

# Widespread ancient whole-genome duplications in Malpighiales coincide with Eocene global climatic upheaval

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## Summary

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- Whole-genome duplications (WGDs) are widespread and prevalent in vascular plants and frequently coincide with major episodes of global and climatic upheaval, including the mass extinction at the Cretaceous–Tertiary boundary (c. 65 Ma) and during more recent periods of global aridification in the Miocene (c. 10–5 Ma). Here, we explore WGDs in the diverse flowering plant clade Malpighiales.
- Using transcriptomes and complete genomes from 42 species, we applied a multipronged phylogenomic pipeline to identify, locate, and determine the age of WGDs in Malpighiales using three means of inference: distributions of synonymous substitutions per synonymous site ( $K_s$ ) among paralogs, phylogenomic (gene tree) reconciliation, and a likelihood-based gene-count method.
- We conservatively identify 22 ancient WGDs, widely distributed across Malpighiales subclades. Importantly, these events are clustered around the Eocene–Paleocene transition (c. 54 Ma), during which time the planet was warmer and wetter than any period in the Cenozoic.
- These results establish that the Eocene Climatic Optimum likely represents a previously unrecognized period of prolific WGDs in plants, and lends further support to the hypothesis that polyploidization promotes adaptation and enhances plant survival during episodes of global change, especially for tropical organisms like Malpighiales, which have tight thermal tolerances.

## Introduction

Whole-genome duplication (WGD), or polyploidy, is an important evolutionary force that has shaped plant evolution. It has long been appreciated that the formation of recent polyploids in vascular plants is common (Stebbins, 1947; Barker *et al.*, 2016), and mounting evidence suggests that ancient polyploids are more frequent than once thought. WGD has been identified in ferns (Wood *et al.*, 2009) and seed plants, including in gymnosperms (Li *et al.*, 2015) and, more prevalently, in angiosperms. Well-cited examples of ancient WGDs have been associated with the origin of numerous hyperdiverse clades, including in the common ancestor of seed plants, flowering plants, monocots, orchids, core eudicots, mustards (Brassicaceae), legumes (Fabaceae), and sunflowers (Asteraceae; Blanc & Wolfe, 2004; Barker *et al.*, 2008, 2009; Bertoli *et al.*, 2009; Tang *et al.*, 2010; Jiao *et al.*, 2011, 2012, 2014; Cannon *et al.*, 2015; Huang *et al.*, 2016; Zhang *et al.*, 2017). Among these ancient WGDs, several have

been dated to the Cretaceous–Tertiary (KT) boundary (c. 65 Ma), potentially linking these polyploidization events to plants' abilities to survive abrupt global environmental change (Fawcett *et al.*, 2009; Vanneste *et al.*, 2014). Similarly, a large number of WGDs have also been reported in grasses during the late Miocene, when arid, grass-dominated landscapes expanded dramatically (Estep *et al.*, 2014). In these cases, the potential adaptive value of WGDs is thought to arise from the origin of genetic novelties (Ohno, 1970; Lynch & Conery, 2000; Taylor & Raes, 2004; Lynch, 2007) and by masking the effects of deleterious mutations (Gu *et al.*, 2003). Together, this may facilitate plant survival across periods of global disruption. Although debate exists as to the influence of WGD on species diversification rates (Wood *et al.*, 2009; Mayrose *et al.*, 2011, 2015; Soltis *et al.*, 2014; Tank *et al.*, 2015; Kellogg, 2016), it is generally accepted that chromosomal rearrangements from WGDs can significantly accelerate isolating barriers, thus promoting cladogenesis (Werth & Windham, 1991; Lynch & Force, 2000; Husband

*et al.*, 2016). In summary, it is established that WGDs are a prominent feature of vascular plant evolution, yet they remain underexplored in many clades.

Here, we investigate WGDs in the large and diverse angiosperm order Malpighiales, which contains more than 16 000 mostly tropical species with tremendous morphological and ecological diversity. Members of this clade also include numerous economically important crops, such as rubber, cassava, and flax. The Malpighiales have long been recognized as one of the most difficult clades to resolve in the flowering plant tree of life (Davis *et al.*, 2005; Wurdack & Davis, 2009), which has been attributed in part to its rapid radiation in the mid-Cretaceous (Davis *et al.*, 2005; Xi *et al.*, 2012b). However, recent efforts utilizing phylogenomic approaches have greatly increased our understanding of deep-level relationships in the order (Xi *et al.*, 2012b). Additionally, the clade includes numerous species that have previously been targeted for genomic investigation of WGDs. Eight genomes are currently available for interrogation: *Hevea brasiliensis* (rubber), *Manihot esculenta* (cassava), *Linum usitatissimum* (flax), *Ricinus communis* (castor bean), *Jatropha curcas* (Barbados nut), *Populus trichocarpa* (black cottonwood), *Salix suchowensis* (shrub willow), and *Salix purpurea* (purple willow). Notably, three WGDs have been identified in Malpighiales using these data: in the common ancestor of *Populus* and *Salix* (35–65 Ma; Tuskan *et al.*, 2006; Dai *et al.*, 2014); in the common ancestor of *Manihot* and *Hevea* (35–47 Ma; Bredeson *et al.*, 2016); and more recently in *L. usitatissimum* (5–9 Ma; Wang Z. *et al.*, 2012). In addition, studies using transcriptomes have identified an older WGD shared by all blue-flowered *Linum* species, including *L. usitatissimum*, at 20–40 Ma (Sveinsson *et al.*, 2014). More complicated polyploidy histories involving multiple rounds of WGDs and hybridization have also been reported using chromosome count data in the genera *Passiflora* (Mayrose *et al.*, 2009) and *Viola* (Marcussen *et al.*, 2012, 2014). In short, given the apparent propensity of WGDs in Malpighiales and the existing complete sequence data available for the order, this clade is an ideal study system for investigating the frequency and timing of WGDs.

## Materials and Methods

### Taxon sampling and transcriptome sequencing

We collected genomic and transcriptomic data for 36 species representing 21 families of Malpighiales, spanning all major clades *sensu* Wurdack and Davis (Wurdack & Davis, 2009; Chase *et al.*, 2016) (Supporting Information Tables S1–S3). In addition, three closely related outgroups from Celastraceae (Celastrales), Elaeocarpaceae (Oxalidales), and Oxalidaceae (Oxalidales), plus three more distantly related outgroups (*Cucumis sativus* (Eurosoid I), *Theobroma cacao* (Eurosoid II), and *Vitis vinifera* (basal Rosid)) were used for rooting (Chase *et al.*, 2016). We sequenced transcriptomes of 15 Malpighiales species following the protocol described by Xi *et al.* (2012a). Total RNA from leaf tissue was extracted using the RNAqueous and Plant RNA Isolation Aid kits (Ambion Inc.), and treated with the TURBO DNA-free kit

(Ambion, Inc.) at 37°C for 4 h to remove residual DNA. The complementary DNA library was synthesized from total RNA following the protocols of Novaes *et al.* (2008). Illumina paired-end libraries were prepared for complementary DNA following the protocols of Bentley *et al.* (2008). Each library was sequenced in a single lane of a Genome Analyzer II instrument (Illumina, Inc.) with paired-end 150 bp read lengths. We additionally included 13 annotated transcriptomes from the OneKP project (Table S2) to complete our taxon sampling for Malpighiales (Matasci *et al.*, 2014). These sequences were obtained following the protocol outlined by Wickett *et al.* (2014). Finally, we also obtained whole-genome sequence data from eight published genomes of Malpighiales plus three outgroup species: *H. brasiliensis* (Willd. ex A.Juss.) Müll.Arg., *J. curcas* L., *L. usitatissimum* L., *M. esculenta* Crantz, *P. trichocarpa* Torr. & A.Gray ex Hook., *R. communis* L., *S. purpurea* L., *S. suchowensis* W.C.Cheng ex G.H.Zhu, *C. sativus* L., *T. cacao* L., *V. vinifera* L. (Table S3).

### Transcriptome assembly

Raw sequencing reads were first corrected for errors using Rcorrector (Song & Florea, 2015). Reads marked as ‘unfixable’, generally constituting regions of low complexity, were discarded. Sequencing and PCR adapters were identified and trimmed using TRIMGLORE v.0.4.2 (Krueger, 2015). We examined the quality of trimmed reads using FASTQC v.0.11.5 (Andrews, 2010) and then assembled the reads using TRINITY v.2.1.1 (Grabherr *et al.*, 2011). We used the longest isoform from each Trinity assembly and further reduced the redundancy generated from sequencing error or alternative splicing by performing a similarity-based clustering ( $-c$  0.99  $-n$  10, threshold following Yang *et al.*, 2015) via CD-HIT-EST v.4.6.4 (Li & Godzik, 2006). The completeness of our assemblies was assessed by comparison against the single-copy orthologs plant database, BUSCO (Simão *et al.*, 2015; Fig. S1). Coding regions of each putative transcript were predicted following the transdecoder workflow (Haas & Papanicolaou, 2012). Finally, to control for transcriptome quality in our subsequent assessments of WGD, we reanalyzed our combined transcriptome and complete genome data following the methods described later with only high-quality transcriptomes. Here, transcriptomes with more than 40% (382/956) missing BUSCOs were removed, including *Bhesa paniculata*, *Flacourtia jangomas*, *Galearia maingayi*, *Ixonanthes reticulata*, *Podostemum ceratophyllum*, *Rinorea anguifera*, and *Tristellateia australasiae*.

### Gene family clustering and orthology inference

To assign sequences into orthologous gene families, we used an integrated method that takes into account sequence similarity and species phylogeny. We first constructed whole genome/transcriptome homology scans using PROTEINORTHO v.5.13 (Lechner *et al.*, 2011) with default parameter settings. This program extends the reciprocal best BLAST hit method and is computationally efficient. Clusters were searched to identify gene families containing at least 22 (>60%) in-group species. This resulted in

8465 candidate homolog clusters. This similarity-based homology search can sometimes be erroneous due to deep paralogs, misassembly, or frame shifts (Yang & Smith, 2014). To reduce such errors in orthology inference, we further applied a tree-based method to sort genes into orthology groups (Yang & Smith, 2014). This method does not rely on a known species tree, but rather iteratively searches for the subtree with the highest number of in-group taxa to assign as orthology groups (Fig. S2). Here, we first aligned the protein sequences of each homolog cluster with MAFFT v.7.299 (Kato & Standley, 2013) using the local alignment algorithm (-localpair -maxiterate 1000). The resulting protein alignments were converted into the corresponding codon alignments using PAL2NAL v.14 (Suyama *et al.*, 2006). A gene family tree of each codon alignment was then reconstructed using RAxML v.8.1.5 (Stamatakis, 2014b) with 10 random starting points. To sort homologs into ortholog groups, we first pruned exceptionally long and short branches within each gene family tree because we suspected such branches to be incorrect homologs, sequencing errors, or transcript isoforms. In addition, short branches may cause problems of overfitting in the subsequent penalized likelihood dating method (Sanderson, 2004). Along these lines, branches that were 10 times longer than the '5% trimmed mean branch length', or shorter than an absolute value of  $1 \times 10^{-15}$ , were pruned. The '5% trimmed mean branch length' was defined as the mean branch length after discarding the lowest and highest 5% of the branch length distribution in each gene family. Orthology was then inferred based on this pruned gene family tree using the 'rooted tree' method ('prune\_paralogs\_RT.py') following Yang & Smith (2014). The resulting 5113 orthology clusters were realigned as amino acids using the method already described. Finally, the back-translated nucleotide alignments were used for subsequent phylogenetic analysis.

### Phylogeny reconstruction and molecular dating

To infer the phylogeny of each orthology group, we first removed sites containing >80% gaps using TRIMAL v.1.4.15 (Capella-Gutiérrez *et al.*, 2009). We then applied RAxML v.8.1.5 to reconstruct maximum likelihood (ML) trees under the GTR +  $\Gamma$  model with 20 random starting points. We chose the GTR +  $\Gamma$  model because it accommodates rate heterogeneity among sites, whereas the other available GTR model in RAxML, the GTRCAT model, is less appropriate owing to our small taxon sampling size (Stamatakis, 2014a). We then filtered each gene tree to eliminate exceptionally long and short branches using the method outlined earlier. The remaining gene accessions were realigned and a final round of ML tree inference was conducted. Statistical confidence of each gene tree was assessed by performing 100 bootstrap (BS) replicates with branch length (-N 100 -k).

In addition to determining the phylogeny of orthology groups, we also estimated the divergence time of each orthologous tree for the purpose of dating WGDs. We applied the penalized likelihood method for its high efficiency when dealing with large data sets including thousands of genes. By contrast, Bayesian

divergence time estimation is computationally intensive and difficult to implement with a data set of this size. We estimated molecular divergence times for each ML gene tree as well as the BS trees with penalized likelihood as implemented in r8s v.1.7 (Sanderson, 2003). The following four calibration points were applied to each tree: the root age was fixed at 109 Ma, representing the approximate age of the crown group divergence in Malpighiales (Xi *et al.*, 2012b); the minimum age of crown group clusoids (including *Calophyllum macrocarpum*, *Clusia rosea*, *P. ceratophyllum*, *Garcinia oblongifolia*, *Hypericum perforatum*, and *Mammea americana*) was set to be 89 Ma, representing the oldest known fossil in Malpighiales, *Palaeoclusia chevalieri* (Ruhfel *et al.*, 2013); and two additional secondary minimum age constraints from Xi *et al.* (2012b) to constrain stem group Euphorbiaceae (97 Ma) and Salicaceae (69 Ma). For each clade, the age constraint was placed on the most recent common ancestor of all gene accessions forming a monophyletic group for that clade. The optimal smoothing parameter for each gene tree was determined within the range of parameter space ( $1 \times 10^{-4.5}$ ,  $1 \times 10^{4.5}$ ) by cross-validation (Sanderson, 2003). Trees were subsequently dated under the assumption of a relaxed molecular clock by applying a semiparametric penalized likelihood approach using a truncated Newton optimization algorithm in r8s.

### Species tree estimation

We inferred a single reference species tree for our analysis of WGD applying a summary coalescent method as implemented in ASTRAL v.4.10.5 (Mirarab & Warnow, 2015). As input gene trees for our species tree analysis we utilized all 5113 gene trees derived from each orthology cluster described earlier. Before species tree inference, we applied an additional branch trimming process to remove duplicated taxa from individual gene trees following Yang & Smith (2014). At each node where two descendant clades contain overlapping taxa, the branch with the smaller number of taxa was pruned. These pruned gene trees were subsequently used in ASTRAL for species tree estimation. We additionally processed the 100 BS trees for each gene using the same pruning method described earlier. These BS trees are used in ASTRAL analyses to conduct BS replicates for species tree estimation. Molecular divergence time estimates were subsequently inferred for the species tree using the penalized likelihood method described earlier using a concatenated sequence matrix derived from all 40 genes containing at least 34 in-group taxa.

### Overview of the methods applied to infer WGD

We utilized three lines of inference described in the following three subsections to identify, locate, and determine the age of WGDs in Malpighiales. Each of these methods has been commonly applied in angiosperms and elsewhere, and includes distribution of synonymous substitutions per synonymous site ( $K_s$ ) among paralogs (Cui *et al.*, 2006; Barker *et al.*, 2008), phylogenetic (gene tree) reconciliation (Jiao *et al.*, 2011; Li *et al.*,

2015), and a likelihood-based gene-count method (Rabier *et al.*, 2014; Tiley *et al.*, 2016).

### $K_s$ -based method for WGD identification

This method identifies a proliferation of duplicated genes from WGDs under the assumption that synonymous substitutions between duplicate genes accrue at a relatively constant rate. Here, each species was subjected to a reciprocal BLAST search to identify putative paralogous gene pairs in their protein coding sequences. Paralogous pairs were identified as sequences that demonstrated 40% sequence similarity over at least 300 bp from a discontinuous MEGABLAST search (Zhang *et al.*, 2000; Barker *et al.*, 2008). Each paralogous protein sequence pair was aligned using MAFFT (Katoh & Standley, 2013) and then back-translated to their coding sequences using PAL2NAL (Suyama *et al.*, 2006). All sites containing gaps were removed from the alignment.  $K_s$  values for each duplicate pair were calculated using the ML method implemented in codeml of the PAML package (Yang, 1997) under the F3 × 4 model (Goldman & Yang, 1994). To infer WGDs from the  $K_s$  distribution, we employed the one sample Kolmogorov–Smirnov goodness-of-fit test followed by 100 BS resampling to assess statistical confidence (Cui *et al.*, 2006). A significant  $P$  value (<0.05) rejected the null hypothesis of a birth–death process of gene duplication, thus supporting evidence of WGD. Because peaks produced by paleopolyploidy are expected to be approximately Gaussian (Blanc & Wolfe, 2004; Schlueter *et al.*, 2004), we applied the expectation–maximization (EM) algorithm to fit mixtures of Gaussian distributions to our data using the normalmixEM() function in the R package MIXTOOLS (Benaglia *et al.*, 2009). Estimated mean peak values for each taxon are reported in Figs S3, S4. Alternative splicing can confound the signal of WGD using the  $K_s$  method (Barker *et al.*, 2008). To alleviate this concern, sites containing gaps were removed from paralog alignments; thus, transcript isoforms generated from alternative splicing will receive a  $K_s$  value of zero. All pairs with a  $K_s$  value of less than 0.001, which would include these transcript isoforms as well as recent tandem duplications, were discarded and not considered in the  $K_s$  distribution. Finally, to further explore the sensitivity of misassemblies and tandem gene duplications on WGD identification, we conducted the same  $K_s$  analysis but applied a more stringent 0.95 threshold in the CD-HIT-EST analysis (-c 0.95 -r 10).

### Placing and dating WGDs using phylogenetic reconciliation and molecular divergence time estimation

We applied a phylogenetic approach to identify more precise placements of WGDs and to determine the approximate age of these events. We first reconciled each orthology tree to the species tree under the duplication–transfer–loss model in Notung v2.9 (Chen *et al.*, 2000). Here, total numbers of gene duplications inferred from well-supported gene tree nodes (> 70 BS) are summarized onto the species tree. For each branch in the species tree, we calculated the percentage of genes duplicated along that branch (total number of inferred gene duplications along the

branch/total number of genes containing at least one descended copy on that branch (i.e., from both single and duplicated gene copies)). The range of duplicated genes along branches where WGD was inferred was 10.0–84.3% (Table S4). Our threshold percentage for identifying a WGD was 10.0%, which is the lowest percentage for a terminal WGD as determined by our  $K_s$  analysis (in genus *Bhesa*). Terminal WGDs are likely to exhibit a higher percentage of retained duplicated genes, and thus represent a more conservative filter for WGD assessment. This threshold is also well above the percentage identified from fully sequenced genomes and transcriptomes that do not exhibit recent WGDs. These fully sequenced genomes and transcriptomes instead show maximally only 1.2–2.2% of duplicated genes where tandem duplications, not WGDs, have been inferred (calculated from *Chrysobalanus* and *Jatropha*, Table S4). Together, this represents a reasonable approach for estimating WGDs in Malpighiales.

Following our phylogenetic localization of WGDs, we applied a customized R script to extract the divergence times from our r8s analyses to summarize the age of gene duplications along each branch of the species tree. Only ages of nodes supported with > 70 BS were used. The inferred distribution of divergence times was fitted to a mixture of Gaussian models using the R package mixtools as described earlier to estimate mean age of each WGD. In several cases, the best model inferred two WGDs along the same branch. These cases were independently supported by our phylogenetic analysis but with smaller percentages of duplications, presumably due to gene loss or missing data (Table S4). Estimated mean peak values for each taxon are reported in Table S4 and Fig. S5. Confidence intervals for mean age of each WGD were estimated using the 100 bootstrapped trees for each gene as outlined earlier (Table S4). To further explore the impact of root age on divergence time estimation using this penalized likelihood method, we fixed the root age to be 125 Ma, which is the maximum age of crown group eudicots based on the fossil record (Hughes & McDougall, 1990; Doyle & Hotton, 1991; Magallón *et al.*, 2015). Moreover, this age is over 20 Ma older than the optimal age estimates for crown group Malpighiales (103.6 Ma) as inferred by Magallón *et al.* (2015). We applied the same set of minimum age constraints described earlier to conduct the divergence time estimation (Table S4).

### Gene-count-based method for confirmation of WGDs

To further assess statistical confidence for the 24 putative WGDs inferred from the  $K_s$  method and using the aforementioned phylogenetic approach, we applied the ML method to test the number and placement of WGDs using gene count data (Rabier *et al.*, 2014). This method estimates the likelihood of prespecified putative WGDs on a phylogeny using a gene count matrix summarized across all orthologous genes. It is advantageous because it suffers less from false-positive rates due to tandem duplication and assembly error (Rabier *et al.*, 2014), which could create an artificial signal of polyploidization. We first tested the utility of this method by examining three independent WGDs previously identified from fully sequenced genomes using synteny analyses.

These three WGDs were identified in *Populus* and *Salix* (Hanley *et al.*, 2006; Tuskan *et al.*, 2006), in *Hevea* and *Manihot* (Bredeson *et al.*, 2016; Tang *et al.*, 2016), and in *Linum* (Wang Z. *et al.*, 2012). Filtering of the gene count matrix to avoid missing data is critical for this method, which may otherwise lead to biased estimates (Rabier *et al.*, 2014). To accomplish this, our dated species trees were first pruned to contain only species with fully sampled genomes, including these five species, plus *Jatropha* and *Ricinus* (where WGDs have not been previously detected) and *Vitis* (as outgroup). Next, a gene count matrix was summarized for all species across all ortholog trees (Fig. S6). We further conditioned the data matrix for all gene families to contain one or more gene copies descended from the branch along which the WGD was tested. Along these lines, gene families that were missing in the taxa affected by WGD were removed in each test. The conditional likelihoods were subsequently estimated for models with and without the WGD of interest using a prior geometric mean of 1.5 (Tiley *et al.*, 2016). After convergence of the likelihood scores for all runs, we performed a series of likelihood ratio tests (LRTs) to determine the significance of individual WGDs. All three previously identified WGDs were successfully identified with confidence (LRT statistic  $\gg 9.55$ , probability of type I error  $\ll 0.001$ ). In addition, *Ricinus* and *Jatropha* showed no evidence of WGD (Table S4), suggesting a low false-positive rate of this method.

We tested the remaining WGDs by sequentially adding species associated with each WGD to the seven-species phylogeny. In each case, we added all of the species from which a single WGD to be tested were descended and tested for a WGD along the added branch of interest. We generated the conditioned gene count matrix containing varying numbers of gene families as described earlier (Table S4).

### Clustering of WGDs in time

To assess whether WGDs were clustered in time, we tested for the optimum number of clusters in age distribution using the finite Gaussian mixture modeling in R package MCLUST (Fraley & Raftery, 2002). The EM algorithm was used for mixture estimation and the Bayesian information criterion (BIC) was used for comprehensive clustering. In order to assess the confidence of the clustering, we conducted the same analysis on WGD age distributions derived from 100 BS replicates. We conducted this Gaussian mixture modeling for age estimations derived from r8s analyses with fixed root age at 109 Ma as well as age estimations with fixed root age at 125 Ma. To further explore the clustering time of the more ancient WGDs, we excluded WGDs younger than 20 Ma and conducted the same Gaussian mixture modeling analysis.

### Synteny-based assessment of WGD in *Linum*

Synteny analysis serves as the gold standard for inferring WGDs, but it is only amenable to the mostly completely assembled genomes. Our phylogenetic and  $K_s$  approach identified two WGDs in *Linum*. Here, we subsampled gene families containing

three or four gene copies of *Linum* consistent with two duplications. The most closely related paralogous gene pairs of *Linum* were expected to arise from the most recent WGD, while their relationship with the remaining copy(ies) arose from the more ancient WGD (Fig. S7). We then mapped these paralogs onto the genome using MCScanX\_h (Wang Y. *et al.*, 2012) and visualized the result using RCircos (Zhang *et al.*, 2013).

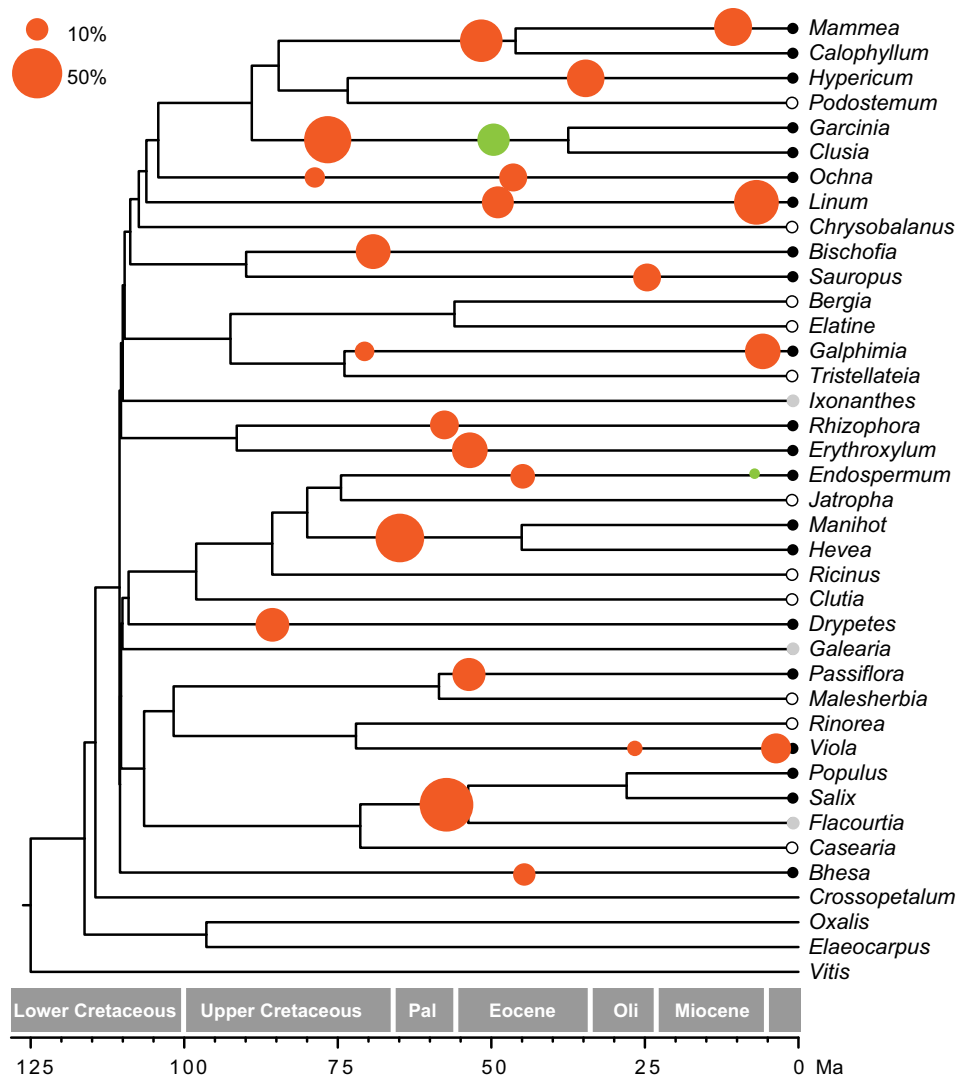
### Results

Our final data set included 36 in-group taxa derived from eight genomes and 28 transcriptomes (15 newly acquired for this study) plus six outgroup species (Tables S1–S3). Summary statistics of the transcriptome assemblies are reported in Table S5. The taxon sampling includes 21 traditionally recognized families in the order, thus representing the broad outline of phylodiversity within Malpighiales (*sensu* Angiosperm Phylogeny Group IV; Wurdack & Davis, 2009; Chase *et al.*, 2016).

Our analyses identified a total of 22–24 WGDs broadly distributed across the Malpighiales phylogeny (Fig. 1). In nearly all cases these events are corroborated by all three methods. Our  $K_s$  analysis identified WGD in 22 species regardless of the sequence similarity threshold (Figs 1, S3, S4 for the 0.99 CD-HIT-EST threshold; Fig S8 for the 0.95 CD-HIT-EST threshold); our phylogenomic reconciliation analysis identified 24 WGDs (Table S4); and 22 WGDs were verified with the likelihood-based gene-count method (LRT statistic  $> 9.55$ ; Table S4). Moreover, these results are robust to data quality and phylogenetic uncertainty (by applying our BS resampling procedure described earlier).

We additionally analyzed a reduced data set containing only completely sequenced genomes and high-quality transcriptomes to further verify these results. Even with this more conservative data set, we still identified 22 WGDs using all three methods (Table S6). Three of the WGDs we identify validate those from previous studies in the common ancestor of *Populus* (Tuskan *et al.*, 2006) and *Salix* (Sterck *et al.*, 2005; Hanley *et al.*, 2006), in the common ancestor of *Manihot* (Bredeson *et al.*, 2016) and *Hevea* (Tang *et al.*, 2016), and in *Linum* (Wang Z. *et al.*, 2012). In addition, recent analyses have inferred two independent WGDs in *Linum* using  $K_s$  distributions (Wang Z. *et al.*, 2012; Sveinsson *et al.*, 2014). These events, however, have not previously been corroborated simultaneously due to the limited power of this method. Our reconciliation and gene-count method, by contrast, identifies these two duplications in the lineage leading to *Linum*. Furthermore, we verified these two independent WGDs using a phylogeny-guided synteny analysis (Jiao *et al.*, 2014) based on 879 gene families containing three or four gene copies in *Linum* (Fig. S7). Here, we identified 15 syntenic regions across large scaffolds reflecting the four-parted paralogous relationship created by two independent WGDs in the *Linum* lineage.

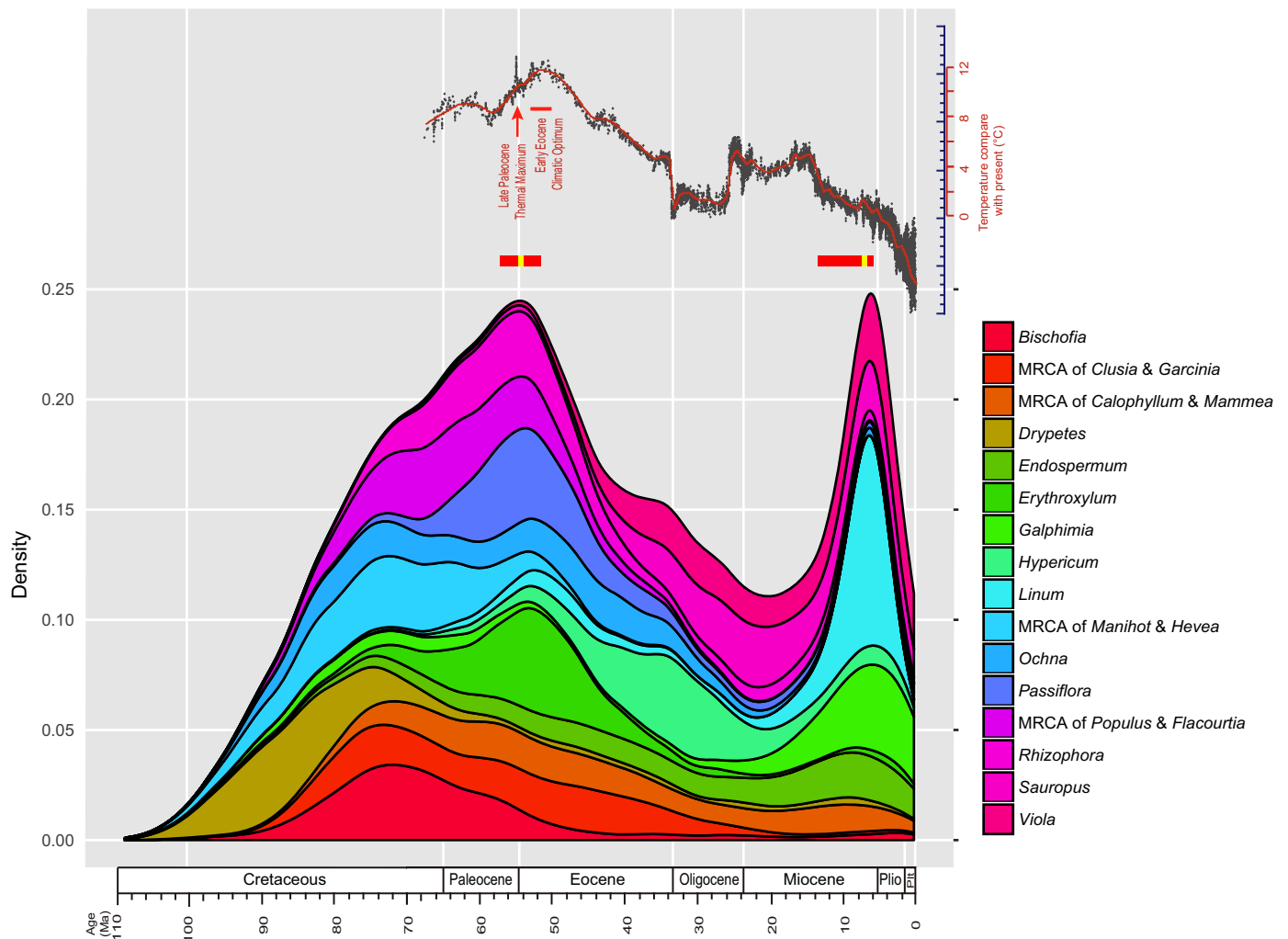
Our divergence time estimates indicated that the timing of WGDs range broadly from 3.2–85.0 Ma (Fig. 2; Table S4). Surprisingly, however, these events are not randomly distributed in time. Instead, our Gaussian mixture model (Fraley



**Fig. 1** Phylogenetic distribution of whole-genome duplications (WGDs) in Malpighiales. Species tree of Malpighiales inferred from 5113 gene trees using a summary coalescent method. WGDs identified from  $K_s$  analysis are illustrated with solid black dots on the terminal branches of corresponding species; species indicated with gray dots have transcriptomes that are potentially insufficient for adequate assessment of WGD using the  $K_s$  method. Two *Salix* species are collapsed into one terminal branch for simplicity. Circles illustrated along branches are dated WGDs (divergence time of parental genomes) from our phylogenomic reconciliation analysis. The radius of each circle is proportional to the percentage of orthologous genes supporting the WGD as determined from phylogenomic reconciliation (scale, top left). Solid red circles are significant WGDs as confirmed by our gene-count analysis; solid green circles indicate WGDs that do not receive significant support using the gene-count method. The photographs on the right, from top to bottom (beginning with the left column), are *Garcinia*, *Ochna*, *Tristellateia*, *Manihot*, *Viola*, *Clusia*, *Linum*, *Rhizophora*, *Passiflora*, and *Salix* and are representatives of the major Malpighiales clades sampled for investigating WGDs.

& Raftery, 2002) indicates that the inferred ages of WGDs display a bimodal distribution (BIC  $-225.05$ , compared with BIC  $-227.30$  for the univariate normal distribution) with peaks at the Eocene–Paleocene (mean age 53.89 Ma) and late Miocene (mean age 7.39 Ma). Our assessment using BS replicates provides confident statistical support for this interpretation: 98% of the replicates support a bimodal distribution of WGDs with the mean age of each cluster ranging from 6.06 to 13.78 Ma and from 52.26 to 57.79 Ma. In our case, an overwhelming number of WGDs ( $n=19$ ) occur during the older, early Eocene time period, vs only five during the more recent late Miocene period. When we excluded WGDs with

mean age younger than 20 Ma, the 19 WGDs were estimated to cluster at 54.21 Ma according to the Gaussian mixture model analysis. The ages of these WGDs are also robust to alternative age constraints. When we fixed the root age of each gene tree at 125 Ma, age estimations of most WGDs were 0–9.9 Ma older than previous estimations except for the recent WGD in *Endospermum*, which is 0.3 Ma younger (Table S4). However, regardless of these older ages, our Gaussian mixture model still supported a clustering age of 56.8 Ma (54.81–60.42 Ma in the 100 BS replicates), which remains closer to the Eocene–Paleocene transition than to the KT boundary.



**Fig. 2** Age distributions of whole genome duplications (WGDs) among clades of Malpighiales. Density of divergence time of duplicated genes by taxa are plotted against time in millions of years ago. The various colors refer to different clades exhibiting WGDs (*Bhesa* excluded for readability). Zachos *et al.* (2001) curve of global temperature fluctuations during the Cenozoic is redrawn at the top. Mean ages of Gaussian mixture model and variations estimated from bootstrap replicates are indicated by yellow and red bars respectively below the Zachos curve. MRCA, most recent common ancestor.

## Discussion

### Massive WGDs in Malpighiales

The WGDs we identified are broadly distributed across 18 branches of the Malpighiales phylogeny. All of these WGDs are inferred in the crown groups of major clades where phylogenetic uncertainty is minimum. As a result, although the backbone of the phylogeny in Malpighiales is still challenging to resolve (species tree with BS support Fig. S9), the phylogenetic placements of these WGDs are clear. Moreover, we identified the same set of WGDs in our full data set compared with the more stringent, filtered data set (cf. *Bhesa*, which we excluded in the more conservative analysis). This result indicates that low-quality transcriptomes do not have a major impact on our assessment of WGD in Malpighiales.

Interestingly, these WGDs are commonly associated with the most diverse clades in the order, including in the clusioids, ochnooids, euphorbioids, phyllanthoids, violets and passion

flowers. Additionally, we note that some species-poor clades show no evidence of duplication (e.g., *Malesherbia*, *Rinorea*, and *Elatinaceae*, among others; Fig. 1), despite having species-rich sister clades. This lends tentative support to suggestions that WGDs may fuel species diversification (Tank *et al.*, 2015), possibly via the establishment of reproductive barriers (Husband *et al.*, 2016). However, other studies have concluded that, although polyploidization is important to cladogenesis in plants, it likely does not enhance species diversification rates (Wood *et al.*, 2009; Estep *et al.*, 2014). We cannot adequately address this question with existing data (Kellogg, 2016); but regardless, our analyses set the stage for establishing the finer scale taxon sampling necessary to pinpoint these events to clarify the association between WGDs and the tempo of diversification in Malpighiales. Namely, do WGDs precede prolific diversification of Malpighiales subclasses?

One WGD requires more detailed exploration. In our phylogenetic reconciliation and gene-count analysis, a WGD is inferred to predate the common ancestor of *Populus*, *Salix*, and

*Flacourtia* (the  $K_s$  analysis is inconclusive for *Flacourtia*; Figs 1, S4). Chromosome count data, however, *do not* support such an early WGD. Instead, the chromosome number of *Populus* and *Salix* are approximately twice that of *Flacourtia* ( $2n=38$ , vs  $2n=20$  or  $22$  in *Flacourtia*; IPCN Chromosome Reports, <http://www.tropicos.org/Project/IPCN>), suggesting that this WGD event likely occurred more recently and is thus restricted to the common ancestor of *Populus* and *Salix*. A similar discrepancy in yeast involves the identification of an older WGD using phylogenetic reconciliation vs a more recent WGD inferred using a gold standard synteny-guided genome comparison (Scannell *et al.*, 2007; Marcet-Houben & Gabaldón, 2015). Here, the authors provide reasonable evidence that the older WGD is spurious and confounded by an allopolyploidization event resulting from hybridization. The nature of this allopolyploidization resulted in a deeper, yet spurious, phylogenetic placement of the WGD. It is important to recognize here that gene-tree data are limited in some respects since they provide an estimate of the divergence times of the parental diploid genomes, but they are less conclusive around exactly when the hybridization and polyploidization event occurred (Gaut & Doebley, 1997; Doyle & Egan, 2010); dates in Table S4 are thus likely older than the actual polyploidization events. In light of these results, a plausible hypothesis is that the WGD shared by *Populus* and *Salix* results from an allopolyploidization in which an ancestor of the *Flacourtia* lineage served as one parental lineage. Testing and evaluating this hypothesis remains a challenge (Goulet *et al.*, 2017), and is an obvious avenue for future research. Regardless, we do not anticipate this phenomenon to be a pervasive problem for our analysis given that the origin of viable polyploids derived from widely disparate phylogenetic lineages appears to be rare, and thus not likely to greatly influence our placement and dating of the large number of WGDs identified here.

These results further point toward a propensity for pervasive and widespread WGDs in angiosperms. Recent and ongoing investigations incorporating vast nuclear genomic data and extended taxon sampling indicate that other similarly diverse clades, including Asteraceae (Barker *et al.*, 2008; Huang *et al.*, 2016), Poaceae (Estep *et al.*, 2014), and Caryophyllales (Yang *et al.*, 2015, 2017), also show histories characterized by prolific WGDs. Collectively, these results suggest that Malpighiales and other previously examined clades represent a more pervasive pattern of WGDs characteristic of possibly hundreds of major angiosperm clades (and extending to other vascular plant clades, including gymnosperms and ferns; Wood *et al.*, 2009; Li *et al.*, 2015). Further investigation is required to address this question more broadly, but this seems a plausible hypothesis in light of recent findings.

#### Timing of WGDs coincides with events of climate upheaval

WGDs in Malpighiales show a bimodal distribution through time, which is supported by 98% of our BS replicates. The majority of the WGDs were identified to cluster around the Eocene–Paleocene transition (mean age 53.89 Ma), during which time the planet was warmer and wetter than any other period in the Cenozoic. The age

of this older peak is estimated to range between 52.26 and 57.79 Ma based on our BS replicates, which falls entirely within the prolonged warming trend from the late Paleocene through the early Eocene (58–50 Ma, Zachos *et al.*, 2008). Moreover, the late Paleocene–early Eocene age of this peak is also robust when we excluded WGDs younger than 20 Ma (mean age 54.21 Ma) or apply an older root age of 125 Ma consistent with the oldest eudicots (mean age 56.8 Ma). Previously, WGD has been hypothesized to buffer plants through episodes of major global and climatic upheavals (Fawcett *et al.*, 2009). Studies have identified WGDs associated with the earlier KT boundary (*c.* 65 Ma) when a large meteor impacted off the Yucatán Peninsula disrupting the global climate, precipitating a major reorganization of the terrestrial biota (Fawcett *et al.*, 2009; Vanneste *et al.*, 2014). Similarly, the late Miocene–early Pliocene (*c.* 10–5 Ma) has been implicated as another period of climatic instability when WGDs were pervasive (Estep *et al.*, 2014). The expansion of  $C_4$  grassland as a result of widespread global aridification (Cerling *et al.*, 1997; Arakaki *et al.*, 2011) has, in particular, been inferred to be correlated with numerous polyploidizations in grasses, which are among the most important members of these arid and cooler habitats that bear their namesake; that is, grassland and steppe biomes (Estep *et al.*, 2014).

By contrast, relatively little is known about WGDs during the Eocene. This is surprising, because the Eocene–Paleocene transition is associated with an extended and prolonged period of intense warming. Most notably, the Paleocene–Eocene Thermal Maximum (56 Ma) and the subsequent Eocene Climatic Optimum (49 Ma) constitute the warmest and most humid period during the Cenozoic. During this time, mean global temperatures increased by 5–10°C due to massive release of  $^{13}C$ -depleted carbon (Pagani *et al.*, 2006; Zeebe *et al.*, 2009). This dramatic climate upheaval is thought to have stimulated profound reshuffling of the terrestrial biome, spurring plant migrations, extensive species turnover, and accelerated species diversification in numerous plant and animal clades (Clyde & Gingerich, 1998; Wilf, 2000; Bowen *et al.*, 2002; Wing *et al.*, 2005; McKenna & Farrell, 2006; Ramírez *et al.*, 2007; Schuettpeiz & Pryer, 2009; Jaramillo *et al.*, 2010). Our results establish a record of at least 19 WGDs during this period, suggesting a role in adaptation during Paleocene–Eocene warming. We hypothesize that this may be common for predominantly tropical groups, like Malpighiales (Davis *et al.*, 2005), which are likely much more impacted by warming given the relatively tight thermal tolerances exhibited by many such groups (Janzen, 1967; Tewksbury *et al.*, 2008; Wright *et al.*, 2009).

What may have stimulated interactions that facilitated increased polyploid formation during the warmer, wetter period of the Eocene beyond the generally increased rates of angiosperm diversification during this window of time? One hypothesis, coined the ‘neutral’ process (Van de Peer *et al.*, 2017), posits that these sorts of upheavals increase the formation of unreduced gametes and therefore result in an excess of polyploids. It is widely appreciated that external stimuli, temperature in particular, has a pronounced effect on unreduced gamete formation (De Storme & Geelen, 2014). In this case, it appears that both high and low temperatures can promote unreduced gametes in various taxa, as demonstrated in *Arabidopsis* (De Storme *et al.*, 2012) and

some roses (Pécricx *et al.*, 2011) respectively. In addition, a recent study in *Arabidopsis thaliana* demonstrated that supernumerary sperm fusion could generate viable polyploid offspring that exhibited vegetative vigor (Nakel *et al.*, 2017). Thus, the extensive early Eocene warming and even the Miocene aridification might have significantly increased unreduced gametes, perhaps contributing to enhanced polyploid formation. This is supported by evidence of increased levels of unreduced gametes in gymnosperms and lycophytes during comparable upheavals, including during the Triassic–Jurassic (Kürschner *et al.*, 2013) and Permian–Triassic (Visscher *et al.*, 2004; Foster & Afonin, 2005) transitions.

Although such polyploidizations might have initially arisen more neutrally, it is also possible that polyploids were adapted for survival in these changing landscapes (the ‘adaptive’ processes; Van de Peer *et al.*, 2017). Of course, both neutral and adaptive processes can contribute simultaneously to the formation and establishment of polyploids. Polyploids are often viewed as evolutionary dead ends owing to their small population sizes, relatively restricted distributions, high extinction risks, and seemingly sparser representation in the deep angiosperm phylogeny (Stebbins, 1970; Comai, 2005; Mayrose *et al.*, 2011). However, these may not apply under less stable environments, such as during major climatic upheavals, when polyploids may outperform their diploid progenitors (Van de Peer *et al.*, 2017). It has been hypothesized that such genomic novelty and epigenetic repatterning may result in phenotypic variability, including variants that confer selective advantages in stressful conditions (Wendel, 2000; Comai, 2005; Madlung, 2013; Van de Peer *et al.*, 2017). In particular, the advantages of especially allopolyploids include altered gene expression leading to hybrid vigor and increased genetic variation (Comai, 2005; Lynch, 2007). Along these lines, polyploids have been reported to be more frequent at higher latitudes and in xeric environments, which may mimic such upheavals (Löve & Löve, 1949; Löve, 1953; Hanelt, 1966; Brochmann *et al.*, 2004). Polyploids also occur with greater frequency among invasive plants, which commonly become established on disturbed grounds (Prentis *et al.*, 2008; te Beest *et al.*, 2011). In support of this argument, Brochmann *et al.* (2004) reported an unexpected overabundance of recently formed polyploids in newly deglaciated areas in the Arctic. Additionally, megafossil plants from Wyoming, USA, indicate that the dynamics of plant community assembly after dramatic warming during the Paleocene–Eocene Thermal Maximum is very similar to late and postglacial floras (Wing *et al.*, 2005), suggesting that observations for the Arctic may represent similar responses to warmer and wetter periods during the Eocene transition. Regardless of these competing ideas, the striking propensity and clustered distribution of WGDs in time strengthens the hypothesis that polyploidization may be an important means of lineage persistence during episodes of major global change.

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## Author contributions

L.C. and C.C.D. designed the research; L.C. analyzed the data; Z.X. collected the data; L.C. and C.C.D. wrote the initial paper draft with input from A.M.A., M.S., J.S.R. and L.L.

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## Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1** BUSCO assessment of transcriptome and genomic completeness of 42 sampled species.

**Fig. S2** Bioinformatic pipeline depicting transcriptome assembly, homology and orthology inferences, and three methods for WGD identification, placement, and dating.

**Fig. S3** Histograms of the synonymous substitutions per synonymous site ( $K_s$ ) of duplicated gene pairs among 22 Malpighiales taxa showing WGDs.

**Fig. S4** Histograms of the synonymous substitutions per synonymous site ( $K_s$ ) of duplicated gene pairs among 11 Malpighiales taxa where WGD is absent (or inconclusive in the case of three taxa).

**Fig. S5** Histograms depicting divergence time estimations inferred using penalized likelihood for 22 species that exhibited WGDs (summarized in Fig. S3).

**Fig. S6.** Gene count matrix for 5113 ortholog groups of 36 Malpighiales species.

**Fig. S7** Phylogeny-guided synteny analyses demonstrate successive WGDs in *Linum*.

**Fig. S8** Histograms of the synonymous substitutions per synonymous site ( $K_s$ ) of duplicated gene pairs among 22 Malpighiales taxa showing WGDs.

**Fig. S9** Cladogram of 50% majority-rule bootstrap tree of Malpighiales inferred from 5113 gene trees using ASTRAL.

**Table S1** Voucher and GenBank information for 15 species in Malpighiales used for *de novo* transcriptome assembly

**Table S2** Taxa sampled from the OneKP data set and the corresponding OneKP library ID

**Table S3** Taxa sampled with complete genomes and corresponding reference

**Table S4** Whole genome duplications (WGDs) in Malpighiales identified with complete taxon sampling, including their phylogenetic placement, percentage of gene families supporting gene duplication, log likelihood, and age estimations of each WGD

**Table S5** Summary statistics for the transcriptome assembly in Malpighiales

**Table S6** Whole genome duplications (WGDs) in Malpighiales identified with more conservative taxon sampling, including their phylogenetic placement, percentage of gene families supporting gene duplication, log likelihood, and age estimations of each WGD

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