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Genome of wild olive and the evolution of oil biosynthesis

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Here we present the genome sequence and annotation of the wild olive tree (*Olea europaea* var. *sylvestris*), called oleaster, which is considered an ancestor of cultivated olive trees. More than 50,000 protein-coding genes were predicted, a majority of which could be anchored to 23 pseudochromosomes obtained through a newly constructed genetic map. The oleaster genome contains signatures of two Oleaceae lineage-specific paleopolyploidy events, dated at ~28 and ~59 Mya. These events contributed to the expansion and neofunctionalization of genes and gene families that play important roles in oil biosynthesis. The functional divergence of oil biosynthesis pathway genes, such as *FAD2*, *SACPD*, *EAR*, and *ACPT*, following duplication, has been responsible for the differential accumulation of oleic and linoleic acids produced in olive compared with sesame, a closely related oil crop. Duplicated oleaster *FAD2* genes are regulated by an siRNA derived from a transposable element-rich region, leading to suppressed levels of *FAD2* gene expression. Additionally, neofunctionalization of members of the *SACPD* gene family has led to increased expression of *SACPD2*, 3, 5, and 7, consequently resulting in an increased desaturation of steric acid. Taken together, decreased *FAD2* expression and increased *SACPD* expression likely explain the accumulation of exceptionally high levels of oleic acid in olive. The oleaster genome thus provides important insights into the evolution of oil biosynthesis and will be a valuable resource for oil crop genomics.

oil crop | whole-genome duplication | siRNA regulation | fatty-acid biosynthesis | polyunsaturated fatty-acid pathway

As a symbol of peace, fertility, health, and longevity, the olive tree (*Olea europaea* L.) is a socioeconomically important oil crop that is widely grown in the Mediterranean Basin. Belonging to the Oleaceae family (order Lamiales), it can biosynthesize essential unsaturated fatty acids and other important secondary metabolites, such as vitamins and phenolic compounds (1). The olive tree is a diploid (2n = 46) allogamous crop that can be vegetatively propagated and live for thousands of years (2). Paleobotanical evidence suggests that olive oil was already produced in the Bronze Age (3). It has been thought that cultivated varieties were derived from the wild olive tree, called oleaster (*O. europaea* var. *sylvestris*), in Asia Minor, which then spread to Greece (4). Nevertheless, the exact domestication history of the olive tree is unknown (5). Because of their longevity, oleaster trees might even be related to Neolithic olive tree ancestors (2). Although the natural long generation time

of olive trees has traditionally hindered breeding in this species, there are a few breeding programs involving sexual crosses that have generated interesting varieties for novel uses, like “Chiquitita,” specifically selected for high-density hedgerow orchards (6).

The olive is tightly associated with the Mediterranean cuisine. However, its consumption also spread to America (United States,

Significance

We sequenced the genome and transcriptomes of the wild olive (oleaster). More than 50,000 genes were predicted, and evidence was found for two relatively recent whole-genome duplication events, dated at approximately 28 and 59 Mya. Whole-genome sequencing, as well as gene expression studies, provide further insights into the evolution of oil biosynthesis, and will aid future studies aimed at further increasing the production of olive oil, which is a key ingredient of the healthy Mediterranean diet and has been granted a qualified health claim by the US Food and Drug Administration.

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Reviewers: R.M., University of Illinois at Urbana–Champaign; and K.S., MPI for Plant Breeding Research.

The authors declare no conflict of interest.

Data deposition: The oleaster genome assembly has been deposited in the GenBank database, <https://www.ncbi.nlm.nih.gov/genbank> (accession no. MSRW00000000); BioProject record ID PRJNA350614. Transcriptome datasets were deposited in the National Center for Biotechnology Information Sequence Read Archive, <https://www.ncbi.nlm.nih.gov/sra> (accession nos. SRR4473639, SRR4473641, SRR44742, SRR4473643, SRR4473644, SRR4473645, SRR4473646, and SRR4473647). The genome and annotation files were uploaded to Online Resource for Community Annotation of Eukaryotes (ORCAE), bioinformatics.psb.ugent.be/orcae; Phytozome, <https://phytozome.jgi.doe.gov>; and the olive genome consortium Web site, olivegenome.org.

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Mexico, Brazil, Argentina, and Peru), Asia (China and India), and Australia. Aside from cultural reasons, this expansion was mainly because of the recognition of the beneficial dietetic properties of olive oil as a source of healthy fatty acids and micronutrients (e.g., antioxidants such as phenolic compounds, including vitamin E). In

fact, olive oil has been granted a qualified health claim as reducing the incidence of cardiovascular disease (i.e., coronary heart disease) (7) by the US Food and Drug Administration (FDA; docket no. 2003Q-0559). As such, it represents the third FDA-approved claim for conventional foods, after nuts and omega-3 fatty acids.

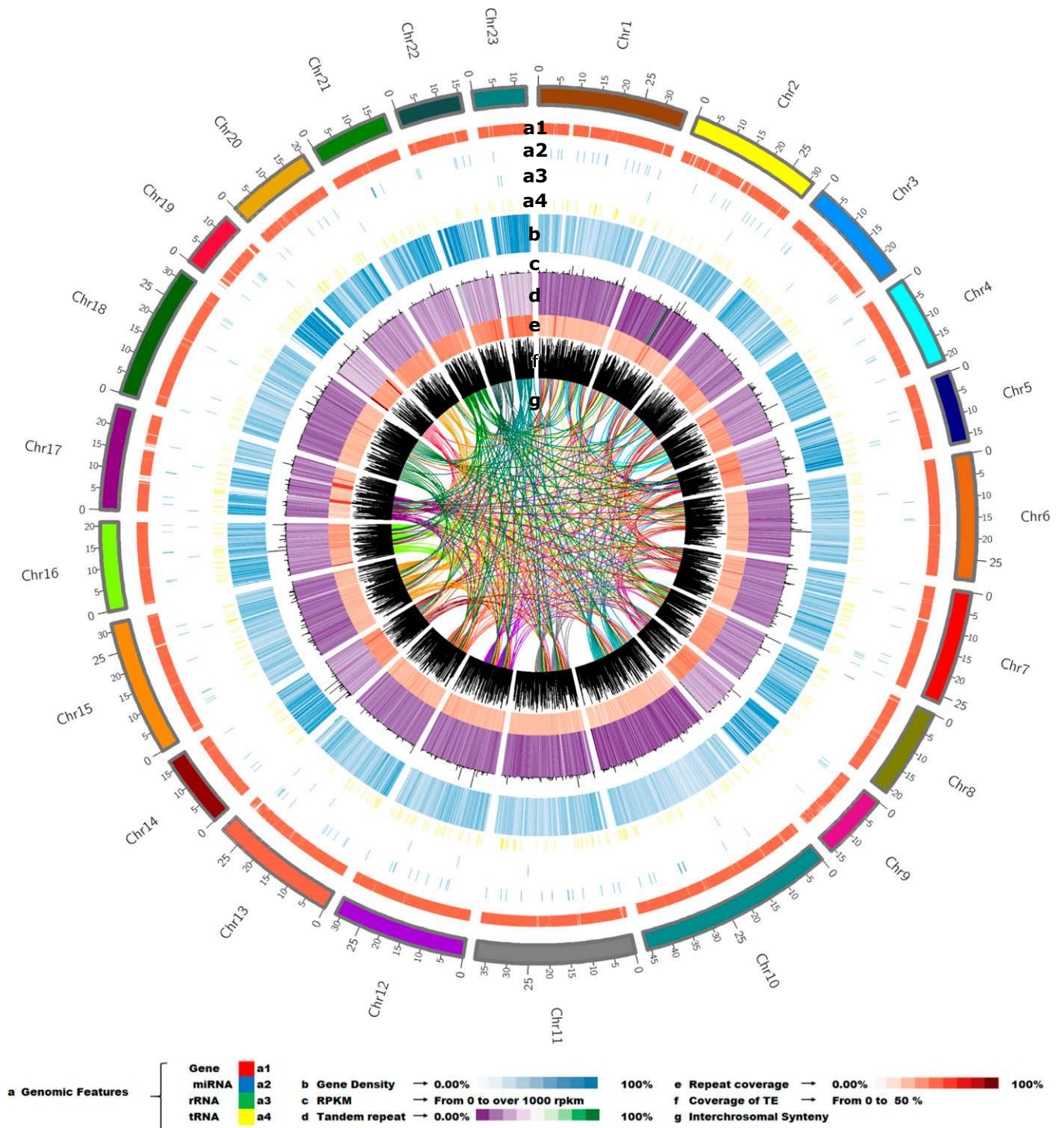


Fig. 1. The genomic landscape of oleaster. The outer layer represents the karyotype ideogram (colored blocks), with minor and major tick marks labeling each 5 Mbp and 25 Mbp, respectively. Genome features across the 23 chromosomes (distinct characters shown as different colors, as indicated in the legend). Gene density per megabase pair. Gene expression patterns in average RPKM (range of RPKM values plotted from 0 to >1,000). Tandem duplication density per megabase pair. Percentage heat map of repeat coverage per megabase pair. Percentage of TEs per megabase pair (ranges of values plotted from 0 to >50). Inner circular representation shows interchromosomal synteny.

Table 1. Statistics of the wild olive tree genome and assembly

Features	Statistics
Genome	
Size (n, Gbp)	1.48
Karyotype (chromosomes, 2n)	46 = 2n
GC content, % (with/without Ns)	36.8/38.8
High-copy repeat no.	
No. LTR/Gypsy and Copia	1,182,454
No. LINE	43,834
No. DNA TE	219,901
No. unknown	42,630
Gene	50,684
Assembly	
No. scaffold >100 bp/>1 kbp	2,356,597/42,843
N50 > 100 bp/>1 kbp	228.62/364.6

N50, shortest sequence length at 50% of the genome assembly.

Moreover, olive tree products and byproducts are also used for pharmaceutical and cosmetic purposes.

Traditionally, olive oil is obtained by pressing olive fruits. Olive fruits consist of 20–30% (wt/wt) oil, 17% cellulose, 4% carbohydrates, 2% protein, and 0.1% micronutrients (1), with the rest (46.9–56.9%) being water. Polyols (mannitol) and oligosaccharides (raffinose and stachyose) are synthesized in olive tree leaves, being further exported with sucrose into the fruits, for general metabolism and as precursors of olive oil biosynthesis (8). Starting from a carbon source such as sucrose, long-chain fatty acids are synthesized, modified, and degraded by the activity of enzymes, including fatty-acid synthases, elongases, desaturases, and carboxylases (9). Fatty acids are the major constituent of triacylglycerols (TAGs). In olive oil, TAGs are mostly composed of monounsaturated oleic acid (C18:1; ~75% of all TAGs), followed by saturated palmitic acid (C16; ~13.5%), polyunsaturated linoleic acid (c18:2 ω -6; ~5.5%), and α -linolenic acid (c18:3 ω -3; ~0.75%) (10).

Results

Assembly of the Oleaster Genome. The wild olive tree genome was shotgun-sequenced (220 \times coverage), generating 515.7 Gbp of data (*SI Appendix, Table S1*). SOAPdenovo (11) was used to assemble the sequence reads, which resulted in a draft genome assembly of 1.48 Gbp, with the scaffold shortest sequence length at 50% of the genome of 228 kbp (*SI Appendix, Table S3*), which is in agreement with genome size estimations from flow cytometry (*SI Appendix, Fig. S1*) and *k*-mer analysis (~1.46 Gbp; *SI Appendix, Fig. S2A and Table S2*). By using a newly constructed genetic map, 50% of sequences longer than 1 kbp (~572 Mbp) could be anchored into 23 linkage groups (Fig. 1 and Tables 1 and 2).

Genome Annotation. The annotation of the oleaster genome was carried out by combining three different approaches, namely ab initio prediction, homology-based prediction, and transcriptome mapping (Fig. 1 and Tables 1 and 2). Approximately 51% of the genome assembly was found to be composed of repetitive DNA (Fig. 1), which is less than what was found for the draft genome of a recently published cultivated olive tree (63%) (12). Genome comparisons between oleaster and nine other plant species showed differences in gene numbers, transcript lengths, and proportions of transposable elements (TEs; *SI Appendix, Table S5B*). TEs and interspersed repeats occupied ~43% of the genome (Tables 1 and 2 and *SI Appendix, Table S7*). LTRs were the most abundant type of TE (40.3% of genome), which is in agreement with a previous analysis of a cultivated olive tree (38.8% of genome) (13), followed by DNA-type TEs (4.6%; *SI Appendix, Table S7*). A total of 50,684 protein-coding genes were predicted on the current assembly, of which 47,124 genes (93%) were confirmed by RNA sequencing

(RNA-seq) data. Further, 31,245 genes were located on the anchored pseudochromosomes (Fig. 1 and *SI Appendix, Fig. S6 and Tables S8 and S9*).

Approximately 90 million small RNA (sRNA) reads from six different tissues were used for noncoding RNA (ncRNA) annotation (*SI Appendix, Figs. S8 and S9 and Tables S10 and S11*). A total of 498 conserved miRNA families and 125 novel miRNAs were identified. Considering highly conserved miRNAs and their function, 29,842 miRNA–target pairs, including 7,849 unique target genes, were predicted. Totals of 4,606, 1,937, and 630 miRNA targets were associated with transcription factors, stress-response genes, and metabolism genes, respectively (*SI Appendix, Table S12*).

Oleaster protein-coding genes were functionally characterized through Gene Ontology and Kyoto Encyclopedia of Genes and Genomes (KEGG), which allowed annotation of 72.42% and 50.14% of all genes, respectively (*SI Appendix, Table S13*). KEGG metabolic pathway annotations of oleaster and 11 other plant species, including other oil crops such as *Sesamum indicum* (sesame) and *Glycine max* (soybean), as well as *Populus trichocarpa* (poplar) as a reference tree genome, *Utricularia gibba* (bladderwort) and *Mimulus guttatus* (monkey flower) as close relatives within the Lamiales, and *Fraxinus excelsior* (European ash tree) as a member of the Oleaceae family, showed a majority of oleaster genes to be involved in folding, sorting, and degradation ($n = 4,263$); biosynthesis of secondary metabolites ($n = 2,236$); carbohydrate metabolism ($n = 1,905$); and lipid metabolism ($n = 811$). Protein clustering of predicted oleaster genes with genes of other sequenced plant species resulted in 17,208 gene families, 1,070 of which were oleaster-specific and 7,522 were shared with the Lamiales *F. excelsior*, *S. indicum*, *M. guttatus*, and *U. gibba*. Although the number of gene families is largely consistent across the different species, the oleaster genome contains a large number ($n = 8,986$) of unique genes (*SI Appendix, Fig. S11 and Table S14*).

Genome Evolution. The oleaster genome contains multiple signatures of paleopolyploidy events. Distributions of synonymous substitutions per synonymous site (K_S) for the whole paranome (the set of all duplicated genes in the genome; *SI Appendix, Fig. S12A*) and duplicates retained in colinear regions only (i.e., excluding duplicates from small-scale duplications; *SI Appendix, Fig. S12B*) consistently showed two clear peaks of duplicates at K_S values around 0.25 and 0.75, respectively. Peaks at similar K_S values have been reported for duplicated genes in the genome of European ash (*F. excelsior*, a sister to oleaster in Oleaceae) (14). Most likely, these peaks indicate two rounds of ancient whole-genome duplication (WGD) in the oleaster lineage (15) shared

Table 2. Statistics of wild olive tree genome annotation

Annotation	No.	Total size, Kbp	Size, bp		
			Average	Maximum	Minimum
mRNA	50,684	65,933.6	1,300.9	48,863	99
CDS	50,684	52,756.9	1,040.9	16,602	99
Exon	235,149	65,933.6	223.4	7,913	1
Intron	184,465	87,396.5	473.8	42,191	10
miRNA	411	49,979	113.33	24	21
tRNA	798	59,716	74.83	95	63
rRNA	773	121,906	121	1,804	29
snRNA	422	47,737	113	217	62
Tandem repeat	454,960	372,874.8	819.57	500,000	25
TE protein	428,172	23,958.1	559.54	5,505	24
Transposon	320,201	150,867.9	471.16	5,928	11
5'-UTR	15,172	8,002.1	527.42	38,088	5
3'-UTR	15,075	7,337	486.7	47,263	5

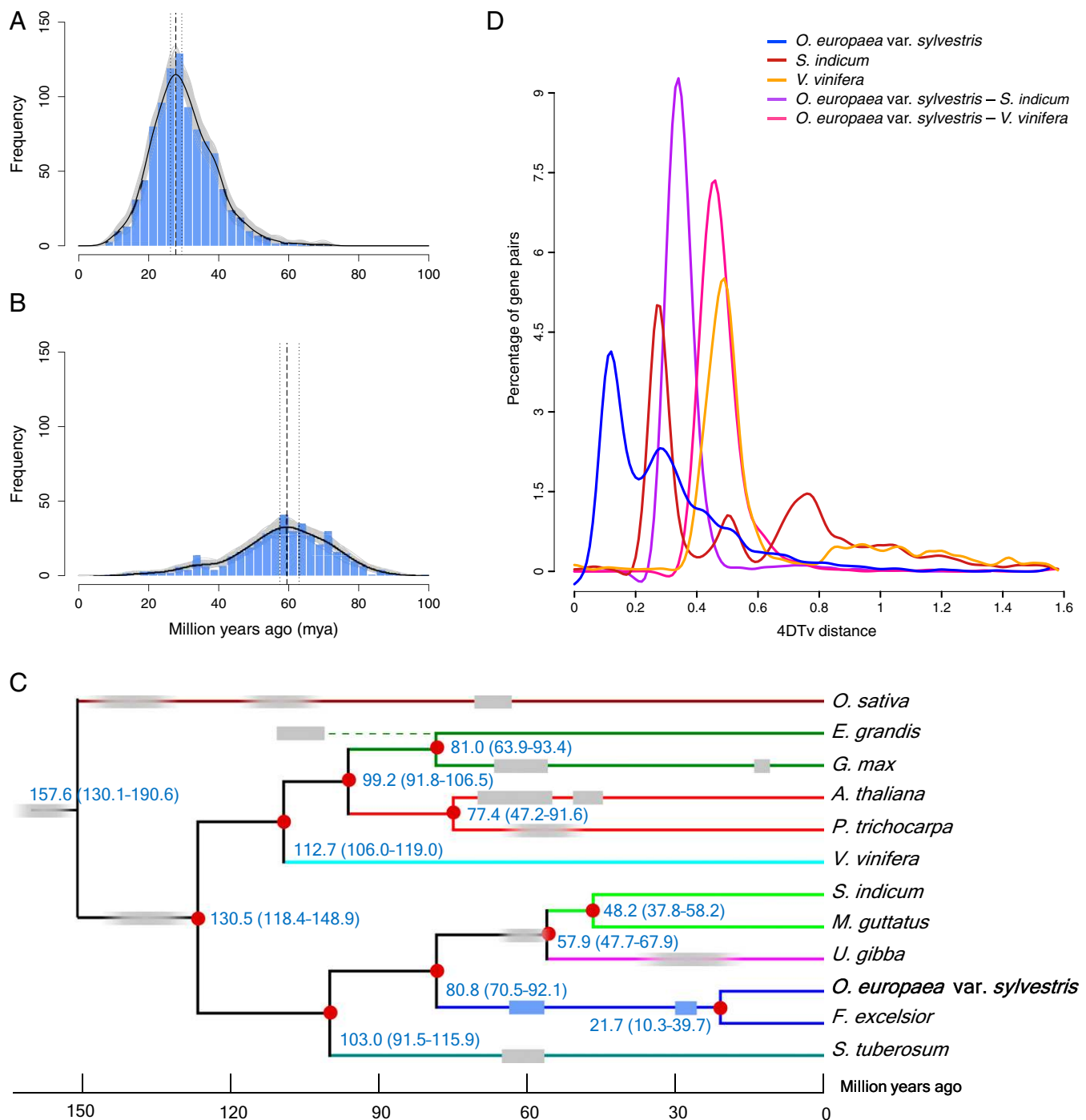


Fig. 2. Oleaster genome evolution. (A and B) Phylogenomic dating of *O. europaea* var. *sylvestris* paralogs. Absolute age distribution for the most recent WGD event (K_s of approximately 0.25; *SI Appendix*, Fig. S12A), with a consensus WGD age estimate of 28 Mya and 90% CI of 26–30 Mya (A). Absolute age distribution for the older WGD event (K_s of approximately 0.75; *SI Appendix*, Fig. S12B), with a consensus WGD age estimate of 59 Mya and 90% CI of 57–63 Mya (B). The solid black line represents the KDE of dated paralogs, and the vertical dashed black line corresponds to its peak, which was used as the consensus WGD age estimate. Gray lines represent density estimates from 2,500 bootstrap replicates, whereas vertical black dotted lines indicate the corresponding 90% CI for the WGD age estimate. Blue histogram shows the raw distribution of dated paralogs. (C) Estimation of divergence time. Blue numbers on the nodes are divergence time to present (in Mya). The two Oleaceae WGDs are indicated on the tree (blue rectangles), as are other known WGDs described in the literature for the species shown (gray rectangles; faded rectangles indicate that an absolute date has not been estimated). Note discussion of phylogenetic relationships in *SI Appendix*, S.3.2. (D) Fourfold degenerate (i.e., 4DTV) distributions for *S. indicum*, *V. vinifera*, and *O. europaea* var. *sylvestris*. Abscissa and ordinate represent 4DTV distance [using the HKY85 (Hasegawa–Kishino–Yano–1985) model] and percentage of homologous gene pairs, respectively.

by olive and ash (14). To establish the age of these two WGDs, absolute phylogenomic dating (16) was carried out. Absolute dating suggests that the most recent WGD had occurred approximately 26–

30 Mya (Fig. 2A) and the older one approximately 57–63 Mya (Fig. 2B). As with many other WGDs in different plant lineages, the latter event seems to have occurred close to the Cretaceous–Paleogene

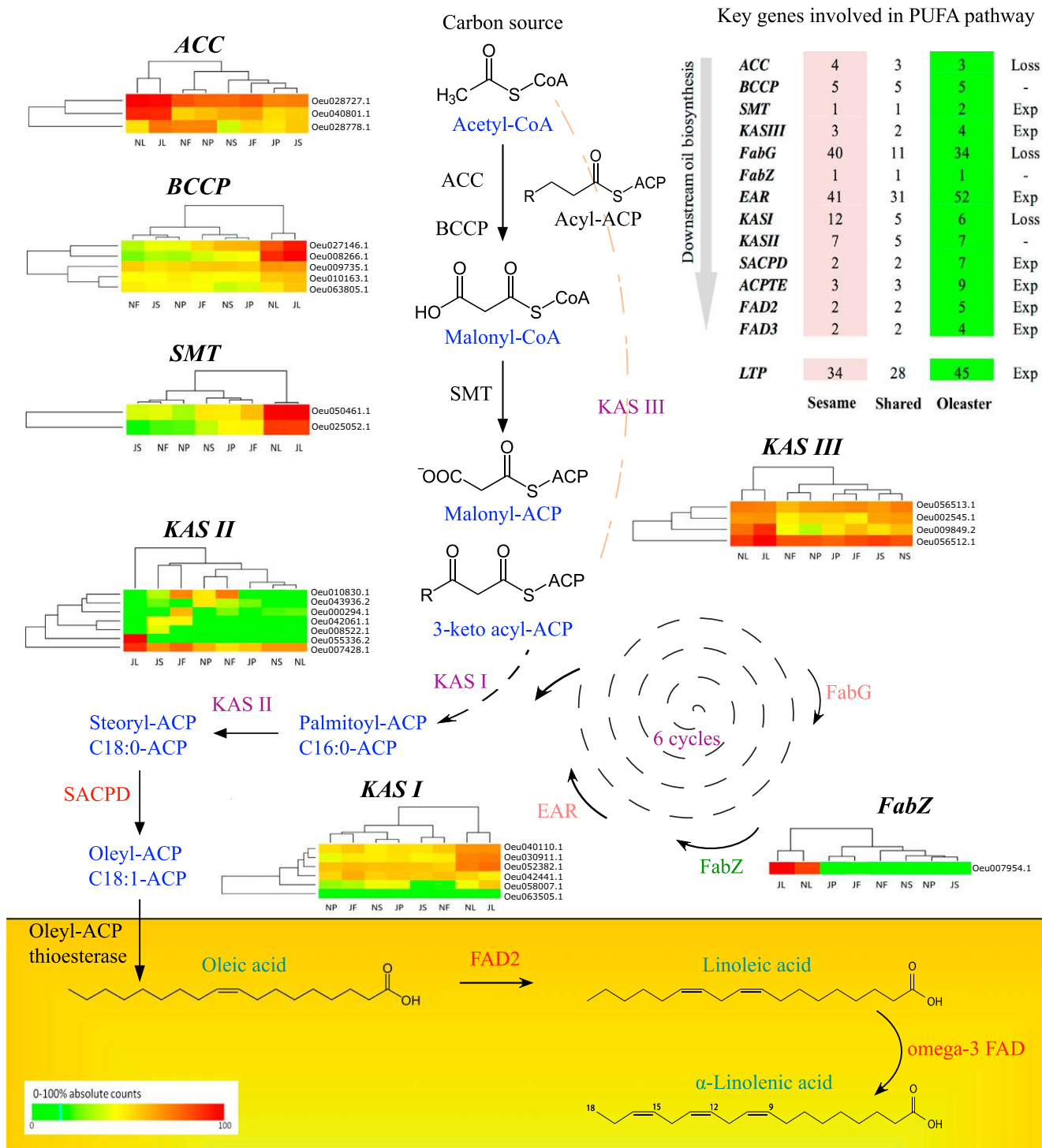


Fig. 3. Oleic-acid biosynthesis pathway in oleaster. Genes involved in oleic-acid biosynthesis with their differential expression patterns in stem (marked as "S"), leaf ("L"), pedicel ("P"), and fruit ("F") tissues are shown. Heat-map data correspond to start (July; "J") and end (November; "N") time points for olive oil biosynthesis. The first step of such biosynthesis is catalyzed by Acetyl-CoA carboxylase (ACC), carboxylating Acetyl-CoA to form malonyl-CoA, which is converted to malonyl-acyl carrier protein (ACP) by S-malonyltransferase (SMT). Malonyl-ACP first reacts with 3-keto acyl-ACP, which is elongated by six reaction cycles in which chain-extender units are added. Then, fatty-acid synthases (FASs) act on that substrate to produce saturated fatty-acid 16-carbon palmitate, which will be desaturated to form unsaturated fatty acids, such as oleic acid in oleaster. ACPTTE, ACP-hydrolase/thioesterase; BCCP, biotin carboxyl carrier protein; EAR, enoyl-ACP reductase; Exp, expanded; FabG, β -ketoacyl-ACP synthase; FabZ, β -hydroxyacyl-ACP dehydrase; FAD, fatty-acid desaturase; KAS, β -ketoacyl-ACP synthase; SACPDP, stearyl-ACP desaturase; SMT, S-malonyltransferase. Sesame expression data were retrieved from the Sesame Functional Genomics Database (SesameFG; www.sesame-bioinfo.org/SesameFG/).

extinction event, providing additional evidence that WGDs—at least in plants—might be linked with periods of environmental change or upheaval (17).

Paleopolyploidy events of similar age have been reported for other asterids in this period. Within the Solanales, a shared whole-genome triplication has been found in the lineage leading to *Solanum*

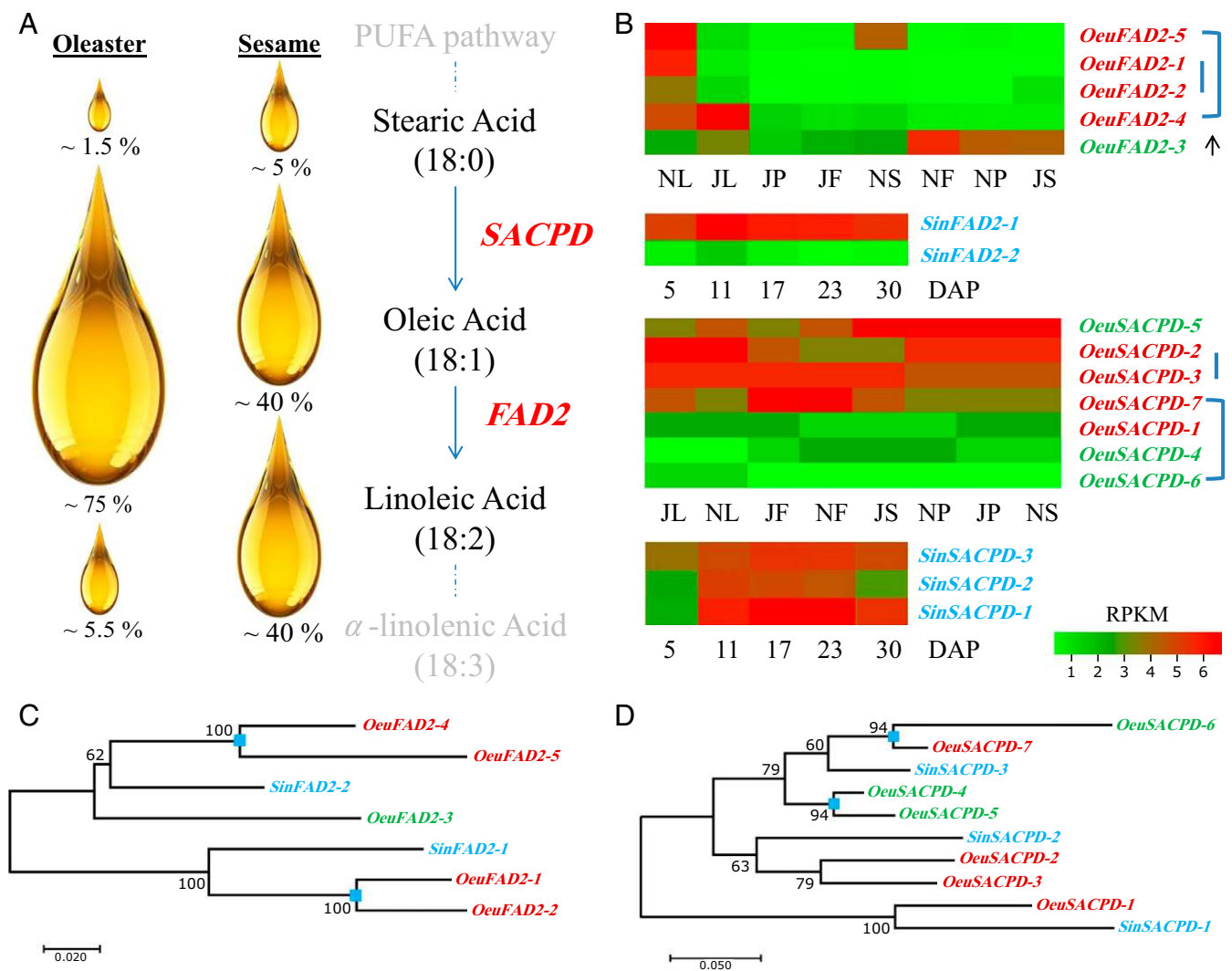


Fig. 4. Oleic-acid biosynthesis pathway in oleaster. (A) Oil content of oleaster and sesame with major genes involved in oil biosynthesis. (B) Heat-map analyses of oleaster and sesame *FAD2* and *SACP2* genes. Blue lines indicate paralogs, which share orthologs with sesame. The arrow represents up-regulation of *FAD2-3* gene, compared with other paralogs, in July unripe and November ripe fruits. Genes with green font color indicate unique genes in the wild olive tree, which have no orthologous counterpart in sesame, whereas red font color represents orthologous genes. Sesame genes are labeled with turquoise color. (C and D) Phylogenetic trees showing the duplication history of sesame and oleaster *FAD2* and *SACP2* genes. Blue squares show duplicated genes after WGD and tandem duplications (SI Appendix, Fig. S28A). DAP, days after pollination.

tuberosum (potato) and *Solanum lycopersicum* (tomato), with an estimated age approximately 57–65 Mya, using methods similar to the ones used here (16). Within the Lamiales, multiple WGDs independent from the paleopolyploidy in the Solanales have been described: two or three in the lineage leading to *U. gibba* (one of which could be shared with *M. guttatus*) (18) and one in the lineage leading to *S. indicum* (estimated age similar to tomato) (19). This latter one and the oldest WGD in *U. gibba* could denote the same event, possibly even shared with the older WGD in the oleaster and ash lineage, or both could be independent ones, partly depending on their phylogenetic relationship (SI Appendix, S.3.2). Mean estimates for the divergence of oleaster from *S. indicum* are 69–74 Mya (20–22) or even older (23, 24) (Fig. 2C). Duplication and speciation events analyzed using fourfold synonymous third-codon transversion rates (4DTv) also showed that there were probably two WGDs in oleaster and one WGD in *S. indicum*, and that these likely occurred after their divergence (Fig. 2D). Thus, the aforementioned dates and 4DTv patterns suggest that both WGD events inferred from the oleaster genome (as well as from the ash one) are specific to Oleaceae and occurred independently of

the WGD in the lineage leading to *S. indicum*, *M. guttatus*, and *U. gibba* (Fig. 2C; see also ref. 14). This seems further supported by a phylogenomic analysis of duplicates from the older oleaster WGD, in which a majority of trees supported an Oleaceae lineage-specific event (SI Appendix, S.3.4, Fig. S13, and Table S15). High colinearity among oleaster chromosomes forms additional evidence for WGDs. At least 78 duplicated homologous genomic segments, 12 of which are intrachromosomal, were identified in the oleaster genome. Among them, chromosomes 1 and 12 (4,743 genes), 7 and 14 (2,300 genes), and 6 and 21 (1,361 genes) are remarkably colinear (SI Appendix, Fig. S14 and Table S16).

Evolutionary Analysis of Oil Biosynthesis. Olive oil is mainly composed of TAG formed by fatty acids (10). Here, genes involved in oil biosynthesis were annotated and grouped according to their sequence identity, pathway, and enzyme codes. KEGG pathway analysis of genes related to oil biosynthesis in oleaster and 11 other species showed that the oleaster genome has the highest fraction of pathways related to lipid metabolism and secondary metabolite biosynthesis. Among 308 described pathway annotations, some of

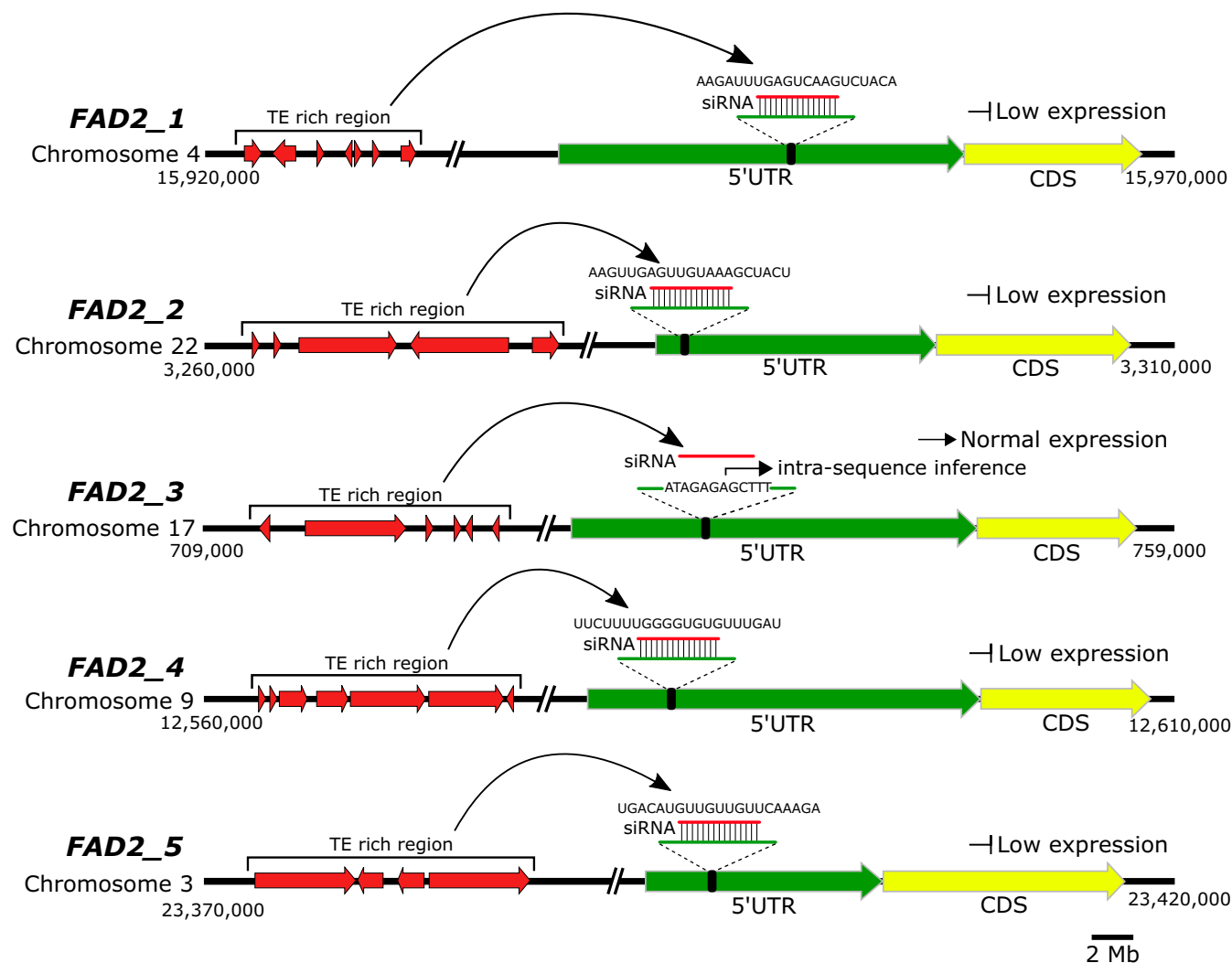


Fig. 5. Regulation of *FAD2* gene expression by siRNA. Possible siRNA-binding sites are marked on 5'-UTRs. Interestingly, siRNA can bind to *FAD2-1*, *FAD2-2*, *FAD2-4*, and *FAD2-5* transcripts but cannot bind to *FAD2-3* transcripts because of the presence of 12 additional nucleotides in the binding site (*SI Appendix, Fig. S27*). Red lines show siRNA molecules. CDS, coding sequence.

them, such as Ca^{2+} -transporting ATPase (K01537), acyl-CoA oxidase (K00232), and phosphatidylserine decarboxylase (K01613), are highly represented in the oleaster genome compared with others. To further compare the evolution of oil biosynthesis between oleaster and another major oil-bearing crop, oleaster and sesame genes were subjected to InParanoid ortholog analysis (25). Among 2,327 oil biosynthesis genes in oleaster, 2,025 seem to have homologs in sesame. After excluding outparalogs, 911 groups of orthologs could be built, with 1,232 inparalogs for olive tree and 1,171 inparalogs from sesame. Interestingly, 563 oil biosynthesis genes showed a strict one-to-one orthology between oleaster and sesame (despite independent WGD in oleaster and sesame), whereas the rest of the inparalogs (669 in oleaster and 608 in sesame) were the result of independent and lineage-specific duplication events (see Fig. 2 C and D). Furthermore, 94 of 267 genes (35%) were found to be unique to oleaster, in comparison with sesame, in terms of oil biosynthesis metabolic pathway annotation. Comparing orthologous genes between oleaster and sesame showed that a large proportion of genes required for oil biosynthesis have been maintained as duplicated genes in the oleaster genome (1,962 genes in 221 families). In contrast, only a small number of gene families (54 genes in 27 families) showed contraction in oleaster.

Fatty-acid biosynthesis is one of the major steps of complex oil biosynthesis (26). It includes elongation, degradation, and biosynthesis of unsaturated fatty acids and is carried out through the activity of a large number of genes encoding fatty acid synthases, elongases, desaturases, and carboxylases. Although the polyunsaturated fatty acid (PUFA) pathway is common in plants, and a considerable number of orthologous gene families ($n = 911$, as detailed above) are shared between oleaster and sesame, many important gene families involved in the oil biosynthesis pathway were found to be expanded in the oleaster genome compared with sesame (Fig. 3 and *SI Appendix, Fig. S17*). Besides the expansion of some oil biosynthesis gene families in the oleaster genome, the contraction of gene families encoding degrading/catabolic enzymes (such as dehydrogenases and hydrolases) may also be responsible for the differential fatty-acid accumulation in oleaster and sesame. For instance, the number of linoleic acid metabolism genes was found to be significantly smaller for oleaster ($n = 20$) than for sesame ($n = 164$).

To explore functional divergence following duplication, expression analyses were performed in different tissues collected from ripe and unripe fruits. Interestingly, it was observed that the expression of duplicated oleaster fatty-acid desaturase (*FAD2*) genes (*FAD2-1*, *FAD2-2*, *FAD2-4*, and *FAD2-5*) was down-regulated in fruit tissues,

especially during the lipid-accumulation ripening stage. Suppression of the expression of these genes causes reduced desaturation of oleic acid into linoleic acid (Fig. 4). *FAD2* genes underwent at least two rounds of WGD events in oleaster, but only one duplication event in sesame (19) (Fig. 4 *B–D*). By mapping sRNA reads to 10-kbp regions encompassing the oleaster *FAD2* genes (*SI Appendix*, Fig. S26), we discovered that an siRNA, which originated from a TE-rich region (27), may bind specifically to the 5'-UTR region of duplicated copies of the *FAD2* gene transcripts, repressing expression in the fruit tissues. Because of the presence of an additional 12 nt at the siRNA-binding site, the *FAD2-3* transcript, unlike the other *FAD2* transcripts, may not be regulated by the activity of the siRNA in ripe fruit (Fig. 5 and *SI Appendix*, Fig. S27). The *FAD2-3* gene is actively expressed in fruits and is responsible for the conversion of only a relatively low amount of oleic acid into linoleic acid (Fig. 4*B*). Sesame seeds also showed a differential expression pattern for *FAD2* genes (*FAD2-1* and *FAD2-2*), but with low diversity (*FAD2*, $\pi = 0.0016$), as reported previously (19). Consequently, silencing effects caused by siRNA on *FAD2* olive gene transcripts (*FAD2-1*, -2, -4, and -5; Fig. 5), and the low diversity in *FAD2* genes of sesame (19), are likely responsible for the higher accumulation of oleic acid in oleaster with respect to sesame.

Oleic acid as a major component of olive oil is formed by dehydrogenation from stearic acid by stearoyl-ACP desaturase (*SACPD*), after which it is desaturated into linoleic acid by *FAD2* (7). Expression measurement of oleaster *SACPD* genes showed that *SACPD1* and 2 have up-regulated expression in leaf tissues, whereas *SACPD7* is highly expressed in fruit tissues. On the contrary, *SACPD5* was found to be overexpressed in stem and pedicel tissues. Additionally, expression patterns of *SACPD1*, 5, and 6 were found at relatively low levels in other tissues (Fig. 4*B*).

It appears that the oleaster key genes involved in the PUFA pathway such as enoyl-ACP reductase (*EAR*), β -ketoacyl-ACP synthase II (*KASII*), β -ketoacyl-ACP reductase (*FabG*), acyl carrier protein (ACP)-hydrolase/thioesterase (*ACPT*), *SACPD*, and *FAD2* have been expanded by WGD and/or segmental duplications (*SI Appendix*, Figs. S28 and S29 and Table S17). Synteny analysis suggests that oleaster *FAD2-1/-2* and *SACPD6/7* paralogs have been duplicated through WGD (*SI Appendix*, Fig. S29*A*). Furthermore, *EAR* (52 genes), *ACPT* (9 genes), *FabG* (34 genes), and *KASII* (7 genes) were shown to be expanded by WGD (*SI Appendix*, Figs. S28 and S29 *B–E*) and tandem and segmental duplications and now have different expression patterns (Figs. 3 and 4).

Discussion

To date, besides the wild olive tree, the sequencing and assembly of two cultivated olive tree genomes have been reported, namely *O. europaea* cv. Leccino (13) and *O. europaea* cv. Farga (12), at $\sim 4\times$ and $\sim 150\times$ coverage, respectively. The latter, with a size of 1.31 Gbp, was preliminary annotated solely by using RNA-seq data, which resulted in more than 56,000 protein-coding genes (12). Compared with the oleaster genome presented here, the cultivated olive tree has a smaller genome size, albeit with a higher number of genes. Unlike some previous reports on olive tree genome data, which lacked chromosome anchoring and genome-wide functional annotation (12, 13), our study includes a near-complete representation and localization of genes, repeat elements, and sRNA, as well as functional and metabolic annotations and an evolutionary analysis of oil biosynthesis genes.

Absolute dating of the two identified WGD events in oleaster and 4DTv patterns suggest that both WGDs, which seem to be shared with the ash tree, are specific to Oleaceae and independent from WGDs reported in other non-Oleaceae Lamiales, including *S. indicum* (sesame; Fig. 2*C*). This is also consistent with synteny results from the ash tree genome (14). The age of the older WGD is close to the Cretaceous–Paleogene boundary. Additional Oleaceae genomes will be required to determine which of the other lineages

within Oleaceae, apart from ash, share either of the two WGDs, and whether one or both are related to patterns of diversification within the family (28).

The expansion of gene families and the functional divergence of genes playing important roles in oil biosynthesis may explain the higher accumulation of oleic acid ($\sim 75\%$ of olive oil) rather than linoleic acid ($\sim 5.5\%$ of olive oil) in oleaster (10). In sesame seed oil, both types of fatty acids are more evenly present ($\sim 40\%$) with lower variation ($\pm 5\%$; Figs. 3 and 4*A*) (19, 29). As a result of gene expansion and loss events in oleaster with respect to the PUFA pathway genes responsible for the accumulation of oleic and linoleic acids, the fatty-acid content of olive oil greatly differs from that of sesame seed oil (10, 19) (Fig. 4*A*).

Here, consistent with a previous report (27), we also describe an siRNA sequence that originated from a TE-rich genomic region. To inhibit expression of duplicated copies of *FAD2* gene transcripts, this regulatory siRNA may specifically bind to the 5'-UTR region of the transcripts in fruit tissues during the oil production period. In a previous study (30), it was reported that mutations associated with a duplication of the Oleate Desaturase (*OD*) gene caused its silencing by binding of an siRNA, further promoting accumulation of high levels of oleic acid in sunflower seeds. Similarly, suppression of *FAD2* gene expression as a result of gene expansion probably leads to the high oleic acid content in oleaster.

Based on expression analysis, *SACPD6/7* may have evolved through subfunctionalization or neofunctionalization events following their duplication (Fig. 4*B*). On the contrary, *FAD2-1/-2* have probably retained similar functions, as their expression patterns have not changed (Fig. 4*B*). Compared with sesame, expansion of *SACPD* genes (*SACPD1–7*) in oleaster has likely led to increased desaturation activity and increased expression through neofunctionalization of *SACPD2*, 3, 5, and 7 in fruit and stem tissues (Fig. 4*B*). Thus, neofunctionalized *SACPD* gene copies in oleaster are likely also responsible for the differences in oleic and linoleic acid contents of olive and sesame (19, 30). Recently, it was observed that mutations in the soybean *SACPD-C* gene promote higher accumulation of leaf stearic acid content, as well as changes in leaf structure and morphology (31). Therefore, *SACPD1* and 2, which are highly expressed in leaves, might be related to leaf morphology as well as oleic acid accumulation in fruit with overexpressed levels of *SACPD7* (Fig. 4*B*).

Methods

A full description of the study methods is provided in the *SI Appendix*.

Plant Material. A wild olive tree ($2n = 46$) was selected for whole-genome shotgun and transcriptome sequencing. Genomic DNA was isolated from leaf tissue (32).

Genome and Transcriptome Sequencing. Sequencing libraries were prepared and sequenced on the Illumina HiSeq 2000 platform, followed by assembly with SOAPdenovo (11). Transcriptome libraries of four tissues including leaf, stem, pedicel, and fruit (ripe and unripe), collected from two different seasons, were also sequenced.

Genome Assembly. All sequence reads were assembled with the SOAPdenovo software (11, 33) producing a reference sequence of the oleaster genome. A total of 319.39 Gbp of clean data were assembled into contigs and scaffolds by using the de Bruijn graph-based assembler of SOAPdenovo with the following four steps: (i) building contigs and scaffolds, (ii) filling gaps, (iii) removing redundancy, and (iv) reconstructing scaffolds.

Genetic Map Construction and Chromosome Anchoring. DNA samples of each F1 individual and parents were digested with PstI-MseI restriction enzymes and then ligated with enzyme-compatible adapters. To increase the number of PstI-MseI fragments, PCR amplifications were performed as described (34). The DArTseq (35) genotyping-by-sequencing (GBS) approach was used to identify SNPs. GBS data were analyzed by using a regression-mapping algorithm of JoinMap 4.0 software (Kyazma) to enable linkage-map construction.

MapChart 2.0 (36) was used for the graphical presentation of linkage maps. Genetic linkage maps were constructed to develop the integrated genome map for anchoring the scaffolds by using 94 individuals from a cross-pollinated population of a cross between cultivars Memecik and Uslu. For chromosome-scale pseudomolecule construction, two maps were established from two progenies: both F1 progenies of 92 individuals (Memecik × Uslu). An integrated map including 1,307 markers was established (37) based on double heterozygous loci (38, 39). Genetic markers were mapped onto the scaffolds by using the Burrows–Wheeler Aligner software module for alignment (40) with default parameters. Afterward, anchoring of assembled scaffolds to genetic maps was achieved by applying the ALLMAPS software (41).

Repeat Element Analyses. Homology-based and de novo approaches were used to find TEs in the oleaster genome. The homology-based approach involved applying commonly used databases of known repetitive sequences, along with programs such as RepeatProteinMask and RepeatMasker (42). RepeatModeler (www.repeatmasker.org/RepeatModeler.html) was used with two ab initio repeat-prediction programs (RECON and RepeatScout) to identify repeat-element boundaries and family relationships among sequences. Tandem repeats were also searched for in the genome by using Tandem Repeats Finder (43).

Gene Prediction. Homology-based and de novo methods, as well as RNA-seq data, were used to predict genes in the *O. europaea* var. *sylvestris* genome. GLEAN (44) was used to consolidate results. Protein sequences of *Arabidopsis thaliana*, *S. indicum*, *S. tuberosum*, and *Vitis vinifera* were aligned with TBLASTN and genBLASTA (45) against the matching genomic sequence by using GeneWise (46) for accurate spliced alignments. Next, the de novo gene-prediction methods GlimmerHMM (47) (<https://ccb.jhu.edu/software/glimmerhmm>) and Augustus (48) were used to predict protein-coding genes, with parameters trained for *O. europaea* var. *sylvestris*, *A. thaliana*, *S. indicum*, *S. tuberosum*, and *V. vinifera*.

Genome Annotation. Functional annotation was achieved by comparing predicted proteins against public databases, including UniProt, the UniProt Knowledgebase, KEGG, and InterPro. Results are available online at the Olive Genome Browser (olivegenome.org) and Online Resource for Community Annotation of Eukaryotes (ORCAE; bioinformatics.psb.ugent.be/orcae). Gene-family clustering was performed by OrthoMCL (49).

Evolutionary Analyses. The GTR+gamma evolutionary model was applied to reconstruct a phylogenetic tree by using 231 single-copy orthologous genes from 12 different plant genomes. K_S -based age distributions of oleaster were also constructed to unveil WGD events in oleaster (15). Absolute dating of two identified WGD events in the oleaster genome was performed as previously described (16) (*SI Appendix, S.3*). SyMAP (50) was used to identify synteny with other species (i.e., *S. indicum*, *V. vinifera*, *P. trichocarpa*, and *S. tuberosum*). Circos (51) was applied to generate a circular visualization of the oleaster genome features. InParanoid was used to identify orthologs and paralogs with sesame involved in the oil biosynthesis pathways. Additional information is provided in *SI Appendix, S.3*.

Availability of Data. The oleaster genome assembly has been deposited in the National Center for Biotechnology Information (NCBI) GenBank database (<https://www.ncbi.nlm.nih.gov/genbank>; accession no. MSRW00000000; BioProject record ID PRJNA350614). Transcriptome datasets were deposited in the NCBI Sequence Read Archive (<https://www.ncbi.nlm.nih.gov/sra>; accession nos. SRR4473639, SRR4473641, SRR44742, SRR4473643, SRR4473644, SRR4473645, SRR4473646, and SRR4473647). The genome and annotation files were also uploaded into ORCAE (bioinformatics.psb.ugent.be/orcae), Phytozome (<https://phytozome.jgi.doe.gov>), and the olive genome consortium Web site (olivegenome.org).

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