Ericales, primuloids, polemonioids and Lecythidaceae, reported in Zhang *et al.* (2020).

The estimated stem and crown ages of Theaceae are largely consistent with previous studies (M. M. Li *et al.*, 2013; Magallón *et al.*, 2015; Foster *et al.*, 2017; Yu *et al.*, 2017b; Rose *et al.*, 2018; Yan *et al.*, 2021) (Supplementary Data Table S4). Our study supported the findings of Yan *et al.* (2021) that Theaceae are of boreotropical forest origin. Under the deep reticulation scenario (Fig. 3), *Franklinia alatamaha* (the only extant species of the genus, distributed in eastern North America) descends from a putative ancient hybridization event between the common ancestor of *Gordonia* (endemic to North and Central America) and the common ancestor of *Schima* (endemic to Asia). The molecular dating analysis suggests a date of reticulation during the Late Oligocene, ~25.1 Mya (24.1–26.6) (Fig. 4; Table S4). A recent study also suggested that gene flow between species

of Theaceae from different continents (e.g. North America, Eurasia) must have occurred during the Oligocene and possibly until the mid-Miocene (Yan *et al.*, 2021). In the study by Yan *et al.* (2021), the ancestral distribution of the crown of Gordonieae was inferred as Nearctic + Sino-Japanese (NS); it is likely that the ancestors of *Gordonia* and *Schima* co-occurred across the NS region during the late Oligocene (~25.6 Mya), and hence ancient gene flow between them was possible. This is a plausible scenario as land bridges (e.g. the Bering land bridge) existed during the Late Oligocene, and the eastern Asia and eastern North American flora was probably continuous across high latitudes of the Northern Hemisphere (Tiffney, 1985; Tiffney and Manchester, 2001; Milne, 2006), allowing for species contact and opportunities for hybridization, which is no longer possible under today's climate and geography. Fossils of Theaceae are known from mid-latitude Northern Hemisphere localities such as western Kentucky and Tennessee in North America and Germany in Europe during the Eocene and Oligocene (Grote and Dilcher, 1992; Kvacek and Walther, 1998; Wilde and Frankenhauser, 1998; Kvaček, 2004).

Overall, our work supports ancient introgression within Theaceae, and is consistent with biogeographical patterns and the fossil history of the group, adding to an increasing list of ancestral hybrids among currently allopatric taxa that yield unique evidence of past biogeographical distributions (e.g. Folk et al., 2018). Interspecific gene flow of an intercontinental scope, although perhaps less geographically remarkable considering that the past distribution of these plants was probably higher in latitude, has likewise been reported in other plant groups distributed across Eurasia and North America. For example, phylogenetic and fossil evidence supports a North American origin for *Picea* (Pinaceae); the Bering land bridge may have facilitated the introgression between species from North America and Eurasia during the Miocene and Pliocene (Ran et al., 2015). Discordance of the mtDNA tree with nuclear and cpDNA trees of Abies (Pinaceae) also indicated intercontinental migration and introgressive hybridizations in this genus during the Miocene (Semerikova et al., 2018). More detailed studies are needed to explore the degree and frequency of intercontinental gene flow between Eastern Asia and North America in other lineages. In particular, a continued recovery of a primarily Miocene date for past hybridization would be of interest in understanding when plant biogeographical connections ceased among these areas.

The best-fit network suggested the clade comprising C. huana and C. szechuanensis was formed through two rounds of hybridizations, with Camellia tsingpienensis and C. amplexifolia as parents. All other PhyloNet analyses (reticulations = 1, 2, 3) consistently indicated a clear pattern of intrageneric gene flow within Camellia (Fig. 3). Camellia tsingpienensis is found in Guangxi, south-eastern Yunnan of China and northern Vietnam, and C. amplexifolia is only present in Hainan of China. Recent studies have suggested that the flora of Hainan is of continental origin and has the highest floristic affinity with Vietnam, and periodic emergence of land bridges between Hainan and Vietnam during Quaternary glacial cycles might have resulted in their floristic affinity (Ali, 2018; S. L. Lin et al., 2021). Even though there are no distribution overlaps between C. tsingpienensis and C. amplexifolia, interspecific gene flow was possible when Hainan and the neighbouring landmasses including Vietnam were connected during the glacial periods of the Quaternary.

Previous studies have likewise found evidence of hybridization in *Camellia*. First, cultivated ornamental camellias resulting from hybridization have been widely used in horticulture (Nishimoto *et al.*, 2003; Tanaka *et al.*, 2005; Xu *et al.*, 2018). Second, Cambod tea (cultivated tea of *C. sinensis* var. *assamica*) was suggested to have originated through hybridization between different tea types (Meegahakumbura *et al.*, 2016; Meegahakumbura *et al.*, 2018). Introgression was also detected between the cultivated *C. sinensis* var. *assamica* and *C. taliensis*, with the latter possibly genetically involved in the domestication of *C. sinensis* var. *assamica* (Li *et al.*, 2015). Given our taxon sampling decisions, further work with increased taxon sampling will be needed to uncover further introgression patterns within *Camellia*.

No Theaceae-specific whole-genome duplication event

Two ancient WGD events, namely At- γ and Ad- β , have been identified in the tea plant (C. sinensis var. assamica) genome (Xia et al., 2017, 2020). However, analysis of genic collinearity reveals that a recent WGD event occurred after the divergence of the tea and kiwifruit lineages, based on the genome of another variety of tea (C. sinensis var. sinensis) (Wei et al., 2018); Larson et al. (2020) later named this WGD Cm-α. Based on a chromosome-scale genome assembly of C. sinensis var. sinensis, the authors suggested one recent Camellia tetraploidization event occurred after the divergence of C. sinensis and Actinidia chinensis from their common ancestor (J. D. Chen et al., 2020), but the time of the Camellia tetraploidization event (58.9-61.7 Mya) was very close to the divergence time between C. sinensis and A. chinensis at 61.2-65.3 Mya. Here, we identified two WGD events shared by all genera of Theaceae and also representatives of other related families (Symplocaceae, Styracaceae, Pentaphylacaceae and Diapensiaceae; Fig. 4; Supplementary Data Fig. S5). We have clarified that Cm- α proposed by Wei *et al.* (2018) and Larson et al. (2020) and the more recent tetraploidization event found by J. D. Chen *et al.* (2020) were actually Ad- β , which has been recently revised to characterize the core Ericales $(ACCH\beta + DIOS\alpha)$ according to genome collinearity and also analyses based on the MAPS pipeline (Leebens-Mack et al., 2019), and more specifically to the clade including core Ericales, primuloids, polemonioids and Lecythidaceae, using deep asterid phylotranscriptomic analyses (C. Zhang et al., 2020). Thus, our results support the hypothesis that the tea family experienced two WGD events (i.e. At- γ and Ad- β) in its evolutionary history, with both shared by other families. There is no evidence for any Theaceae-specific WGD event. Thus, this study sheds light on the significance of broad phylogenetic sampling for inferring the number and placement of WGD events.

SUPPLEMENTARY DATA

Supplementary data are available online at https://academic.oup. com/aob and consist of the following. Figure S1: Gene annotation of 610 orthologous low-copy nuclear genes. Figure S2: Gene annotation of 21 putative paralogous low-copy nuclear genes. Figure S3: Species tree topology, inferred by ASTRAL, from the 610 low-copy nuclear genes; posterior probability values are shown beside the nodes. Figure S4: Co-phylogenetic comparison between ASTRAL-III nuclear coalescent species tree and the RAxML plastome topology of Theaceae. Figure S5: Histograms of the distribution of gene duplications with mixture models of inferred WGDs for all sampled ingroup and outgroup species. Table S1: Statistics of sampled transcriptomes. Table S2: Results of AMAS summaries on 610 low-copy nuclear genes. Table S3: Functional annotation of 610 and 21 low-copy nuclear genes. Table S4: Age estimates of Theaceae (millions of years ago) for selected nodes and comparison with results from other studies.

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