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SUPPLEMENTARY INFORMATION

2	Oak genome reveals facets of long lifespan				
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72 **1. Information about the genus** *Quercus*

73 Oaks are the dominant tree species in many temperate ecosystems and landscapes. Their species diversity and geographic distribution underlie this predominance. There are about 350 74 to 450 oak species worldwide¹, although species delineation remains a matter of debate due to 75 76 considerable phenotypic variation within species and frequent hybridization. However, oak 77 species diversity is much greater in North and Central America (about 200 species) than in Asia (about 150 species), and Europe (about 30 species)². On all three continents, a few 78 79 species have a continent-wide distribution: Q. petraea and Q. robur in Europe, Q. macrocarpa in North America³, and *Q. acutissima* and *Q. mongolica* in Asia⁴. The IUCN lists 80 81 13 oak species as critically endangered and 16 as endangered, mostly due to land use conflicts or overexploitation⁵. About 240 species are maintained in *ex situ* collections, in arboreta and 82 83 botanical gardens. Unlike other important temperate forest species, such as conifers, oaks are 84 not intensively cultivated in artificial forest plantations. Forest renewal is driven mostly by 85 natural regeneration, and oak plantations often target specialist output markets, for veneer, cork or truffles. In many countries, oaks are also used outside their native distribution range, 86 87 in urban forestry, for example, in which they are planted in parks and streets. Horticultural 88 cultivars have occasionally been selected for these highly specialized purposes.

89 Oaks provide major ecosystem services, ranging from the provision of raw construction 90 materials and the regulation of natural resources, to the conservation of biodiversity and the 91 provision of recreational and cultural services⁶. Oaks have been making an invaluable 92 contribution to human society since humans first reached the Northern Hemisphere, when acorns were a regular part of the human diet⁷. Timber can be obtained from most temperate 93 oak species, but oaks have always fulfilled multiple functions in human societies, by 94 providing a combination of habitat, economic and cultural services⁸. With recent increases in 95 96 public awareness of the environment, forest ecosystem services have been extended to include

97 the enhancement of carbon sequestration and biomass production for bioenergy purposes. Oaks produce numerous raw materials, including wood, cork, fiber, biomass, and 98 99 biomolecules; these raw materials are used to produce diverse manufactured products for the construction, food, pharmaceutical, and cosmetics industries⁹. This tremendous utility of oaks 100 101 is illustrated by the diverse uses of wood, bark (cork), leaves, and even acorns. Oak wood is 102 frequently used for fuel, timber frames, interior paneling, veneers, and barrels for wines and 103 spirits, whereas cork is use to make stoppers for bottles and in coverings for floors, walls and 104 ceilings. Tannins have been extracted from oaks for centuries, for use in the leather industry, 105 and oak secondary metabolites are now used in the cosmetics industry. Finally, there is a 106 renewed interest in the possible use of acorns in the human diet, for both nutritional and 107 ecological reasons, to meet the challenges raised by human population growth in a context of substitution for food products with a high carbon cost¹⁰. Oaks also generate other important 108 109 biological products by providing a habitat for associated species. Edible mushrooms, such as 110 boletes and truffles in particular, are the fruiting bodies of mycorrhizal fungi that grow in 111 association with oak roots and are harvested in many countries for their gastronomic and 112 nutritional qualities. Iron gall ink, which was used for centuries for the writing of official 113 documents and parchments, to ensure that they did not fade, is made from iron salts and gallic 114 acid from oak galls. Oaks also provide ecological services as single trees and as forests, by 115 offering shelter to a very large range of fungi, insects, birds, and other wildlife, the list of 116 species benefiting from these services being continually updated and lengthened. In many 117 parts of the world, oak forests have been assigned functions in habitat conservation, 118 contributing to the preservation of natural resources, such as water or soils.

119 Oaks also occupy a special position in science, for case studies of tree biology and evolution, 120 and as a major research tool in the fields of archeology, history and climatology¹¹. Oaks are 121 very long-lived, and oak-ring width series in Central Europe have been reconstructed as far

122 back as 8,480 BC. These ring series are used as a standard dating tool with a yearly resolution in archeology, and, in some cases, as a tool for dendro-provenancing¹². This resource is 123 124 continually updated with data from archeological remains, opening up additional possibilities for applications in climate reconstruction¹³. Oak microfossil remains are frequent and widely 125 126 distributed, due to the past and present widespread distribution of these trees, and may 127 therefore serve as biological indicators of previous plant distributions. The use of oak-ring 128 series also poses new research questions concerning the stasis or microevolution of oaks 129 across the Holocene and Anthropocene. Climate reconstruction, inferred from ring series, 130 assumes the conservation of a climate-growth relationship, which may be challenged by the 131 plastic or evolutionary response of oaks to environmental changes. These questions have 132 triggered genetic and genomic investigations of the ability of long-lived species to adapt to 133 rapid environmental changes.

134 2. Reference genome sequencing, assembly and anchoring

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2.1. BAC sequence analysis

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2.1.1. Construction and screening of libraries, sequencing and annotation

137 BAC library construction

Two BAC libraries were constructed from high-molecular weight genomic DNA from the Q. *robur* "3P" accession, partially digested with *Eco*RI for one library, and *Hin*dIII for the other. The *Eco*RI library, which was obtained from Clemson University (CUGI), was generated by a standard procedure¹⁴, as previously described¹⁵. This library comprises 92,160 BAC clones with a mean insert size of 135 kb, corresponding to approximately 12x coverage of the haploid pedunculate oak genome. A second BAC library was constructed, with *Hin*dIII digestion, to increase genome coverage and reduce the bias associated with the uneven

145 distribution of restriction sites. The HindIII library was constructed at the French Plant 146 Genomic Resource Center (CNRGV, http://cnrgv.toulouse.inra.fr/). Nuclei were isolated from young leaves¹⁴. High-molecular weight DNA was partially digested with *Hin*dIII and 147 148 subjected to size selection and the ligation of appropriately sized fragments into the 149 pIndigoBAC-5 HindIII-Cloning Ready vector (Epicentre Biotechnologies, Madison, 150 Wisconsin, USA). This library contained 98,304 clones with a mean insert size of 120 kb, 151 providing 14x coverage of the haploid genome. These two BAC libraries are available from 152 the CNRGV (http://cnrgv.toulouse.inra.fr/Library/Oak) under accession codes Qro-B-153 EnglishOak 3P (EcoRI) and Qro-B-3Ph (HindIII).

154 Screening of BAC libraries

155 The two BAC libraries were arranged in plate and row pools for PCR screening, as described by Chalhoub et al.¹⁶. Three sets of BAC clones were screened (summarized in 156 157 Supplementary Data Set 10 sheet #1): i) allelic BACs, to validate the assembly procedure 158 for the diploid genome sequence, i.e. the presence of distinct scaffolds (haplotypes) 159 corresponding to the allelic BACs, ii) BACs carrying expressional or positional candidate 160 genes coinciding with QTLs for water-use efficiency, bud burst or epinasty, for further studies 161 aiming to characterize QTLs associated with adaptive traits, and iii) BACs selected at random 162 or on the basis of BAC end sequences. The primer pairs for library screening were designed 163 from expressed sequence tags, gene sequences, genetic markers or BAC end sequences. PCR was performed as described by Faivre-Rampant et al.¹⁵. The success of PCR amplification 164 165 was checked by subjecting the PCR to electrophoresis in 2% agarose gels. Positive plate pools 166 were used to identify potential clones, which were subsequently validated by a second PCR 167 analysis of individual clones. For the identification of allelic BAC clones, the pools were 168 screened with primer pairs specific to single-copy microsatellite loci shown during genetic mapping experiments to be heterozygous in the reference "3P" genotype¹⁷. Each allelic BAC 169

170 clone was selected by visualizing length polymorphisms between PCR products or by direct171 sequencing of the products of PCR amplification for biallelic markers.

172 Sequencing and annotation of BAC inserts

In total, 34 BAC inserts were fully sequenced and annotated. Selected clones were cultured individually on LB medium and DNA was isolated by the standard alkaline method¹⁶. DNA inserts were sequenced in pools at 40x coverage, with the 454 mate-pair (5 kb) procedure. The sequence reads were assembled with Newbler (version MapAsmResearch-04/19/2010-patch-08/17/2010). Additional sequencing was carried out with the Illumina MiSeq platform (paired-end overlapping reads of 2×250 bp) at a coverage depth of 400x, and GapCloser (-V1.12-6) software was used to reduce the gaps between contigs.

180 Repeated elements were identified and classified in a two-step approach: i) Censor software and Repbase V21.08¹⁸ were initially used (see Plomion et al.¹⁹), but detection was limited to 181 182 BAC insert regions displaying identity to sequences in Repbase; ii) searches for the remaining 183 repeated elements were performed with the library of consensus repetitive elements presented 184 in section 3.1. Structural and functional gene annotations were added to the BAC sequence, as 185 described in the approach presented in section 3.3, using: i) Eugene to integrate *ab initio* and similarity-based gene finding programs²⁰, and ii) FunAnnotPipe, an in-house bioinformatic 186 pipeline based principally on InterproScan²¹. The data were then manually curated with 187 BLAST tools from the NCBI website and NetGen2²² for the confirmation of exon/intron 188 boundaries. Transcript evidence (ESTs and oak unigenes²³ were used to establish gene model 189 structures. We also used FGENESH²⁴ and Augustus²⁵ software to confirm or update Eugene 190 191 predictions, with Vitis vinifera or Theobroma cacao as the model. Some short-gene models 192 (encoding < 50 amino acids) were removed. Manually curated genes were then compared 193 with gene models predicted from the genome sequence.

194 Haplotype diversity analysis

We compared sequences between allelic BACs previously reported by Plomion et al.¹⁹, using 195 Dotter Yass software (http://bioinfo.lifl.fr²⁶). Local alignments were generated with NUCmer 196 package²⁷ 197 from the MUMmer and visualized with the Easyfig pipeline http://easyfig.sourceforge.net²⁸. 198

199 Data availability

BAC sequences were deposited in the European Nucleotide Archive under accession numbers LT99005-LT99038 (see **Supplementary Data Set 10 sheet #1** for the accessions). The 34 BAC sequences, with their annotations, are available from the oak genome browser (https://urgi.versailles.inra.fr/WebApollo_oak_PM1N/PseudoMolecule.html). The track "Gene_BAC_manual" provides manually curated gene models.

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2.1.2. **Results of BAC sequence analysis**

206 Eight of the 34 sequenced BAC inserts were assembled into continuous sequences without gaps. The others formed a set of oriented contigs separated by stretches of 100 nucleotides. 207 208 Each BAC corresponded to one or two scaffolds of the diploid version of the oak genome 209 sequence (Supplementary Data Set 10 sheet #1). Gaps were flanked by low-complexity 210 repeat sequences. One of the 34 BACs sequenced (#10P13) corresponded to chloroplast 211 DNA, nine corresponded to randomly picked clones from the libraries and 24 corresponded to 212 selected clones identified by PCR screening with single-copy genetic markers or candidate 213 genes. BLAST-n analysis with primer, genetic marker or candidate gene sequences confirmed 214 the presence of these sequences in the targeted BAC. The nuclear BAC assembly covered 215 4,282,332 bp. The mean G+C content of nuclear BAC sequences was 35.9%, and all BAC 216 sequences had G+C contents close to this mean. A similar G+C content was reported in a previous study for 20,056 BES¹⁶. 217

The 33 BAC sequences corresponding to nuclear regions were screened for simple sequence repeats (SSRs). In total, 1,342 perfect SSRs with a motif length of two to five nucleotides were detected within the 4,282,332 bp analyzed, corresponding to one SSR every 3,200 bp. A previous study had already shown the density of SSRs within the oak genome to be high. As previously described for BAC end sequences¹⁶, AT/TA dinucleotide motifs were the most abundant.

Repeat masking resulted in the masking of 24.7% of the BAC sequences, 99% of which corresponded to transposable elements (TEs), the other 1% corresponding to other types of sequence repeats. The percentage repeat content varied from 32% (clone #138D21) to 49% (#108022). Retrotransposons were the most abundant repeated elements, with 27% of the Gypsy type and 15% of the Copia type (**Supplementary Table 23**). A *de novo* repeat search detected 38.5% TEs, a value lower than that estimated for the whole genome (52%).

230 Putative genes were predicted with a combination of Eugene, trained on the oak genome, 231 FGENESH and Augustus (Supplementary Table 24). In total, 322 gene models were 232 predicted. Manual annotation was performed, with BLAST queries against NCBI non-233 redundant proteins, oak unigenes and oak ESTs available from the NCBI Short Read Archive. 234 Exon and intron structures were manually optimized on the basis of evidence for splice sites. 235 After manual curation, 44 of the 322 predicted genes were found to be located at the end of 236 BAC sequences and were not curated, 28 were deleted (corresponding to gene models 237 encoding < 50 amino acids), and intron/exon structure remained unresolved in 50, which were 238 therefore considered "problematic". Thus, 200 predicted genes with a resolved intron/exon 239 structure were finally approved. Intron/exon structure was modified for 37 of these 200 genes, 240 merged for 25 genes, and 138 genes (i.e. 69%) had already been accurately predicted by the 241 automatic annotation procedure described in section 3.3. This proportion is close to that of validated CDS from the set of 1,714 manually curated genes (79%, see section 3.5). We thus 242

found a mean of six genes per 100 kb, corresponding to one one protein-coding gene per 16.7
kb (Supplementary Table 24), a value twice the mean gene density across the genome (3.2
genes/100 kb). This bias probably results from the selection of genes for insertion into BACs.

Gene function was assigned on the basis of sequence identity to proteins within the phytozome and NCBI non-redundant protein database and/or the presence of Pfam domains (**Supplementary Data Set 10 sheet #2**).

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2.1.3. Comparison of genomic structure between haplotypes

250 Primer pairs of mapped simple sequence repeat (SSR) markers, PIE033, PIE260, PIE275, 251 PIE257, and ZQR111 and the CL4 candidate gene, were used to screen the two BAC libraries 252 (Supplementary Data Set 10 sheet #1) for allelic BAC identification. Allelic BAC clones 253 were identified by sequencing of the PCR products. Six sets of homologous BAC clones 254 (50E24-177A20/38C23, 27L3-48K1/72H20, 5E10-107I07, 64H03-30P1, 4E16/12J1-121F1, 255 and 4N17-11F22) were selected and sequenced. Except for BACs 111F22 and 64H03, an 256 analysis of BAC sequences confirmed the presence of the markers within the BAC clones. 257 The sequences of BACs 4E16 and 12J1 overlapped fully. We therefore removed 4E16 from 258 further analyses. Surprisingly, 4N17 and 111F22 did not overlap, and neither was therefore 259 considered in subsequent analyses. The overlap between the remaining allelic BACs ranged 260 from 22 kb to 84 kb and the mean sequence identity in overlapping regions was 97% (Evalue 261 =0.0) (Supplementary Table 25 and Supplementary Table 26). Pairwise sequence 262 alignment revealed insertions and deletions within the intergenic regions, for all pairs 263 (Supplementary Fig. 26). We identified TE insertion/deletion as the main factor accounting 264 for the considerable structural polymorphism observed between allelic BACs. Gene order and 265 structure were nevertheless well conserved.

266 **2.2. Comparison of the V1 and V2 assemblies and assembly validation**

We compared the previous release (version 1: V1¹⁹) of the diploid assembly with the current 267 268 release obtained by the addition of synthetic long reads generated by highly parallel library 269 preparation and the local assembly of short read data. Standard metrics revealed a huge 270 difference in terms of contiguity (see Supplementary Table 27). Indeed, the N90 of our 271 assembly was six times better than that of the previous assembly, and the proportion of 272 ambiguous bases was only 4.6% for our assembly, whereas it was 11.6% for the previous 273 assembly. We used Busco to assess the degree of gene completion for the two assemblies. The 274 V2 release presented a completeness of 90.8% (Supplementary Table 27), i.e. greater than 275 the V1 assembly (90.4%). Standard metrics also suggested that haplotypes were better 276 resolved in the V2 release (as indicated by the cumulative sizes of the assemblies). We then 277 validated the better differentiation between the two haplotypes of the V2 assembly, by 278 mapping a dataset of Illumina paired-end reads (2x250 bp) on both assemblies. Collapsing the 279 two haplotypes should increase the observed coverage by a factor of 2, whereas keeping the 280 two haplotypes separate should yield identical observed and expected coverages. As expected, 281 we observed fewer regions with twice the coverage in the V2 release (Supplementary Fig. 282 1). We also aligned the V1 (Supplementary Fig. 27, Supplementary Fig. 28, 283 Supplementary Fig. 29) and V2 (Supplementary Fig. 11, Supplementary Fig. 12, 284 Supplementary Fig. 13) releases with three pairs of BACs, each pair corresponding to the two alleles of the same genomic region. We first aligned the whole assembly against each 285 BAC with BLAT alignment tool²⁹ and default parameters, retaining only the scaffold with the 286 highest alignment score. Alignment positions were then extracted from a NUCmer³⁰ 287 288 alignment (identity > 90%) between each BAC and the corresponding scaffold. We selected 289 SNPs between allelic BACs, using a sliding window of 100 bp, and we used a seed sequence 290 of 41 bp (20 bp on either side of the SNP) to retrieve the allelic variants of the scaffolds. We 291 generated graphical representations (see Supplementary Fig. 11, Supplementary Fig. 12, 292 Supplementary Fig. 13, Supplementary Fig. 27, Supplementary Fig. 28, Supplementary 293 Fig. 29) highlighting the advantages of long reads for differentiating between haplotypes. The 294 V1 release often merged the two haplotypes into a single scaffold for the three genomic 295 regions, whereas the V2 release contained a pair of scaffolds for each pair of BACs. 296 Furthermore, the V2 scaffolds showed fewer switches between haplotypes than the V1 release 297 (see Supplementary Fig. 11, Supplementary Fig. 12, Supplementary Fig. 13, 298 Supplementary Fig. 27, Supplementary Fig. 28, Supplementary Fig. 29).

299

2.3. Pseudomolecule construction

300 Scaffolding can extend the contiguity of a genome sequence assembly by orders of magnitude 301 relative to contigs, but the construction of a chromosome-scale genome requires either 302 physical or genetic maps to anchor the scaffolds. We used a genetic map for this purpose. A composite genetic map was first established with LPmerge software³¹, bringing together 5,589 303 already mapped EST-SSR and SNP markers from eight individual linkage maps^{17,32} 304 305 (Supplementary Data Set 2 sheet #1), including one map for accession '3P' used to establish the reference genome sequence. Gene model sequences for the 5,589 mapped loci were then 306 aligned with the 1,409 scaffold sequences, using $BLAT^{29}$ (> 95% identity). In total, 2,615 307 308 unique scaffold/marker relationships (Supplementary Data Set 2 sheet #2) were identified 309 and classified into four categories (Supplementary Table 28). Overall, the scaffold-310 anchoring strategy (taking into account 2,285 markers from the most reliable assigned 311 scaffolds, i.e. from categories 1, 2 and 3) delivered 612 (43%) anchored scaffolds, covering 312 624.8 Mb (77%) of the haplome. Additional scaffolds were then anchored onto the 12 oak linkage groups, according to the synteny-driven strategy illustrated in Supplementary Fig. 2 313 (see Pont et al.³³ for the details). To this end, the 1,409 scaffold sequences were aligned 314 315 (BLAST-n, >70% identity) with the eight chromosomal sequences of Prunus persica (a 316 species phylogenetically related to Q. robur). This approach yielded a set of 653 scaffolds, 317 including 259 scaffolds anchored by synteny only (i.e. locally ordered according to the gene 318 order in Prunus persica), and 394 scaffolds already anchored and ordered with markers 319 (Supplementary Data Set 2 sheet #3). These scaffolds highlighted links shared between the 320 peach genome and the oak map and made it possible to intercalate the 259 scaffolds initially 321 anchored on the basis of synteny alone. Using the second set of 394 scaffolds, and comparing 322 gene order between the Prunus and Quercus genomes, we estimated the accuracy of the 323 syntenomic approach for the correct positioning and orientation of the first set of scaffolds 324 (anchored on the basis of synteny alone) at 86%. The 12 oak pseudomolecules (hereafter referred to as chromosomes and numbered according to the SNP-based linkage map³²) were 325 326 then constructed on the basis of 871 (62%) anchored and oriented scaffolds, with the filling in 327 of 100-nucleotide tracts between consecutive scaffolds (Supplementary Data Set 2 sheet 328 #4): i) 218 scaffolds anchored and ordered by genetic markers only, ii) 259 scaffolds anchored 329 by synteny only, with local ordering according to gene order in peach, and iii) 394 scaffolds 330 anchored and ordered by both procedures. Overall, the 871 scaffolds cover 716.6 Mb (i.e. 331 88% of the haplome) and contain 23,220 (90%) genes. The 12 chromosomes and the 538 332 unanchored the scaffolds are available from oak genome JBrowse 333 (https://urgi.versailles.inra.fr/WebApollo oak PM1N/PseudoMolecule.html).

Based on scaffold order and orientation on the 12 chromosomes, the oak genome browser was populated with a "marker" track including an optimized set of markers tolerant of inversions between physical and genetic positions within a maximum window of 5 cM. This track was designed to project the position of any quantitative trait locus (QTL) from the eight individual linkage maps onto the oak genome sequence, to facilitate subsequent biological interpretation of their genetic bases. The track was created according to the procedure described in **Supplementary Fig. 30**. We found that 2,127 of the 2,615 markers (retained for scaffold

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341 anchoring) fitted the criteria presented in Supplementary Fig. 30 (referred to as set#1 in 342 Supplementary Data Set 2 sheet #1), and 1,943 were retained from the other set of 2,974 343 markers initially excluded from scaffold anchoring (set#2 in Supplementary Data Set 2 sheet #1). As a result, the "marker" track included 4,070 markers spanning the 12 linkage 344 345 groups (red horizontal lines in Supplementary Fig. 31). The alignment of each marker set 346 with the 12 chromosomes is shown in Supplementary Fig. 32. Overall, the rank correlation between genetic and physical positions ranged from 0.991 to 0.999 (Supplementary Table 347 348 29).

349 3. Genome annotation

350 3.1. Detection and annotation of transposable elements

As in other sequenced plant genomes, the class I retrotransposon fraction predominated (70% of TE sequences), consisting of 53% LTRs (long terminal repeats: 26% Gypsy-like and 21% Copia-like) and 16% non-LTR retrotransposons (mostly LINE). Class II DNA transposons accounted for 15% of TE sequences, and 92% of the transposons in this fraction were TIRs (terminal inverted repeats) (**Supplementary Fig. 3**, **Supplementary Table 4**).

Thirteen of the 1,750 consensus sequences (0.6% of the TE content) were further characterized as Caulimoviridae sequences (see section 3.2). **Supplementary Table 30** shows a comparative analysis of TEs across the 16 species (including oak) used for the comparative genomic analysis in section 4. We found no correlation between TE content and the phylogeny of these species (based on NCBI Taxonomy Browser findings) (**Supplementary Fig. 33**).

362 **3.2. Identification and preliminary characterization of endogenous Caulimoviridae**

363 Plant viruses can have a major impact on the populations and genomes of their hosts. Paleovirology approaches can provide insight into virus-host associations by detecting 364 fragments of viral genomes integrated in host genomes³⁴. Caulimoviridae is a major family of 365 plant viruses with deleterious effects on plant populations and crop production³⁵. 366 367 Caulimoviridae do not need to integrate into the host genome during their replication cycle, 368 but such integration occurs randomly and repeatedly, resulting in the presence of significant numbers of Caulimoviridae genome fragments in plant genomes^{36,37}. We screened the oak 369 370 genome for the presence of genomic fragments from endogenous Caulimoviridae. Reverse 371 transcriptase (RT) is the best conserved domain of the Caulimoviridae family, so we began by searching the oak genome for RT domains displaying the highest levels of identity to 372 373 homologs from known Caulimoviridae genera. Protein clustering (>80% identity) identified 374 eight groups including seven comprising several sequences corresponding to RT sequences from Caulimoviridae. This viral family contains eight genera. Phylogenetic analysis revealed 375 376 that one of the RT cluster from endogenous oak Caulimoviridae belonged to the genera 377 Petuvirus, whereas the other seven belonged to the recently discovered Florendovirus genera³⁷ (Supplementary Fig. 34). 378

379 We then performed targeted clustering (98% identity and 95% length) on the nucleotide 380 sequences corresponding to putative Caulimoviridae loci in the oak assembly and built 381 consensus sequences based on the multiple sequence alignment (MSA) for each cluster. We 382 then clustered the consensus sequences with the closest evolutionary relationships to 383 Caulimoviridae into seven families, each of which displayed at least 90% local identity. In 384 five families, the longest consensus sequence accounted for a complete, or almost complete 385 Caulimoviridae genome and was, thereafter, considered the representative sequence for each 386 family. Remarkably, we noticed that, while representative consensus sequences were built 387 from the MSA of only a few highly similar copies, we found cases in which consensus 388 sequences corresponding to truncated variants of the representative Caulimoviridae genomes 389 were generated from the MSA of hundreds of almost identical copies (Supplementary Fig. 390 35). We compared the representative sequences with the library of repetitive elements built by 391 TEdenovo and found that most were well represented in this library (see section 3.1). 392 Collectively, copies of consensus sequences from the TEdenovo library corresponding to 393 fragments of the Caulimoviridae genome accounted for 4.4 Mb of the REPET annotation, 0.6 394 % of the haplome, and were distributed evenly over the 12 chromosomes (Supplementary 395 Fig. 36).

396 3.3. Gene prediction and functional annotation of protein-encoding genes

397 We retained a core set of 25,808 high-confidence genes (listed in Supplementary Data Set 398 1). The total gene space was 74 Mb in size, with a density of 0.32 genes/10 kb on average 399 (Supplementary Table 31). This density is lower than that reported for other species, such as A. thaliana (2.3 genes /10 kb; TAIR 10^{38}), P. persica (1.22 genes /10 kb³⁹), M. domestica 400 $(0.78 \text{ genes } /10 \text{ kb}^{40})$, but similar to that for species with a similar genome size and TE 401 content, such as *E. grandis* (817 Mb; 50% TE; 0.45 genes /10 kb⁴¹), *C. papava* (815 Mb; 52% 402 TE; 0.34 genes/10 kb⁴²) and C. clementina (816 Mb; 43% TE; 0.3 genes /10 kb; ⁴³). Overall, 403 99% of the predicted *Q. robur* genes (i.e. 25,516) were found to encode proteins, with at least 404 405 domain/motif, localization/targeting signal, or similarity-based evidence (Supplementary 406 Fig. 37).

407

3.4. TEs and genome dynamics

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3.4.1. Estimation of the age of TE families from consensus sequences

409 The consensus sequences used to annotate the TE copies in the oak genome represent 410 common ancestral structural variants (TE families) of TEs transposing in the oak genome in

the past⁴⁴. Indeed, they were constructed from highly repeated genome segments (see section 411 412 3.1). We investigated the evolution of TEs in the oak genome, by plotting and comparing the 413 observed divergence (1-identity %) of TE copies from their respective consensus sequences, to estimate their relative age⁴⁵. We performed this analysis separately for different orders of 414 415 TEs (LTR and Non-LTR retrotransposons, TIRs and Helitron DNA transposons) and 416 superfamilies (LTR Gypsy and Copia superfamiles). Most TE copies (i.e. 62% representing 417 44% of the TE space) displayed more than 15% divergence from the corresponding consensus 418 sequence (Supplementary Fig. 38), whereas only 6.7% of TE copies (17% of the TE space) 419 displayed low levels of divergence (<5%) from their respective consensus. This result 420 suggests that all the TEs present in the oak genome are relatively ancient, contrasting with 421 findings for A. thaliana, in which 73% of TE copies (52% of the TE space) display more than 422 15% divergence and 10.5% of the copies (26% of the TE space) display less than 5% divergence⁴⁵. By contrast, the divergence of the LTR retroelement superfamilies Gypsy and 423 424 Copia suggests that TE activity continued until fairly recently for the elements of these 425 families.

426

3.4.2. **Retrotransposition dynamics**

427 We first refined the annotation of LTR retrotransposons with the dedicated LTR Harvest tool⁴⁶, retaining the 5,904 complete elements that displayed more than 90% reciprocal overlap 428 429 with those from the general annotation of TEs with the REPET pipeline. We then classified 430 these elements into families by sequence clustering of their left LTRs with SiLiX, as previously described⁴⁷. We analyzed retrotranspositional history on a subset of 4,333 elements 431 432 from families of more than 200 elements. The insertion date of each element was calculated 433 from the sequence similarity between its left and right LTRs, as determined by LTR Harvest, as follows: date = $((1 - (\% \text{ identity}/100)) / 2.6) \times 10^{8}$ We plotted the data as density 434 histograms representing the distribution of insertion dates within each family, together with a 435

436 curve representing local density estimates (Supplementary Fig. 39). We observed a general 437 asynchronism of retrotranspositional history, with families displaying one of three contrasting 438 patterns of activity history: ancient (e.g. fam #6 and #10), constant (e.g. fam #8) or recent 439 (e.g. fam #12, #14 and #29). This result suggests that the complexity of the oak genome 440 developed through repeated bursts of retrotransposition over the last five million years, with 441 no clear increase in such activity in the recent past. These findings differ from those for 442 annual plants of similar genome size, for which genome complexification has occurred more recently, through concomitant bursts of transposition (over the last 1-2 million years⁴⁹). 443

444

3.4.3. **Distribution of TEs and genes in the oak genome**

445 TEs are often associated with genome rearrangements. They have been found in breakpointcontaining windows in comparisons of A. thaliana and A. lyrata³⁸. In maize and A. thaliana, 446 447 the pericentromeric regions of the chromosomes are highly enriched in LTR retrotransposons 448 of the Gypsy superfamily, and maize also displays an accumulation of TEs from the Copia superfamily in regions of euchromatin^{50,51}. We investigated whether TEs, particularly those of 449 450 the Gypsy and Copia superfamilies, were evenly distributed throughout the oak genome. We 451 calculated the percentage of TEs and annotated genes in sliding windows (300 kb, with a 200 452 kb overlap). We found that TEs accumulated in gene-poor regions. We also identified a 453 region of chromosome #2 displaying strong TE accumulation, potentially corresponding to 454 the centromeric region (Supplementary Fig. 40). The Copia elements tended to accumulate 455 away from the potential centromere, both upstream and downstream. This pattern was 456 particularly marked for chromosome #2 (Supplementary Fig. 41). Below, we consider the 457 potential role of TEs located in close proximity to genes.

18

458

3.4.4. Role of TEs in gene expansion and tandem duplication

459 We investigated the possible role of TEs in gene duplication and/or gene family expansion, by 460 comparing the genomic environment (in terms of proximity to TEs) of several categories of 461 genes according to their distance to the closest TE. The distance from each gene to the closest TE was calculated with getDistance.py from the S-MART package⁵². Only distances up to 5 462 463 kb were considered. We assessed the dependence between the different classes of distances 464 and belonging to an expanded gene family, for oak genes. We found that genes from 465 expanded gene families (see section 4.1.2) were closer to TEs than other genes (Chi-squared *p*-value $< 2.2e^{-16}$; Supplementary Fig. 42). TE-mediated gene family expansion has been 466 described in multiple species^{53,54}. We obtained similar results for tests of the dependence of 467 468 different classes of distances and membership of the TDG (tandem duplicated genes), LDG 469 (long distance-duplicated genes) and SG (singleton genes) classes. TDGs were closer to TEs than SGs or LDGs (Chi-squared *p*-value $< 2.2e^{-16}$; illustrated for SG in Supplementary Fig. 470 43), but no significant difference was observed for comparisons of LGD and membership of 471 472 the SG class. This result suggests that TEs may favor tandem duplications leading to gene 473 family expansion.

474

3.4.5. Horizontal transfer of TEs

475 We studied the horizontal transfer of TEs (HTT), by performing an in silico analysis on all 476 plant genomes available from the NCBI and Phytozome databases, focusing on LTR 477 retrotransposons. We chose one element for each family identified in the annotation step. 478 BLAST-n searches were performed to identify high levels of nucleotide sequence identity, 479 with the **NCBI** nr (http://www.ncbi.nlm.nih.gov/) and Phytozome v9.0 480 (http://www.phytozome.net/) databases. Candidates for HTT (listed in Supplementary Table 32) were detected by applying a 90% identity threshold⁴⁷, to ensure that we detected 481

482 horizontally, as opposed to vertically inherited TE sequences. Eight horizontal transfers of 483 LTR-retrotransposons were identified. All potential candidates were validated by checking 484 that the LTR retrotransposon sequences were located on large contigs and not on isolated, 485 short sequences in genome assemblies, and that the high degree of sequence identity was 486 limited to the elements themselves and not in their flanking sequences, to eliminate possible 487 contamination during genome assemblies and annotation errors. Moreover, analysis with 488 Dotter software confirmed that all the horizontally transferred elements harbored both the 489 LTR and an internal sequence in the two species involved. We identified six HTT events 490 involving oak and grapevine (Vitis vinifera), with sequence identities of 90 to 94%. We found 491 one HTT event involving oak, grapevine and peach (Prunus persica). The HTT event between grapevine and peach had already been identified (BO6) in the analysis by El Baidouri et al.⁴⁷, 492 493 and 92% identity was found between the corresponding sequences from the two species. We also identified one HTT event between oak and poplar (Populus trichocarpa), with 91% 494 identity. HTTs have been shown to occur frequently between flowering plants⁴⁷, but our 495 496 findings for the oak genome provide the first evidence of multiple HTTs in a single species.

497 **3.5.** Gene prediction, functional annotation of protein-encoding genes and manual 498 curation

The total gene space of the 25,808 predicted proteins was 74 Mb in size, with a density of 0.32 genes/10 kb on average (**Supplementary Table 31**). This density is lower than that reported for other species, such as *A. thaliana* (2.3 genes /10 kb; TAIR10³⁸), *P. persica* (1.22 genes /10 kb³⁹), *M. domestica* (0.78 genes /10 kb⁴⁰), but similar to that for species with a similar genome size and TE content, such as *E. grandis* (817 Mb; 50% TE; 0.45 genes /10 kb⁴¹), *C. papaya* (815 Mb; 52% TE; 0.34 genes/10 kb⁴²) and *C. clementina* (816 Mb; 43% TE; 0.3 genes /10 kb⁴³). Overall, 99% of the predicted *Q. robur* genes (i.e. 25,516) were 506 found to encode proteins, with at least domain/motif, localization/targeting signal, or 507 similarity-based evidence (**Supplementary Fig. 37**).

Experts manually checked (using WebAppolo) the protein-coding sequence structures of 1,714 mRNAs. They validated 79% of the transcripts without the need for additional modification, whereas the remaining 21% had to be corrected (**Supplementary Table 12**). We then aligned the coding sequences of these 1,714 mRNAs, to validate 2,067 genes of the *Q. robur* genome (diploid V2). Finally, 1,176 of these 2,067 genes were recovered in the *Q. robur* haplome. In the following sections we provide information concerning some of the gene families manually curated.

515

3.5.1. Aquaporin

516 Forty genes encoding putative aquaporins were identified in the Q. robur haplome. 517 Aquaporins are intrinsic channel proteins found in all organisms. Their overall structure is 518 highly conserved, with six transmembrane helices connected by five loops, a tetrad of amino-519 acids (helix 2, helix 5 and loop E) forming an aromatic/arginine constriction region (Ar/R 520 filter), and two membrane embedded half-helices with an asparagine-proline-alanine signature (NPA motif)^{55,56}. Five conserved amino-acid residues discriminated aquaporins from other 521 major intrinsic proteins⁵⁷. One *Q. robur* gene was invalidated due to the absence of a key 522 signature (Supplementary Table 33). 523

Q. robur was found to have aquaporins from the five subfamilies found in higher plants (**Supplementary Fig. 44**), with 14 plasma membrane-intrinsic proteins (PIPs), nine tonoplastintrinsic proteins (TIPs), eight nodulin-26 intrinsic proteins (NIPs), three small basic intrinsic proteins (SIPs) and five unrecognized X intrinsic proteins (XIPs). Two subclasses of XIPs were identified in *Q. robur*, with a particular mapping pattern for *XIP2*, suggestive of local amplification on Qrob_H2.3_Sc0000154. Except for the XIPs, the composition of the *Q*. *robur* aquaporin family was similar to those of *Arabidopsis* and maize^{58,59}. However, the fulllength *TIP3* gene was missing from the *Q. robur* genome. In several species, TIP3s have been reported to be specific to maturing and dry seeds⁶⁰. Variations at key motifs and in gene structure between the *Q. robur* aquaporin subclasses were consistent with published findings^{58,61}. The global rate of tandem duplication in this gene family was 37.5%, which similar to the overall rate for the oak genome (35.6%).

536

3.5.2. **MYB**

537 MYB genes are characterized by a highly conserved DNA-binding domain (MYB domain) 538 consisting of up to four imperfect repeats of a sequence of about 52 amino acids in length (R). 539 They constitute one of the largest families of transcription factors in plants, with members 540 regulating many key biological processes, including cell fate, developmental processes, primary and secondary metabolism, and responses to biotic and abiotic stresses⁶². MYB 541 542 proteins can be classified into several classes on the basis of the number of contiguous repeats 543 of the MYB domain. The most abundant of these classes contains MYB proteins with two 544 repeats of the MYB domain (R2R3-MYBs).

545 We identified 139 R2R3-MYBs, five 3R-MYBs and one 4R-MYB (Supplementary Table 546 34). This distribution of MYB proteins is similar to that in other species, such as A. thaliana 547 (126 R2R3-MYBs, five 3R-MYBs, and one 4R-MYB), E. grandis (141 R2R3-MYBs) and V. 548 vinifera (123 R2R3-MYBs). We performed a comparative phylogenetic analysis of the R2R3-549 MYB sequences from Q. robur, P. trichocarpa, E. grandis, V. vinifera, A. thaliana and O. 550 sativa (Supplementary Fig. 45). The topology of the phylogenetic tree was similar to that described for Arabidopsis⁶², with most of the subgroups conserved. However, like other 551 552 woody perennial plants, oak presented subgroups with more members than in herbaceous 553 annual plants such as Arabidopsis or rice (Supplementary Fig. 45). These expanded clusters in woody plants include the so-called "woody preferential subgroups", which are completly 554

555 absent from the basal lineages of bryophytes and lycophytes and from the more recent Brassicaceae and Monocot lineages⁶³. We investigated the possible role of the MYB gene 556 557 family in tree habit specialization by classifying R2R3-MYB genes according to their 558 duplication and expansion profiles in woody perennials. The global rate of tandem duplication 559 in the R2R3-MYB family (32.4%) was slightly lower than the overall rate for the oak genome 560 (35.6%). However, the tandemly duplicated MYBs were remarkably enriched within the 561 woody-expanded subgroups (Supplementary Fig. 46, Fisher's exact test *p*-value < 0.0001). 562 A substantial enrichment of tandemly duplicated genes belonging to woody expanded 563 subgroups has been also observed in other woody plants, such as eucalyptus, poplar and grapevine⁶⁴. 564

565 The few genes from subgroups expanded in woody perennials that have been characterized 566 seem to regulate phenylpropanoid metabolism, mostly controlling flavonoid biosynthesis, 567 although, in some cases, they also directly or indirectly alter the content of lignin and other soluble compounds, such as oligolignols or salicinoid phenolic glucosides^{64–69}. During 568 569 evolution, tandemly duplicated genes have a greater likelihood of being retained if they are involved in responses to environmental factors⁷⁰. Unlike herbaceous annuals, which die after 570 571 reproduction, perennial plants, such as trees and shrubs, must survive many periods of 572 challenging stressful environmental conditions over their long lifespans. Woody perennial 573 plants may, therefore, contain more elaborate stress resistance mechanisms. The large number 574 of tandemly duplicated genes regulating the biosynthesis of flavonoids and other 575 phenylpropanoid-derived compounds, mostly known to be protective, may enable oak trees to 576 develop complex protective mechanisms and to adapt woody growth to environmental 577 conditions. It is also possible that the production of the some of the many phenolic 578 compounds accumulating in oak heartwood, such as ellagitannins, or the gallotannins found in 579 oak galls, are controlled by these genes.

3.5.3. **SWEET**

581 The host plant supplies the mycorrhizal fungi with hexoses, which support the production of 582 the external fungal mycelium, a prerequisite for effective nutrient acquisition by hyphal networks. Plant sugar transporters of the SWEET superfamily deliver sugars to microbes⁷¹, 583 584 and the microbe-specific modulation of SWEET gene expression may alter sugar efflux at the site of colonization⁷². We therefore analyzed the phylogeny of the pedunculate oak SWEET 585 586 superfamily, and performed RNAseq analyses to determine whether the abundances of *Q*. 587 robur SWEET transcripts were altered by inoculation of the oak clone DF159 with the ectomycorrhizal fungus (EMF) Piloderma croceum⁷³, the mycorrhizal helper bacterium 588 589 (MHB) Streptomyces sp. AcH 505, and the causal agent of oak powdery mildew, Erysiphe 590 *alphitoides*⁷⁴. Oak clone DF159 was micropropagated and rooted for gene expression analysis as described by Herrmann et al.⁷⁵, and cultivated in gamma-sterilized soil-based microcosms, 591 as described by Herrmann et al.⁷⁴. Culture and inoculation conditions, RNA extraction, 592 sequencing and data processing for fungi and bacteria were as described by Tarkka et al.⁷³, 593 Kurth et al.⁷⁶ and Herrmann et al.⁷⁴. Sequence data were deposited in the NCBI Short Read 594 595 Archive (accessions for P. croceum: SRX383906, SRX383899, SRX383898, SRX798260, 596 SRX798261, SRX798262; for AcH 505: SRX976815, SRX976817, SRX976819, 597 SRX976827, SRX976829, SRX976831; for E. alphitoides: SRX2398909, SRX2398916, 598 SRX2398917, SRX2398913). T. magnatum ectomycorrhizae were sampled on 6-month-old inoculated Q. robur plantlets produced by Robin nurseries (St Laurent du Cros, France), 599 600 following their standard protocols. Total RNA was extracted from ectomycorrhizal root tips 601 of T. magnatum/O.robur using the RNeasy Plant Mini Kit of Qiagen with DNase step and 602 addition of 20 mg/ml polyethylene glycol to the RLC extraction buffer. Three replicates were 603 used for RNA-seq. Preparation of libraries from total RNA and 2 x 100bp Illumina HiSeq 604 sequencing (RNA-Seq) was performed at the GET platform (Génopole Toulouse MidiPyrénées, Auzeville, France) following their standard protocol. Quality filtered reads were aligned to the *Q.robur* haplome reference transcripts using CLC Genomics Workbench 9 (Qiagen). To identify transcripts differentially regulated in ectomycorrhizae compared to control roots (greenhouse grown non-mycorrhizal roots; ERX1916509-11) the test from Baggerly et al. implemented in CLC Genomic Workbench and *p*-values from the differential expression tests were adjusted for false discovery rate (Benjamini & Hochberg). The *T. magnatum* RNA-Seq data are available at NCBI/GEO as Series GSE97122.

We identified 14 *SWEET* genes in the oak genome (**Supplementary Fig. 47**), belonging to the four clades identified in *A. thaliana*⁷¹. Clade IV seems to have been expanded, with six members in oak, *versus* only one in *Malus domestica*, two in *Arabidopsis thaliana* (SWEET16 and SWEET17), two in *Eucalyptus grandis* and three in *Solanum tuberosum*. Biochemical characterization of the SWEETs of *Arabidopsis thaliana* showed that the members of clade I and II preferentially encoded monosaccharide transporters, whereas the members of clade III encoded disaccharide transporters, mostly for sucrose^{71,77}.

619 In total, five SWEET genes from clades I, III and IV were differentially expressed in oak with 620 the EMF Piloderma croceum and Tuber magnatum, the MHB Streptomyces sp. AcH 505, or 621 Erysiphe alphitoides. Oak clade I transcript Qrob P322480.2, homologous to SWEET1, was 622 upregulated by the EMF P. croceum and T. magnatum. Consistent with these findings, 623 arbuscular mycorrhiza (AM) formation also leads to SWEET1 induction in potato and in Medicago truncatula⁷⁸. By contrast, the abundance of the oak SWEET1 transcript was 624 625 decreased by Erysiphe alphitoides infection or inoculation with the MHB Streptomyces sp. AcH 505. The SWEET1 gene therefore displayed differential regulation as a function of the 626 biotic interaction. 627

The oak clade I *SWEET3* homolog *Qrob_P321700.2* was induced by *P. croceum* and *Streptomyces*, and a related gene was among those upregulated in the potato AM symbiosis⁷⁹.

All the clade III SWEETs have sucrose transporter activity, and the oak clade III transcript $Qrob_P657550.2$, homologous to *SWEET12*, was upregulated upon interaction with *P*. *croceum* and *Streptomyces*. Interestingly, a related gene is upregulated in the protocorms of the orchid *Serapias vomeracea* during interaction with an orchid mycorrhizal fungus⁸⁰. By contrast, most of the transcripts repressed in the arbuscular mycorrhizal symbiosis of potato corresponded to clade III SWEETs⁷⁹, suggesting that clade III SWEETs are differentially regulated in different types of mycorrhizal interactions.

Clade IV SWEETs are vacuolar glucose, fructose and sucrose carriers in A. thaliana⁸¹. The 637 638 oak clade IV SWEET17 homolog Qrob P216890.2 was upregulated upon interaction with P. 639 croceum and Streptomyces, as also reported for the closely related potato StSWEET17a and StSWEET17b in AM symbiosis⁷⁹. By contrast, Streptomyces treatment led to the 640 downregulation of Qrob P546940.2 in leaves, and the expression of a related gene, 641 StSWEET17c, was suppressed in potato AM symbiosis⁷⁹. Different expression patterns were 642 643 observed for clade IV genes during mycorrhizal interactions with oak, suggesting that 644 SWEET17 genes are regulated in a complex manner in beneficial symbioses. Thus, the predicted expansion of clade IV SWEET sugar efflux carrier genes in the oak genome and the 645 646 differential abundances of oak SWEET transcripts, may reflect the adaptation of oak to a 647 remarkably rich spectrum of biotic interactions.

648

3.5.4. Thioredoxin, glutaredoxin and glutathione transferase

Redox changes are major cellular disturbances that affect a range of processes throughout the organism's lifetime, through their involvement in various stages of development and in stress responses, in particular. Post-translational redox modifications of proteins are increasingly being recognized as a rapid, targeted mechanism for initiating cellular responses in a very short timeframe⁸². For example, light is known to control carbon metabolism enzymes through a cascade of electron exchange reactions, including dithiol-disulfide exchanges.

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655 Within cells, these reactions are controlled by thioredoxins (TRX) and glutaredoxins (GRX), 656 encoded by two multigenic families containing 20 to 40 genes and constituting the reducing systems^{83,84}. Different isoforms are often present in different subcellular compartments, 657 probably because they catalyze different reactions or have different protein partners. In 658 659 addition to these regulatory functions, TRX and GRX are also required for the regeneration of 660 some detoxification enzymes, particularly those requiring a reactive catalytic cysteine residue. This residue oxidized upon reaction with the substrate, as in peroxiredoxins and methionine 661 662 sulfoxide reductases, must be recycled for the next turnover. We have also investigated the 663 glutathione transferase (GST) family, the members of which have certain structural and biochemical features in common with GRXs. An analysis of the possible expansion of this 664 665 gene family was also particularly enlightening, because its members are involved in 666 secondary metabolism and in xenobiotic detoxification, and display transcriptional regulation 667 in very diverse stress conditions. The TRX, GRX and GST gene content of Q. robur was thus 668 analyzed at the level of defined subclasses, with comparisons with other photosynthetic 669 organisms, including multiple tree species (Populus trichocarpa, Prunus persica, Citrus 670 clementina and Eucalyptus grandis) (Supplementary Table 35). In most pairwise 671 comparisons, the number of genes remained remarkably constant, indicating that these 672 systems are essential for plants.

The genes of the GRX family displayed the greatest variation in number in plant-specific class III, with 9 to 24 genes in angiosperms, the oak genome having an average number of these genes (N=14), 50% of them being tandem-duplicated. Hence, variations in this class can be accounted for mostly by species-specific duplications. Similarly, the oak genome has a number of genes from the TRX/TRX reductase family similar to that in other organisms, and none of the subclasses are missing. The most striking characteristic is the presence of six genes for NADPH-TRX reductase a/b type (NTRa/b), rather than the one or two in other species, with four of the genes in oak identified as tandem duplicate genes. In line with this
observed expansion, the associated orthogroup (#2778) was found to be enriched in GO term
(GO:0004791) analysis.

Fourteen classes of GST genes were identified in the last phylogenetic analysis performed 683 with photosynthetic organisms⁸⁵. Only 11 of these classes are present in angiosperms, the 684 685 other three classes being found in *Physcomitrella patens*. The total number of genes (88) in oak is in the upper part of the range, as are those for *P. trichocarpa*, *E. grandis* and *S. bicolor*. 686 687 A detailed subclass analysis revealed that the difference between oak and other organisms resulted principally from the presence of a larger number of GST Tau (GSTU) family 688 689 members. This finding was confirmed by the orthoMCL analysis (see section 4.1.2), with an 690 expansion observed for four clusters, including one with 19 GSTU genes (red branches in 691 Supplementary Fig. 48). From this very variable gene content (there is no GSTU in P. 692 patens, but 21 to 62 GSTU genes in the analyzed angiosperms) and the presented 693 phylogenetic tree (many sequences cluster by species), it seems clear that GSTU genes evolved relatively recently and in a species-specific manner in plants. This situation differs 694 695 from that for most GRX and TRX classes, for which photosynthetic organisms usually have the same number of each isoform (Supplementary Table 35)^{83,84}. This difference is also 696 697 highlighted by the 76.1% rate of tandem duplication for the GST family (mostly due to the 698 largest classes, GSTF and GSTU), much higher than the 22.0% and 28.0% reported for the 699 TRX/TRX reductase and GRX families, respectively, and the value obtained for the oak 700 genome (35.6%).

701 3.5.5. **MLO**

Studies of the MLO (mildew locus O) family of disease resistance genes are particularly
 relevant in *Quercus*, the plant genus infected by the largest number of powdery mildew

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species (16 from six different genera⁸⁶). This large group of obligate plant pathogenic fungi 704 infects almost 10,000 species of angiosperms⁸⁷. The first MLO gene was isolated from 705 barley^{88,89}. It was found to act as a susceptibility gene, with recessive loss-of-function alleles 706 707 (mlo) associated with broad-spectrum resistance (i.e. to all genotypes/races) to the fungal pathogen *Erysiphe graminis f. sp. hordei*, one of the causal agents of powdery mildew⁹⁰. 708 709 Unlike many of the resistance genes used in crop plants, which have been rapidly overcome 710 by virulent races of pathogens after deployment, *mlo* resistance has remained durable in the 711 field for decades, despite its widespread use. Mildew-resistant mlo mutants have also been described in Arabidopsis thaliana, tomato and pea⁹¹. The Mlo gene encodes a protein of 712 713 unknown biochemical activity, with seven transmembrane domains, located at the plasma 714 membrane. Mlo genes have been found into small families (often about 15 genes) in the genomes of many higher plant species, including Prunus persica⁹², Vitis vinifera, Cucumis 715 sativus, and others⁹³. Functional studies in Arabidopsis have shown that MLO function is not 716 717 restricted to plant-powdery mildew interactions. Instead, these proteins are also involved in pollen perception⁹⁴ and root thigmotropism⁹¹. 718

719 We found 19 MLO genes in the haplome of Q. robur (Supplementary Table 36, 720 Supplementary Table 37, Supplementary Fig. 49), including seven belonging to clade V. 721 This clade contains the genes associated with powdery mildew susceptibility/resistance in Arabidopsis thaliana⁹⁵ and some other species. The large number of MLO genes in oak, 722 723 particularly in clade V, is only surpassed by soybean, cotton and apple, all of which have 724 undergone recent whole-genome duplication events. Most of the MLO genes are located on 725 chromosomes #8 (5 genes, 4 of which belong to clade V), #10 (4 and 1) and #1 (3 and 2). We also found seven incomplete genes with strong homology to MLO. As MLO genes are 726 susceptibility genes, these incomplete genes may confer resistance⁹⁰. There are three 727 728 incomplete genes on chromosome #10, at the 5' and 3' ends of a complete clade V mlo gene.

If we consider all complete and partial genes, the overall rate of tandem duplication of MLOgenes is 46.2%, slightly higher than the overall rate for the oak genome (35.6%).

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3.5.6. **NB-LRR**

NLR-parser (⁹⁶, https://github.com/steuernb/NLR-Parser) was used to identify disease 732 733 resistance genes encoding nucleotide-binding leucine-rich repeat proteins (NB-LRRs or 734 NLRs) and related proteins from the oak genome (haplome). We identified an initial set of 735 1,431 genes. Based on the orthoMCL analysis, 81 proteins from NB-LRR-related classes (i.e. 736 orthogroup #1000, 1004, 1015, 1031, 1084, 1140, 1187, 1269, 1540, 1697, 2368, 2399, 4549, 737 5011, 7397, 14497 and 15991, see Supplementary Data Set 3 sheet #1) were added to the 738 set of putative disease resistance genes. The accuracy of domain prediction was checked with the NCBI Conserved Domains CD-search website (⁹⁷, Batch CD-search version). Each protein 739 740 was manually inspected and attributed to a given category based on the presence of the 741 following canonical domains: Toll-interleukin receptor-like (TIR), NB and LRR. The non-742 TIR domains found in oak putative NLRs consisted of coiled-coil (CC) and resistance to 743 powdery mildew protein (RPW8) domains, referred to as CNL and RNL, respectively. 744 Finally, we also recorded any other domains (X) potentially representing integrated domains^{98,99}. After curation and removal of mispredictions, we recovered a total of 1,091 745 746 putative NB-LRR-related protein-encoding genes (Supplementary Data Set 5), 834 of which 747 had a putative complete or partial NB domain and 54 showed a non-canonical and putative 748 integrated domain. Many of these integrated domains, possibly acting as decoys for pathogen effectors^{100,101}, are DNA-interacting domains such as zinc-finger or Myb/SANT family 749 750 domains. Other notable integrated domains had signaling functions (e.g. WD40) or were 751 previously reported in secreted proteins from animal parasites and pathogens, i.e. the 752 Rhomboid protease family (Pfam PF1694).

753 Beyond the large number of single domains retrieved (i.e. 14 CC, 151 TIR, 1 RPW8, 61 NB 754 and 85 LRR, Supplementary Table 8), the total complement of NB-LRR genes in the oak 755 genome is remarkable by comparison to those of other species. The list LRR genes is 756 probably incomplete, as this category is inherently very difficult to characterize due to the 757 highly variable number of LRRs and the abundance of other LRR-related proteins (e.g. LRR-758 RLK or LRR-RLP), a group that is also expanded in the oak genome. If we exclude single 759 domains, then, for the 1,091 genes, TIR-related NB-LRR proteins account for 43% of the 760 remaining disease resistance genes (335 of 779 genes). This ratio of TIR- to non-TIR- NB-761 LRRs close to 1 indicates that the disease resistance gene content of the oak genome is more balanced than reported for other eudicots¹⁰²⁻¹⁰⁵. One group of non-TIR NB-LRRs, the 762 763 expanded set of RNLs (orthogroup #1140) may also reflect the evolutionary history of 764 pedunculate oak with the fungus *Erysiphe alphitoides*, responsible for oak powdery mildew. 765 No disease resistance gene for this disease has been identified and cloned or described in oak 766 trees, but this gene complement suggests considerable potential for resistance to these 767 pathogens and represents a valuable source of genetic information.

768 We investigated the expansions detected in the oak genome by the orthoMCL/CAFE analysis 769 in more detail, by retrieving protein sequences with an NB domain from classes that 770 displaying marked expansion relative to other plant species. We focused in particular on 771 orthogroup #1000 (labeled as #1 in Fig. 3d and 4b). Multiple alignments were constructed for selected proteins, with the hmmalign program, from the HMMER 3.0 package¹⁰⁶, and the 772 773 Pfam NB-ARC domain (PF00931) seed alignment converted into a hidden Markov model 774 profile by hmmbuild. Collected NB domains were manually inspected and truncated domains 775 and obvious outliers were discarded. Orthogroup #1000 genes encoding TNL-related proteins 776 accounted for 1,927 sequences from the 16 plant genomes used in the orthoMCL analysis, 777 only 1,641 of which had an NB domain suitable for alignment. There were 308 oak genes in 778 the final set: 174 TNLs, 115 NLs, 16 TNs and 3 Ns. The Homo sapiens apoptotic protease-779 activating factor-1 (APAF-1) sequence, a commonly used outgroup for NB phylogenetic 780 analysis, was added to orthogroup #1000 for tree rooting. A global alignment was obtained 781 with Clustal-Omega in Seaview and conserved sites were selected manually with G-block implemented in Seaview¹⁰⁷. The maximum likelihood tree was estimated in RAxML 7.7.2, 782 783 with the standard algorithm, the PROTGAMMAIWAG model of sequence evolution and 1,000 bootstrap replicates¹⁰⁸. The phylogenetic tree was designed with FigTree v1.4.3 784 785 (http://tree.bio.ed.ac.uk). The TNL-containing orthogroup #1000 (Supplementary Fig. 6) 786 displayed two major specific expansions in oak that were well supported by bootstrap values. 787 Within these two clades, several small physical clusters containing more than three 788 contiguous genes were identified. These physical clusters were well supported by bootstrap 789 values and consisted of numerous tandem duplicates (see Supplementary Data Set 5 for 790 details). With 75 genes in total, chromosome 9 was found to have the largest number of TNL 791 clusters distributed along its length. Although based only on 85% of the genes of orthogroup 792 #1000 showing a correct NB domain for alignment, the phylogenetic analysis highlights the 793 obvious expansion of TNLs and related resistance proteins in woody species, shown in brown, 794 relative to other selected plants, shown in green. Other notable large expanded clades 795 corresponded to *E. grandis* and *M. domestica* (Supplementary Fig. 6).

796

3.5.7. **RLK**

Receptor-like kinases (RLKs) constitute one of the largest gene families in plants. The functions of most RLKs are unknown, but the functions described for members of this family include innate immunity, pathogen response, abiotic stress, development, and, in some cases, multiple functions. RLKs usually consist of three domains: an N-terminal extracellular domain, a transmembrane domain, and a C-terminal kinase domain (KD). Leucine-rich

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repeat-receptor-like kinases (LRR-RLKs), which contain up to 30 leucine-rich repeat (LRRs)
in their extracellular domain, constitute the largest RLK family.

804 We identified a genome-wide repertoire of oak RLKs containing a KD (PF00069.16), with the hmmsearch program¹⁰⁹. The KDs of oak RLKs were then aligned with those of RLKs from 805 Arabidopsis thaliana (623) and Oryza sativa (1,147), using MAFFT¹¹⁰. Alignments were 806 cleaned with trimAl (gt 0.2,¹¹¹) and used to build an approximate maximum-likelihood 807 phylogenetic tree (Fastree 2.1.8,¹¹²). The quality of the alignments was systematically 808 809 manually checked around all sites on which a positive selection footprint was detected. If the 810 alignment was dubious (less than 4 sequences, presence of numerous gaps, or too divergent 811 sequences), the site was not considered.

812 With Arabidopsis and rice genes as references, this tree was used to classify the oak RLKs into subfamilies, and into 20 subgroups (SG) for LRR-RLKs¹¹³. We identified 1,247 RLK 813 814 genes, corresponding to 4.83% of the gene repertoire, versus only 2.28% in Arabidopsis and 815 2.06% in rice (Supplementary Data Set 6). Two RLK subfamilies are clearly 816 overrepresented in oak: SD1 (0.88% of the oak gene repertoire, versus 0.11% in Arabidopsis 817 and 0.04% in rice) and LRR-RLK (1.69% of the oak gene repertoire, versus 0.83% in Arabidopsis and 0.67% in rice). A comparison with the LRR-RLK repertoire of 31 other 818 angiosperm species¹¹³ showed that two subgroups, SG-XIIa and SG-XIIb, displayed the 819 820 highest overall expansion rates relative to the estimated number of genes in the angiosperm 821 last common ancestor (102 copies in oak, expansion rate of 6.8 for SG-XIIa, and 50 copies in 822 oak, expansion rate of 10 for SG-XIIb). As for NBS-LRR genes, a large proportion of LRR-823 RLK expansions were caused by tandem duplications: 72% and 79% for SG-XIIa and SG-824 XIIb, respectively. In addition, LRR-RLKs from SG-XIIa, most of which belonged to 825 orthogroup #1006, displayed significant expansion in oak (labeled as #6 in Fig. 3d) and in 826 trees more generally (labeled as #5 in Fig. 4b), whereas LRR-RLKs from SG-XIIb, mostly from orthogroup #1003, displayed significant expansion only in trees (labeled as #3 Fig 4b).
The few known genes in SG-XIIa include *FLS2* (FLAGELLIN-SENSITIVE 2) and *EFR* (EFTU RECEPTOR) in Arabidopsis, and *Xa21* in rice. The SG-XIIb subgroup includes *XIK1*(Xoo-induced kinase 1). All these receptors are involved in the response to bacterial
aggression.

832 The detection of a positive selection signature provides direct objective evidence of the 833 adaptive role of lineage-specific duplications. We therefore investigated whether, and to what 834 extent, the lineage-specific expanded LRR-RLKs and LRR-RLPs in oak harbored positive selection signatures in oak, as they do in other species¹¹³. Indeed, the detection of positive 835 836 signature of selection is a direct and objective evidence of the adaptive role of lineage specific 837 duplications. We used the orthoMCL families built from the same set of 16 species (15 838 species plus oak). We realigned the proteins from three significantly expanded families of 839 LRR-RLKs (orthoMCL orthogroups #1003, #1006 and #1016 in Supplementary Data Set 3), and two of LRR-RLPs (#1009 and #1049 in Supplementary Data Set 3) using MAFFT¹¹⁰ 840 and trimAl (gt 0.2)¹¹¹. Phylogenetic trees were built for each family (PhyML 3.0¹¹⁴ and 841 842 groups of oak ultraparalogs (*i.e.* sequences only related by duplication) were identified using a 843 tree reconciliation approach (between the gene trees and species tree), as described by Fischer et al.^{113,115}. For each group of ultraparalogs, sequences were aligned to preserve the coding 844 phase (using Prank with the 'codon' option¹¹⁶ and Guidance for cleaning¹¹⁷. We used the 845 EggLib package¹¹⁸ to infer the maximum likelihood phylogeny at the nucleotide level for 846 every alignment, with PhyML 3.0¹¹⁴, under the GTR substitution model. We ran the codeml 847 site model implemented in PAML 4 software¹¹⁹ to infer positive selection on codons under 848 several substitution models (for more details about the models used, see Fisher et al.¹¹³). The 849 850 significance of positive selection was assessed in likelihood ratio tests (LRT). The sites at 851 which positive selection was detected were checked manually and we identified the domain to 852 which they belong, including the specific residue of the LRR when required. In the five gene 853 families identified as displaying significant expansion in oak, 24 groups of oak ultraparalog 854 genes containing up to 28 sequences were identified. Nineteen of these groups had a 855 significant strong signature of positive selection (11 corresponding to LRR-RLKs and 8 to 856 LRR-RLPs, Supplementary Data Set 9). The 11 LRR-RLK groups of ultraparalogs 857 belonged to four previously defined subgroups: SG-VIII-2 (1 group), SG-XI (1 group), SG-858 XIIa (5 groups) and SG-XIIb (4 groups). The two SG-XII subgroups were shown to have 859 undergone species-specific expansion events in a study of 31 angiosperm genomes¹¹³. Most of 860 the SG-XIIa genes described to date are involved in responses to biotic stresses. After manual 861 curation, 260 sites were confirmed to be targets of positive selection (175 in LRR-RLK, and 862 85 in LRR-RLP genes). We found that 78% (205) of the 260 sites were located in the LRR 863 domain (150 in LRRs of LRR-RLK genes and 55 in LRR-RLP genes). An investigation of the 864 precise location of the 150 sites within the LRR of LRR-RLK genes revealed that four amino 865 acids in particular (6, 8, 10 and 11), were more frequently targeted by positive selection (121 866 of the 150 sites, i.e. more than 80%, Supplementary Fig. 8). These variable amino acids lie in the unconserved part of the LXXLXLXX β -sheet/ β -turn structure typical of LRRs that is 867 involved in protein-protein interactions^{120,121}. The residues targeted by positive selection were 868 869 solvent-exposed^{122,123}.

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3.5.8. **Biosynthesis of hydrolysable tannins**

Oak tissues have a very high hydrolyzable tannin (HTs) or gallotannin content, and have been one of the chief sources of HTs for leather tanning and dye manufacture for centuries. We studied the oak genome, to find potential clues to the ability of oak to synthesize HTs, which are esters of gallic acid with a polyol (typically β-D-glucose). Gallic acid is a derivative of the shikimate pathway generated by the dehydrogenation of a 5-dehydroshikimate intermediate¹²⁴. The first committed step in HT biosynthesis is the formation of β-glucogallin 877 (1-O-galloyl- β -D-glucose), which is generated by the esterification of gallic acid and glucose 878 followed by transesterification to generate di-, tri-, tetra-, and pentagalloylglucose 879 Supplementary Fig. 50). Ellagitannins and gallotannins are derived from pentagalloylglucose by the addition of further galloyl residues or oxidation¹²⁵. The UDP-glucose:gallic acid 880 881 glucosyltransferase UGT84A13 has recently been identified as a candidate enzyme in the 882 biosynthesis of β -glucogallin in *Q. robur*¹²⁶. However, the genes and enzymes involved in 883 further esterification steps to generate di-, tri-, tetra-, and pentagalloylglucose remain 884 unknown.

885 A first set of genes potentially involved in the biosynthesis of HTs was annotated on the basis 886 of sequence similarities to genes involved in the chorismate pathway in Arabidopsis thaliana¹²⁷. Uridine diphosphate (UDP) glycosyltransferases (UGTs) mediate the transfer of 887 888 glycosyl residues from activated nucleotide sugars to acceptor molecules, and a superfamily of over 100 genes encoding UGTs has been identified in A. thaliana¹²⁸. Based on the recent 889 characterization of UGT84A13 in *Q. robur*¹²⁶, we focused on the members of the neighboring 890 UGT 74, 75, 83 and 84 families¹²⁹ (http://www.p450.kvl.dk/At_ugts/family.shtml). We 891 892 identified 91 genes potentially associated with HT biosynthesis in the oak genome and we 893 performed phylogenetic analyses of the relationships between these genes and their 894 Arabidopsis orthologs from the chorismate pathway (Supplementary Fig. 51a) and from the 895 UGT 74, 75, 83 and 84 families (Supplementary Fig. 51b). We detected significant 896 expension of the UGT 74, 75 and 83 families in the oak genome, and most of the duplications 897 appeared to be in tandem arrays. Thus, tandem duplications seem to have driven the 898 expansion of these UGT families. Conversely, neither the genes of the UGT84 family nor 899 those involved in the chorismate pathway were expanded in the oak genome.
900 3.5.9. Laccases

The so-called "laccases" (EC 1.10.3.2) are a particularly disparate group of multicopper oxidases (MCOs) in plants, also known as laccase-like multicopper oxidases or simply laccase-like proteins¹³⁰. Laccases can oxidize multiple substrates, whereas other enzymes from the MCO family, such as ascorbate oxidases (EC 1.10.3.3), oxidize only specific substrates. Ascorbate oxidases and laccases are structurally related but different MCOs¹³¹, and ascorbate oxidases are often used as an outgroup in phylogenetic analyses of plant laccases.

907 Little is known about the functions of plant laccases. They can polymerize various phenolic 908 compounds to form insoluble polymers with possible roles in wound healing, plant defense, lignification and the oxidation of seed coat tannins¹³². Three Arabidopsis laccases (AtLAC4, 909 910 11 and 17) were recently shown to play a role in lignin polymerization, and one (AtLAC15) was implicated in the polymerization of flavonoids in the Arabidopsis seed coat^{133,134}. These 911 912 results suggest, at least for lignin polymerization, that laccases are functionally redundant, 913 with multiple mutations required to have a significant effect. In Arabidopsis and poplar, the laccases involved in lignification are targets of miR397a¹³². This micro-RNA downregulates 914 915 laccases and transgenic poplars displaying miR397 overexpression have been produced. 916 These trees displayed low levels of expression for 17 laccases and decrease of up to 40% in the laccase activity of the stem xylem¹³⁵. 917

We found 27 laccase genes in the haplome of *Q. robur* and performed a comparative phylogenetic analysis of the laccase protein sequences from *Q. robur*, *P. trichocarpa*, *E. grandis*, *V. vinifera*, *A. thaliana* and *O. sativa* (**Supplementary Fig. 52**). Sequences were retrieved from Phytozome (https://phytozome.jgi.doe.gov) and the Rice Genome Annotation Project (http://rice.plantbiology.msu.edu/cgi-bin/putative_function_search.pl) by BLAST-p searches. Protein name aliases were used in place of gene model names for *A. thaliana*¹³³ and *P. trichocarpa* (**Supplementary Table 38**). On the phylogenetic tree, *Q. robur* sequences 925 were distributed across the seven phylogenetic groups already described in Arabidopsis. 926 Supplementary Table 39 shows the number of laccases within each phylogenetic group for 927 Arabidopsis, poplar and pedunculate oak. Pedunculate oak was found to have about twice as 928 many laccase genes as Arabidopsis, and about half as many as poplar. Our results therefore 929 suggest that the laccase gene family has undergone expansion in oak, but to a lesser extent 930 than in poplar that however shows a recent whole-genome duplication. Groups #2 and #6 931 displayed a clear expansion of the laccase gene family in tree species relative to *Arabidopsis*, 932 with group #6 displaying stronger expansion in oak. Group #2 corresponds to laccase 933 homologs of ATLAC4, 11 and 17, essential for lignification in Arabidopsis. This biological 934 function is of primary importance for wood cell lignification in trees, so the patterns of 935 duplication and functionalization may differ between trees and herbaceous plants. Group #6 936 contains seven laccases, including AtLAC14.

937 **3.6.** Non-coding RNA prediction and annotation

938 The prediction of long non-coding RNAs was based on 13 RNAseq libraries (listed in 939 Supplementary Table 13). Paired fastq files for the different libraries were aligned with the reference genome fasta file with STAR (version STAR 2.4.0i)¹³⁶. The 13 libraries included 940 941 29 million to 72 million sequence pairs and originated from different tissues and conditions: 942 six from buds, four from roots, one from xylem, one from leaf and one from callus tissues. PCR duplicates were pruned from alignment files (SAMtools rmdup, Version: 1.1)¹³⁷, which 943 were then merged (SAMtools merge, Version: 1.1) before new transcript and gene calling 944 (Stringtie v1.0.1)¹³⁸. The unique alignment rate for read-pairs exceeded 82 %. The Stringtie 945 946 model included 158,714 genes and 215,270 transcripts. The resulting GTF file was processed 947 with FEELnc (https://github.com/tderrien/FEELnc, Version 26/05/2015) to remove known 948 genes and transcripts and to calculate the coding potential of the remaining sequences. The 949 predicted lncRNAs were classified with FEELnc_classifier.pl. FEELnc predicted 16,017

genes and 27,147 transcripts not overlapping with the existing haplome gene model and with more than one exon. Using the FEELnc coding potential program, we identified 12,327 long non-coding RNA genes (corresponding to 19,712 transcripts) and 4,312 new protein-coding genes (corresponding to 7,299 transcripts). FEELnc classified one third of the long noncoding RNA candidates as sense and two thirds as antisense, one third as genic and two thirds as intergenic. A track called 'lncRNA' was added to the genome browser with the FEELnc candidate_feelnc_lncRNA.gtf.lncRNA.gtf file.

957 Other non-coding RNA genes were predicted and annotated with 12 paired RNAseq datasets 958 (listed in Supplementary Table 14). In total, 28,001 loci corresponding to ncRNAs were predicted and annotated with tRNAscan-SE¹³⁹, RNAmmer¹⁴⁰, cmsearch¹⁴¹ with RFAM 959 covariance models¹⁴² and sRNA-PlAn (Supplementary Table 15). Transfer RNA (tRNA) 960 961 and ribosomal RNA (rRNA) genes were predicted with tRNAscan-SE and RNAmmer, 962 respectively. The tRNAscan-SE software predicted 827 tRNAs, including 757 tRNAs 963 decoding standard amino acids, 57 pseudogenes, 12 tRNAs of unknown isotypes and one 964 possible suppressor tRNA. RNAmmer software found 82 rRNA loci, including 13 large 965 subunit (LSU) rRNA genes, 20 small subunit (SSU) rRNA genes and 49 rRNA 5S genes. We 966 used the cmsearch program from the Infernal suite with a selection of covariance models 967 relating to families found in eukaryotic genomes, including tRNAs, rRNAs, small nucleolar 968 RNAs (snoRNAs), small nuclear RNAs (snRNAs), miRNAs, SRP RNAs, RnaseMRP RNAs, 969 telomerase RNAs and Vault RNAs. The Infernal cmsearch program found no LSU, but 970 predicted 14 rRNA 5.8S loci, 52 SSU rRNA loci, and 65 rRNA 5S loci. Thus, considering all 971 rRNA predictions, 136 loci in total were predicted and annotated as ribosomal RNA, 972 including 70 rRNA 5S and 61 LSU and/or SSU rRNAs. In total, 44 predicted rRNA 5 S genes 973 were common to the cmsearch and RNAmmer analyses. Seven of the 13 LSU loci predicted 974 by RNAmmer overlapped the 5.8S rRNA predictions calculated by cmsearch. The Infernal

975 cmsearch program predicted 815 tRNAs. In total, 852 tRNAs were predicted, 790 of which 976 were detected by both tRNAscan-SE and Infernal cmsearch; 25 were specific to Infernal 977 cmsearch and 37 were specific to tRNAscan-SE. Among the other ncRNA genes, a total of 978 412 C/D box and 74 H/ACA box snoRNA genes were predicted, corresponding to 73 and 17 979 different families of snoRNA, respectively. With 146 predicted candidates, the C/D box 980 snoRNA71 family was the most heavily represented. An analysis of snoRNA gene 981 organization showed that 190 of these genes were organized into 59 clusters containing two to 982 11 snoRNA genes. Other snoRNA genes were also identified by eye in the clusters 983 (Supplementary Fig. 14, sequence of a snoRNA H/ACA gene conserved in A. thaliana). The 984 cmsearch program also predicted 263 pre-miRNA loci, two RNase MRP RNA genes, 31 SRP 985 RNA genes, 225 spliceosomal snRNA genes including 34 U1 snRNA genes, one U11 snRNA 986 gene, 55 U2 snRNA genes, one U12 snRNA gene, 33 U4 snRNA genes, 24 U5 snRNA genes, 987 64 U6 snRNA genes and 13 U6atac snRNA genes. We retained only one of the pre-miRNA 988 predictions made on both strands at same positions, resulting in the consideration of 204 pre-989 miRNA loci in the set of pre-miRNA gene predictions. We also predicted miRNA genes with 990 sRNA-PlAn (source code available as a workflow at https://forgemia.inra.fr/genotoul-991 bioinfo/ngspipelines/tree/master/workflows/srnaseq) on the 12 paired small RNAseq 992 datasets. sRNA-PlAn implements a model of miRNA biogenesis. Loci are built by 993 considering the regions of the genome to which reads produced by sRNA-seq experiments 994 map. Candidate loci are subjected to the miRNA prediction procedure, which considers the 995 expected pre-miRNA stem-loop structure, the size of the pre-miRNA sequence, the size of 996 pre-miRNA loops (bulges, internal loops, stem loop), the size of the most represented 997 sequence (20-24 nt), the alignment of this most represented sequence with the stem of the pre-998 miRNA and the expected expression profile of the pre-miRNA. A score is assigned to each 999 predicted pre-miRNA locus, taking into account the characteristics described above. Each 1000 predicted pre-miRNA locus is then subjected to an annotation procedure in which it is aligned with miRBase¹⁴³ and RFAM¹⁴² sequences with BLAST+¹⁴⁴, to differentiate between known 1001 1002 ncRNA families and new candidate miRNA families. In total, 26,109 miRNA loci were 1003 predicted by sRNA-PlAn, from which 1,508 mature miRNA loci predicted by sRNA-PlAn 1004 with a high score were annotated as miRNAs on the basis of strong similarities (one error 1005 allowed) to sequences in the miRBase or RFAM databases (fasta files). We found that 145 of 1006 the related pre-miRNAs were specific to the RFAM cmsearch program and that 64 of these 1007 pre-miRNAs encoded members of the mir-69 gene family. Interestingly, 59 of the pre-1008 miRNA loci predicted by cmsearch contained one or both of the mature miRNAs predicted 1009 and annotated with sRNA-PlAn. Different tracks relating to ncRNA predictions/annotations 1010 were added to the genome browser, according to the software used.

1011 Finally, the 12 paired small RNAseq datasets as well as ncRNA predictions, lncRNA genes 1012 and TEs were used to assign expression evidence to the whole set of predicted noncoding 1013 regions. Reads obtained from small RNAseq datasets were mapped onto the genome with Bowtie2¹⁴⁵, using default parameters and retaining only one alignment. Transcription 1014 1015 evidence and count data were obtained with Featurecounts for each predicted non-coding RNA locus¹⁴⁶. Predicted lncRNAs were used to confirm expression at predicted non-coding 1016 loci and to identify clusters of shorter ncRNA genes. SAMtools¹³⁷ and BEDtools¹⁴⁷ functions 1017 were used to manipulate alignments and to identify regions of overlap between the predicted 1018 1019 lncRNA and ncRNA genes. We found that 212 of the predicted lncRNA genes overlapped 1020 annotated ncRNA loci (strand not considered). Fourteen overlapped 14 rRNA predictions, 1021 three overlapped six SRP RNA predictions, with two lncRNAs containing two and three SRP 1022 RNA predictions, respectively; 114 overlapped 124 mature miRNA or pre-miRNA 1023 predictions, with six lncRNAs overlapping two or more pre-miRNA predictions; 34 1024 overlapped 82 snoRNA, with 19 of lncRNAs overlapping two to 11 snoRNA predictions; 22 1025 overlapped 22 tRNA loci; seven overlapped 10 U6 snRNA predictions, with one lncRNA 1026 overlapping four U6 snRNA predictions; one overlapped one U6atac snRNA prediction; three 1027 overlapped three U1 snRNA predictions; six overlapped 13 U2 snRNA predictions, with five 1028 lncRNAs overlapping two to five U2 snRNA predictions; five overlapped six U4 snRNA 1029 predictions, with one lncRNA overlapping two U4 snRNA predictions. The content of small 1030 RNAseq datasets was analyzed for non-coding elements, such as predicted lncRNAs, other 1031 predicted/annotated ncRNAs and predicted TEs (Supplementary Table 16). Bowtie2 aligned 1032 83.34% of the 383,274,162 reads on scaffolds. Using Featurecounts with non-coding 1033 elements, such as TEs, lncRNA and ncRNA annotations and predictions, we were able to 1034 assign a total of 231,211,802 reads, corresponding to 72.4 % of the mapped reads, to 1035 annotated non-coding elements.

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1037 4. Mutational landscape

1038 **4.1. Estimate of genetic diversity and** $\pi 0/\pi 4$ ratio

1039 The genetic diversity of oak (π) was 0.011 at synonymous sites (π_4), and 0.005 at non-1040 synonymous sites (π_0), with a mean π_0/π_4 ratio of 0.44 (**Supplementary Table 9**). For 1,176 1041 manually curated genes, we recovered the $\pi_0/\pi 4$ ratio equals to 0.43 ($\pi_0 = 0.00429$ and $\pi_4 =$ 1042 0.00990). Oak has a higher genetic diversity and π_0/π_4 ratio (Fig. 2a) than the other woody perennial species studied by Chen et al.¹⁴⁸. Further comparisons between "Expanded", 1043 1044 "Contracted", and "Unchanged" gene families showed that π_0 estimates were significantly higher for expanded gene families in oak (0.007, p-value $< 2 \times 10^{-16}$) whereas π_4 values were 1045 1046 similar for all types of gene families (0.012, Supplementary Fig. 54), resulting in a higher 1047 π_0/π_4 ratio (0.56). Contracted family genes had a significantly lower π_0/π_4 ratio (0.30) than unchanged families (0.32, *p*-value= 5.2×10^{-3}). TDGs also had a higher π_0/π_4 ratio (0.53), and 1048

an even higher π_0/π_4 was found in the families expanded in oak (0.62). Similar estimates were 1049 1050 obtained from the analysis of the pool-seq dataset, i.e. average $\pi_0/\pi 4$ of 0.50 ($\pi_0 = 0.00538$ 1051 and $\pi_4 = 0.0108$), increasing to 0.59 for TDG, and 0.60 for expanded, 0.30 for contracted and 1052 0.32 for unchanged genes. Higher π_0/π_4 values suggest a potential accumulation of deleterious 1053 mutations in expanded gene families relative to contracted or unchanged families. We 1054 calculated the frequency of mutations (as unnormalized pairwise differences) likely to cause 1055 protein malfunction (e.g. premature stop codons, start/stop codon changes). Genes from 1056 expanded families displayed significantly more potentially deleterious mutations (mean =0.23, *p*-value< 2×10^{-16}) than those from contracted (0.09) or unchanged families (0.06). 1057

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4.2. Detection of somatic mutations

1059 We compared the three libraries L1, L2 and L3 (i.e. 6 pairwise combinations, Supplementary 1060 Table 20) and detected 61 reliable somatic mutations. A total of 46 somatic mutations were 1061 completely absent from the poolseq dataset (40 SNPs), or and had MAFs below 0.5% (6 1062 SNPs), *i.e.* below the minimum threshold used to exclude sequencing errors in the poolseq 1063 dataset). Considering our high sequencing depth and the number of individual pooled (20 1064 genotypes), each allele is expected to be near 2.5%. As a consequence, low allele frequency variants are expected to be related to sequencing errors (estimated at 2.4%, see 1065 1066 Supplementary Figure 25). Thus, to be conservative, we filtered out all candidate somatic 1067 mutations with an allele frequency above 0.005. As a result, 75% of the somatic mutations (46/61) could be considered to be detected exclusively in the "3P" accession (Supplementary 1068 1069 Table 5).

1070 Noteworthy, one of the 40 somatic mutations detected exclusively in "3P" was found in a 1071 gene coding sequence: Sc0000066_1207928 in Qrob_T0204900.2 corresponded to a member 1072 of the large cytochrome P450 superfamily encoding a protein of the CYP4/CYP19/CYP26 subfamilies with annotations relating to secondary metabolite biosynthesis, transport and
catabolism, lipid transport and metabolism. This mutation was synonymous (ACC->ACT,
corresponding to a threonine residue in the protein).

1076 **5. Comparative and evolutionary genomics**

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5.1. Macroevolutionary analysis

1078

5.1.1. Oak karyotype evolution and genome organization

1079 Considering grape to be the closest modern representative of the n=21 rosid ancestor (derived 1080 from a post- γ ancestor with 7 protochromosomes (shown in color on the *y*-axis of the dotplots 1081 of **Supplementary Fig. 16**) the comparisons between grape-eucalyptus and grape-watermelon 1082 shows a clear 1:2 relationships, while that between grape-coco, grape-peach and grape-oak 1083 genomes shows a clear a 1:1 relationships see dotplot diagonals in each chart, shown with 1084 green circles in **Supplementary Fig. 16**.

1085 5.1.2. Gene family expansion/contraction in oak

1086 For the total of 541,339 gene models across the 15 species (Supplementary Table 21, 1087 Supplementary Fig. 17) plus Q. robur, 435,095 were classified into 36,844 orthogroups 1088 (gene families) (Supplementary Data Set 3 sheet #1), with 106,444 genes remaining 1089 singletons after clustering (Supplementary Table 7). In total, 4860 orthogroups were 1090 common to all species. For the 25,808 oak proteins, 22,498 clustered into 11,813 orthogroups, 1091 479 of which were oak-specific and contained 1,737 oak proteins. There were also 3,310 1092 singleton proteins for oak. From the 36,844 orthogroups 524 and 72 were found to be 1093 expanded and contracted in oak, respectively (Supplementary Data Set 3 sheet #2 and #4). 1094 A total of 154 orthogroups were specific to oak (Supplementary Fig. 18), whereas 65 were 1095 common to all species (Supplementary Fig. 18). We found that 73% of the genes within expanded orthogroups were tandemly duplicated genes (TDGs), this percentage increasing to
99% if we also included long distance-duplicated genes (LDGs), whereas the 72 contracted
orthogroups contained only 47% TDGs (Supplementary Fig. 19).

1099

5.2. Identification of tandemly duplicated genes in oak

1100 Speciation and duplication events in the pedunculate oak genome were identified using the K_s 1101 distribution of orthologous gene pairs between oak and peach (green bars in Supplementary 1102 Fig. 20) and paralogs in oak (purple bars in Supplementary Fig. 20), respectively. The 1103 oak/peach ortholog K_s distribution defines the position of the speciation event between these 1104 two species, with a single ancestral triplication event (γ) common to grape, peach, cocoa and 1105 oak and predating the speciation event. The burst of tandem duplicates highlighted by the 1106 purple K_s peak occurred after oak/peach speciation and appears to be an oak-specific event. 1107 The dot plot representation of tandemly duplicated genes (TDGs) in oak is depicted in 1108 Supplementary Fig. 21. We identified 9,189 TDG (Supplementary Data Set 4) using the 1109 threshold and methodology presented in the method section. They were validated based on (i) 1110 the comparison with polymorphism of allelic gene pairs (Supplementary Fig. 22) and (ii) 1111 sequence coverage analysis (Supplementary Fig. 23). Besides, we identified 8,797 genes as 1112 long distance duplicated genes (LDGs) and 7,822 genes as single genes (SGs) 1113 (Supplementary Data Set 4).

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5.3. Challenges in the identification of genes related to tree habit

The increasingly rapid rate at which full genome sequences are being published opens up exciting possibilities, but the 16 species for which suitable genome sequences were available for this study represents only a small proportion of the worldwide diversity of plants (there are currently ~350,000 accepted angiosperm species (http://www.theplantlist.org/). It was recently estimated that almost 50% of vascular plants, most of which are angiosperms, are

woody¹⁴⁹. There are probably, therefore, many genomic changes associated with shifts in 1120 1121 growth form not captured in our analyses. Given the modest number of genomes included in 1122 this study, it remains unclear whether the patterns highlighted here can be generalized to 1123 larger numbers of species and clades. It is also unclear whether the same sets of expanding 1124 and contracting gene families would be identified in all evolutionary transitions from 1125 herbaceous to woody forms. Fortunately, even with the limited number of genomes available, 1126 it is possible to identify the branch points within the phylogeny at which additional targeted 1127 sequencing would help to provide an answer to this question. The most dynamic aspects of 1128 growth form shifts within the angiosperm phylogeny have occurred within the eudicots, a group consisting largely of rosids and asterids^{149–152}. The sequencing of genomes for 1129 1130 additional tree species in this part of the phylogeny would be particularly informative. Fitziohn et al.¹⁴⁹ used the distribution of growth forms from Zanne et al.¹⁵² to estimate the 1131 1132 proportion of woody taxa across vascular plants at the genus, family and order levels. The clades highlighted by Fitzjohn et al.¹⁴⁹ as both variable in growth form (defined here as clades 1133 1134 with 30-70% of species considered to be woody according to the strong prior) and diverse 1135 (defined here as containing >10 species), comprise 470 genera, 41 families and 12 orders. 1136 Paired comparisons of close relatives in these clades, ideally genera with both woody and 1137 herbaceous members, would make it possible to determine whether gene expansions in R-1138 gene families are correlated with evolutionary shifts in growth form. It is clear that certain 1139 clades are extraordinarily variable in terms of growth habit, but it seems unlikely that growth habit per se drives expansions and contractions in R-gene families. Instead, with their longer 1140 1141 lifespans, woody species probably accumulate a greater pathogen load than herbaceous taxa. 1142 It would therefore appear reasonable to consider longevity as a driver of these functional gene 1143 shifts, and growth habit as a correlate of such differences in life history.

1144 We examined the full genome sequences currently or soon to be available for eudicots (as 1145 reported in (https://genomevolution.org/wiki/index.php/Sequenced_plant_genomes) as of 18 1146 December 2016), to identify future targets building on existing genomic resources. We had 1147 access to genome sequences for 46 herbaceous and 27 woody species from 18 orders 1148 (Supplementary Table 40 and Supplementary Fig. 55), 14 of which display growth habit 1149 variation (defined here as clades with 30-70% of woody species according to the strong prior) and sufficient diversity (defined here as >10 species), accordin to Fitzjohn et al.¹⁴⁹. The 1150 1151 sequencing of herbaceous species from four variable genera (Nicotiana, Linum, Genlisea, 1152 Solanum) is underway, but full genome sequences for both a woody and a herbaceous species 1153 from the same genus have yet to be published. In Fabaceae, complete genome sequences have 1154 been released for herbaceous species from eight genera and for one woody species from the 1155 genus Cajanus. Rosaceae also includes four woody and one herbaceous species (Fragaria) for 1156 which complete genome sequences have been released. Three variable orders have fully 1157 sequenced herbaceous species (Ranunculales, Caryophyllales, Solanales). In addition, 1158 Lamiales has three herbaceous and one woody (Fraxinus) species for which full genome 1159 sequences have been obtained, and Fabales has eight herbaceous and one woody (Cajanus) 1160 species with full genome sequences. Additional genome sequences for species from any of 1161 these clades (ideally within genera), considered together with the oak genome sequence, 1162 would improve our understanding of the evolution of genomic features favoring a long 1163 lifespan and woodiness in plants.

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5.4. Gene ontology (GO) enrichment analysis

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5.4.1. **GO term enrichment in three categories of genes**

We assigned a total of 3,433 GO terms (**Supplementary Table 41**): 1,179 for molecular function (MF), 1,867 for biological process (BP) and 387 for cellular component (CC). At 1168 least one GO term was assigned to 16,820 of the 25,808 oak gene models (65.2%). The mean 1169 number of GO terms per gene was 3.73 (ranging from 1 to 18) (**Supplementary Fig. 56**) and 1170 gene counts per GO term are provided in **Supplementary Fig. 57**. We found that 332 GO 1171 terms were associated with only one gene.

GO term enrichment was compared between three categories of genes: (i) TDGs (i.e. genes located in close proximity, see section 4.2 for the method), (ii) LDGs and (iii) SGs. The number of significant GO terms (*p*-value < 0.05) was 97 for TDGs, 144 for LDGs and 240 for SGs (**Supplementary Table 41**).

1176 For TDGs (Supplementary Data Set 8 sheet #1), the best supported GO terms (in terms of 1177 *p*-value and fold-enrichment) highlighted gene products involved in 'protein phosphorylation' (GO:0006468, $P < 1 \times \times 10^{-30}$), 'signal transduction' (GO:0007165, $P < 1 \times 10^{-30}$), 'recognition of 1178 pollen' (GO:0048544, $P < 1 \times 10^{-30}$), 'oxidation-reduction process' (GO:0055114, $P = 7.2 \times 10^{-29}$), 1179 'metabolic process' (GO:0008152, $P=2.2\times10^{-17}$), 'chitin catabolic process' (GO:0006032, 1180 $P=1.9\times10^{-13}$), 'response to biotic stimulus' (GO:0009607, $P=4\times10^{-12}$), 'cell wall 1181 macromolecule catabolic process' (GO:0016998, $P=2.3\times10^{-10}$), 'response to oxidative stress' 1182 (GO:0006979, 10^{-8}), 'drug transmembrane transport' (GO:0006855, $P=1.9\times10^{-7}$) and 'defense 1183 response' (GO:0006952, $P=9\times10^{-7}$). Thus, gene products executing activities related to 'ADP 1184 binding' (GO:0043531, $P < 1 \times 10^{-30}$), 'transferase activity' (GO:0016758, $P = 3.8 \times 10^{-28}$), 'heme 1185 binding' (GO:0020037, $P=1.3\times10^{-27}$), 'protein kinase activity' (GO:0004672, $P=1.2\times10^{-26}$), 1186 'oxidoreductase activity' (GO:0016705, $P=5.8\times10^{-23}$), iron ion binding ('GO:0005506', 1187 $P=1.1\times10^{-17}$), 'chitinase activity' (GO:0004568, $P=2.1\times10^{-13}$), 'protein serine/threonine 1188 kinase activity' (GO:0004674, $P=2.4\times10^{-13}$), and 'nutrient reservoir activity' (GO:0045735, 1189 $P=1.3\times10^{-10}$) were the most frequently detected. Membrane-bound (LRR-RLKs and LRR-1190 1191 RLPs) and cytosolic (NB LRR) receptors, together with UDP-glycosyltransferase,

cytochrome P450, chitinase, peroxidase, and psathogenesis-related protein were among themost frequently detected proteins corresponding to the BP and MF ontologies.

1194 By contrast, a very different molecular signature was obtained for LDGs (Supplementary Data Set 8 sheet #2). The best supported GO terms included 'regulation of transcription, 1195 DNA-templated' (GO:0006355, $P=1.20\times10^{-10}$), 'protein dephosphorylation' (GO:0006470, 1196 $P=3.50\times10^{-9}$, 'small GTPase-mediated signal transduction' (GO:0007264, $P=1.30\times10^{-8}$), 1197 'microtubule-based process' (GO:0007017, $P=2.60\times10^{-8}$), 'translation' (GO:0006412, 1198 1199 $P=1.30\times10^{-7}$), 'response to heat' (GO:0009408, $P=3.50\times10^{-7}$), 'protein folding' (GO:0006457, P=1.70×10⁻⁶), 'fatty acid biosynthetic process' (GO:0006633, P=2.20×10⁻⁶ 1200 'biosynthetic process' (GO:0009058, $P=3.60\times10^{-5}$ and 'protein polymerization' 1201 (GO:0051258, $P=6.90\times10^{-5}$). Thus, gene products with activities relating to 'protein 1202 serine/threonine phosphatase activity' (GO:0004722, $P=3.70\times10^{-13}$), 'DNA binding' 1203 (GO:0003677, $P=1.70\times10^{-12}$ 'sequence-specific DNA binding' GO:0043565, $P=1.90\times10^{-11}$), 1204 'GTP binding' (GO:0005525, $P=9.30\times10^{-11}$), 'structural constituent of ribosome' 1205 (GO:0003735, $P=2.60\times10^{-9}$), 'transcription factor activity, sequence-specific DNA binding' 1206 (GO:0003700, $P=5.90\times10^{-9}$), 'NADH dehydrogenase (ubiquinone) activity' (GO:0008137. 1207 $P=1.30\times10^{-8}$), 'GTPase activity' (GO:0003924, $P=2.40\times10^{-8}$), 'structural constituent of 1208 cytoskeleton' (GO:0005200, $P=8.40\times10^{-7}$), and 'microtubule binding' (GO:0008017, 1209 $P=2.80\times10^{-6}$) were the most frequently detected. Protein phosphatases, proteins with DNA-1210 1211 binding and homeobox domains, transcription factors, elongation factors, ribosomal proteins, 1212 microtubule-associated proteins, and DNA gyrases were among the most widespread proteins 1213 corresponding to the BP and MF ontologies.

For SGs (**Supplementary Data Set 8 sheet #3**), the best supported GO terms concerned 'DNA replication' (GO:0006260, $P=5\times10^{-13}$), 'transcription, DNA-templated' (GO:0006351, $P=8.6\times10^{-13}$), 'DNA repair' (GO:0006281, $P=1.1\times10^{-11}$), 'RNA processing' (GO:0006396,

 $P=1.2\times10^{-9}$), 'photosynthesis' (GO:0015979, $P=1.2\times10^{-9}$), 'pseudouridine synthesis' 1217 (GO:0001522, $P=1.5\times10^{-9}$), 'DNA recombination' (GO:0006310, $P=3.9\times10^{-9}$), 'protein 1218 ubiquitination' (GO:0016567, $P=9.8\times10^{-7}$), 'glycerol ether metabolic process' (GO:0006662, 1219 $P=1.3\times10^{-6}$) and 'translation' (GO:0006412, $P=3.4\times10^{-6}$). Thus, gene products with activities 1220 relating to 'binding' (GO:0005488, P=4.7×10⁻²⁴), 'RNA binding' (GO:0003723, P=6.2×10⁻ 1221 ²⁰), 'nucleic acid binding' (GO:0003676, $P=2.4\times10^{-19}$), 'zinc ion binding' (GO:0008270, 1222 $P=6.9\times10^{-16}$), 'metal ion binding' (GO:0046872, $P=4.8\times10^{-10}$), 'pseudouridine synthase 1223 activity' (GO:0009982, $P=1.5\times10^{-8}$), 'DNA-directed RNA polymerase activity' (GO:0003899, 1224 $P=1.9\times10^{-8}$), 'nucleotide binding' (GO:0000166, $P=4.1\times10^{-8}$), 'ubiquitin-protein transferase 1225 activity' (GO:0004842, $P=8\times10^{-8}$), 'threonine-type endopeptidase activity' (GO:0004298, 1226 $P=1.1\times10^{-5}$), 'DNA helicase activity' (GO:0003678, $P=4.5\times10^{-5}$) and 'DNA binding' 1227 (GO:0003677, $P=9.5\times10^{-5}$) were the most frequently detected. Zinc finger and DNA repair 1228 1229 proteins, as well as DEAD/DEAH box helicase, RNA pseudouridylate synthase and RNA 1230 polymerase were among the most frequently detected proteins corresponding to the BP and 1231 MF ontologies.

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5.4.2. GO term enrichment in orthogroups expanded in pedunculate oak

1233 The 524 orthogroups expanded in oak comprise 5,910 genes (3 to 359 genes per orthogroup, 1234 with a mean of 11.3 genes per orthogroup, **Supplementary Fig. 58**). In total, 366 orthogroups 1235 were annotated with at least one GO term. The number of GO terms per gene family ranged from 1 to 17 (mean value, 2.89). We found that 4,217 of the 5,910 genes (71.4%) were 1236 1237 annotated with at least one GO term (Supplementary Table 42). The annotation used 3,433 1238 unique GO terms, including 1,722 singletons (GO terms used only once) (Supplementary Fig. 59). We identified 58 significantly enriched GO terms (33 MF, 17 BP and 8 CC) 1239 (Supplementary Data Set 8 sheet #4) in orthogroups displaying expansion in oaks. We 1240 compared sample counts (numbers of genes annotated with particular GO terms among the 1241

1242 genes belonging to the orthogroups expanded in oak) with genome counts (number of genes 1243 annotated with particular GO terms among the 25,808 oak gene models; Supplementary Fig. 1244 60). The enriched term with the best statistical support was 'protein kinase activity' (GO:0004672, $P < 10^{-30}$), which was attributed to 726 genes (in the 524 expanded orthogroups) 1245 1246 of the 1,556 genes found in the 25,808 oak gene models, corresponding to two-fold 1247 enrichment. These 726 genes belonged to 31 orthogroups containing genes encoding both 1248 cytosolic (NB-LRRs) and membrane (LRR-RLKs, LRR-RLPs) receptors of the innate 1249 immune system (i.e. R-genes). This overrepresentation of R-genes was also supported by the 1250 enrichment of the orthogroups in the following annotations: 'protein serine/threonine kinase activity' (GO:0004674), 'protein binding' (GO:0005515), 'polysaccharide binding' 1251 (GO:0030247), 'ADP binding' (GO:0043531), 'protein phosphorylation' (GO:0006468), 1252 'signal transduction' (GO:0007165), 'recognition of pollen' (GO:0048544), all with $P < 10^{-30}$ 1253 1254 and fold-enrichments of 1.7 to 3.9 (for 'ADP binding'). The highest fold-enrichment (about 1255 4.4) was observed for the MF 'thioredoxin-disulfide reductase activity' (GO:0004791, $P=8.4\times10^{-5}$) and the BP 'removal of superoxide radicals' (GO:0019430, $P=2.9\times10^{-5}$), with 1256 seven genes annotated as pyridine nucleotide-disulfide oxidoreductases. 1257

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5.4.3. GO enrichment within the gene families expanded in woody perennial trees relative to herbaceous species

Overall, 18,855 of the 36,844 othoMCL orthogroups (**Supplementary Data Set 3 sheet #1**) (51.2%) were annotated with at least one GO term, with 16,703 orthogroups annotated for molecular function (MF), 11,495 for biological process (BP) and 5,073 for cellular component (CC). In total, 3,936 unique GO terms were used in the annotation. Of the 126 orthogroups expanded in "trees" (**Supplementary Data Set 7 sheet #2**), 108 were annotated with GO terms used in the GO term enrichment analysis. We detected significant enrichment for 61

GO terms (38 MFs, 19 BPs and 4 CCs, Supplementary Table 43 and Supplementary Data Set 8 sheet #5).

1268 The functions of the set of gene families expanded in woody species were identified against the background of all orthogroups. The degree of orthogroup size expansion for statistically 1269 1270 significant GO terms, represented by fold-enrichment in woody perennials is depicted in 1271 Supplementary Fig. 7. The term with most statistical support was 'apoptotic process' (GO:0006915, $P=7.4\times10^{-14}$). It was found in 10 of the 126 expanded orthogroups, but only 37 1272 1273 of the total number of 36,844 orthogroups, giving a fold-enrichment of 79 (Supplementary 1274 Fig. 7). These 10 clusters included R-genes with a characteristic NB-ARC domain. 'ATP binding' (GO:0005524, P=2.8×10⁻¹⁰), 'ADP binding' (GO:0043531, P=10⁻⁹), 'protein 1275 serine/threonine kinase activity' (GO:0004674, $P=6.4\times10^{-7}$), 'protein tyrosine kinase activity' 1276 (GO:0004713, P=1.4×10⁻⁶), 'protein phosphorylation' (GO:0006468, P=1.8×10⁻⁶) 'DNA 1277 integration' (GO:0015074, P=3.2×10⁻⁶), 'polysaccharide binding' (GO:0030247, P=1.7×10⁻⁵), 1278 transmembrane signaling receptor activity' (GO:0004888, $P=9.6\times10^{-5}$), 'innate immune 1279 response' (GO:0045087, P= 1.4×10^{-4}) and 'recognition of pollen' (GO:0048544, P= 2×10^{-4}) 1280 1281 ranked among the next most significant GO terms, with fold-enrichments ranging from 7 up 1282 to 83.5 for 'protein serine/threonine kinase activity'. The orthogroups concerned included 1283 almost exclusively cytosolic and membrane receptors of the innate immune system (Supplementary Data Set 8 sheet #5). For instance, the 10 most frequent orthogroups 1284 1285 (orthogroups #1000, 1004, 1021, 1084, 1006, 1010, 1016, 1017, 1037, 1003) cited 115 times 1286 in a total of 367 occurrences, i.e. over 30%, corresponded to the two major types of plant 1287 receptors: leucine-rich repeat-receptor-like kinase/receptor-like proteins (LRR-RLKs, LRR-1288 RLPs) and nucleotide-binding leucine-rich repeat proteins (NB-LRRs).

1289 **6. Web resources**

Genome data	Web access
Oak genome assembly PM1N	Download: https://urgi.versailles.inra.fr/download/oak/Qrob_PM1N.fa.gz
(haploid version: 12	Blast: https://urgi.versailles.inra.fr/blast
pseudomolecules + 538	Pseudomolecule: https://urgi.versailles.inra.fr/WebApollo_oak_PM1N/PseudoMolecule.html
unassigned scaffolds)	JBrowse: https://urgi.versailles.inra.fr/WebApollo_oak_PM1N/jbrowse
	Intermine: https://urgi.versailles.inra.fr/OakMine_PM1N/begin.do
Oak genome assembly V2_2N	Download: https://urgi.versailles.inra.fr/download/oak/Qrob_V2_2N.fa.gz
(diploid version 2)	Blast: https://urgi.versailles.inra.fr/blast
	JBrowse: https://urgi.versailles.inra.fr/WebApollo_oak_V2/jbrowse/
Oak genome assembly V1_2N	Download: https://urgi.versailles.inra.fr/download/oak/Qrob_V1_2N.fa.gz
(diploid version 1, ¹⁹)	Blast: https://urgi.versailles.inra.fr/blast
Oak transcriptome (de novo	Download: https://urgi.versailles.inra.fr/download/oak/OCV4_assembly_final.fsa.gz
assembly, ²³)	Blast: https://urgi.versailles.inra.fr/blast
Oak protein-coding sequences	Download CDS (aa) https://urgi.versailles.inra.fr/download/oak/Qrob_PM1N_CDS_aa_20161004.fa.gz
predicted on PM1N (haploid	Download CDS (nt) https://urgi.versailles.inra.fr/download/oak/Qrob_PM1N_CDS_nt_20161004.fa.gz
version)	Blast: https://urgi.versailles.inra.fr/blast

1290

1291 **7. Data availability**

The oak haploid genome assembly and corresponding annotation have been deposited in the European Nucleotide Archive under project accession code PRJEB19898. Other sequence release data are indicated in Supplementary tables 1, 13, 14 and 19 and Supplementary Data Set 10. We also invite readers to download data stored at the URLs indicated in section 6 (Web resources) as well as in the oakgenome web site: http://www.oakgenome.fr.

1297 **8. References**

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 phylogenetic analysis. *Mol. Biol. Evol.* 17, 540–552 (2000).
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- 1713
- 1714

1715 9. Supplementary Data Sets

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1717 Supplementary Data Set 1 List of 25,808 oak gene models with their annotations.
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Supplementary Data Set 2 Mapping data used to anchor the scaffolds onto the oak
genetic linkage map. Sheet #1: List of 5,589 mapped markers. Sheet #2: Subset of 2,615
markers matching sequence scaffolds, classified by categories according to Supplementary
Table 27. Sheet #3: syntenic relationships between oak markers and peach gene models.
Sheet #4 ordered scaffolds along the 12 chromosomes.

1723 Supplementary Data Set 3 List of orthogroups (orthoMCL analysis) and expanded gene 1724 families (CAFE analysis) in pedunculate oak. Sheet #1 List of clusters obtained with 1725 orthoMCL. The family P-value is provided by CAFE and corresponds to the probability of 1726 observing the data (orthogroup size distribution between taxa). Orthogroups with larger size 1727 variance are expected to have lower P-values. The Qr P-value is the oak branch-specific P-1728 value. It corresponds to the probability of transitions between the parent and child family sizes 1729 for the oak branch. A low P-value indicates a rapidly evolving orthogroup. These data were 1730 provided by CAFE. Sheet #2 List of clusters expanded in oak. Sheet#3 list of outstanding 1731 outlier clusters expanded in oak. Sheet#4 list of clusters contracted in oak.

Supplementary Data Set 4 List of gene categories. Sheet #1: tandemly duplicated genes
(TDG). Sheet #2: list of TDG relationships. Sheet #3: long distance-duplicated genes (LDG).
Sheet #4: singleton genes (SG).

Supplementary Data Set 5 Classification of NB-LRR-related genes. List of oak NB-LRRrelated genes. For each gene, the gene model ID and the proteinID are provided. NB-LRR
genes were classified into categories according to their canonical domains, i.e. CC, coiledcoil; LRR, leucine-rich repeat; NB, nucleotide-binding; RPW8, resistance to powdery mildew

protein; TIR, Toll interleukin receptor-like; X, putative integrated decoy. The position of the genes (gene start and gene end) on pseudomolecules or unassigned scaffolds is indicated. The orthogroup ID is also provided for the orthoMCL analysis, together with the family expansion/contraction status (oak vs. other species, 1% threshold), and tandem duplication status.

Supplementary Data Set 6 Classification of RLK-related genes. Sheet #1: list of RLKrelated genes from oak, Arabidopsis and rice - phylogeny and assignment. Sheet #2: subgroups of RLK-related genes and subgroups within LRR-RLK from sheet #1. Sheet #3 data from Fischer et al.¹¹³ for comparison with oak. Sheet #4: list of oak genes from sheet #1, with their classification into orthoMCL orthogroups and their status (in a tandem array or not). Sheet #5: results from sheet #4.

Supplementary Data Set 7 Summary of orthogroups expanded in 'trees'. Sheet #1: *P*value and FDR for all orthoMCL orthogroups. Sheet #2: orthogroups expanded (FDR<0.05)
in 'trees'. Shee #3: list of outstanding 'tree' orthogroups and their functional annotations.
Sheet #4: orthogroups expanded [contracted] in herbaceous species [trees].

Supplementary Data Set 8 Summary of the Gene Ontology (GO) term analysis showing
significantly enrichment in GO terms for molecular functions (MF), biological processes
(BP), and cellular components (CC). Sheet #1: tandem duplicated genes (TDGs). Sheet #2:
long distance-duplicated genes (LDGs). Sheet #3: singletons (SGs). Sheet #4: orthogroups
expanded in oak. Sheet #5: orthogroups expanded in woody perennials. Sheet #6: expanded
orthogroups in herbaceous species. *P*-values are from Fisher's exact tests.

Supplementary Data Set 9 Footprint of selection in RLK-related genes. Sheet #1: ID of
ultraparalogous genes with their association group, annotation, orthoMCL orthogroup and
- 1762 label from **Fig. 3d and 4b**. Sheet #2: codeml results for 24 groups of ultraparalogs. Sheet #3:
- 1763 results of all the manually validated sites (domain plus position in the LRR motif).

1764 Supplementary Data Set 10 List of pedunculate oak BAC clones used in this study. Sheet

- 1765 #1: list of sequenced BAC clones and matching scaffolds on the diploid version of the oak
- 1766 genome sequence. Sheet #2: gene annotation on the sequenced BACs.

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10. Supplementary Tables

Supplementary Table 1 List of genomic and cDNA libraries used to sequence and annotate the pedunculate oak genome accession "3P".

DNAseq: genomic libraries used to sequence the oak genome. RNAseq: cDNA libraries used to annotate the oak genome.

Project	Material	Filename	# reads	# bases	Library	Technology	Accession ID	Diploid genome (1.5G/2C) coverage
				DNAse	q			
Qr	А	Oak_7_1_sequence.fastq	62418280	4681371000	Paired-end	Illumina (GAIIx)	SRX739064	3x
Qr	В	Oak_5Kb_MatePair_6_1_sequence.fastq	56012498	2800624900	Mate-pairs 5 Kb	Illumina (GAIIx)	SRX739054	2x
AWU	А	AWU_AOSC_3_D11KBACXX.IND3	183557983	35871118970	Mate-pairs 3Kb	Illumina	ERX546778	24x
AWU	А	AWU_AOSF_1_C0BULACXX.IND1	33955645	6731457878	overlapping PE	Illumina	ERX546816	86x
AWU	А	AWU_AOSF_1_D0J4FACXX.IND1	184348978	36406701485	overlapping PE	Illumina	ERX546795	
AWU	А	AWU_AOSF_2_D0J4FACXX.IND1	173912862	34132669204	overlapping PE	Illumina	ERX546803	
AWU	А	AWU_AOSF_4_D0J4KACXX.IND1	118761014	23565095820	overlapping PE	Illumina	ERX546766	
AWU	А	AWU_AOSF_6_C0D1LACXX.IND1	142592731	28210290830	overlapping PE	Illumina	ERX546794	
AWU	А	AWU_AOSN_2_C2MP1ACXX.IND18	112059267	18207771474	Mate-pairs Nextera 3 Kb	Illumina	ERX546793	26x
AWU	А	AWU_AOSN_4_D2BM7ACXX.IND18	130196132	21273821373	Mate-pairs Nextera 3 Kb	Illumina	ERX546821	
AWU	А	AWU_AOSN_1_C2MP1ACXX.IND2	128515390	20882553184	Mate-pairs Nextera 5 Kb	Illumina	ERX546851	35x
AWU	А	AWU_AOSN_4_D2C4BACXX.IND2	128931290	21392347587	Mate-pairs Nextera 5 Kb	Illumina	ERX546756	
AWU	А	AWU_AOSN_7_D25ULACXX.IND2	63924125	10508889688	Mate-pairs Nextera 5 Kb	Illumina	ERX546847	
AWU	А	AWU_AOSN_4_D2C5KACXX.IND4	132901113	21958813992	Mate-pairs Nextera 8 Kb	Illumina	ERX546787	22x
AWU	А	AWU_AOSN_7_D25ULACXX.IND4	65407657	10724020167	Mate-pairs Nextera 8 Kb	Illumina	ERX546842	
AWU	А	AWU_AORS_HLG9GIU01	376951	139474495	Single Reads	454	ERX546760	15x
AWU	А	AWU_AORS_HO6MKXJ01	615920	275537385	Single Reads	454	ERX546817	
AWU	А	AWU AORS HO6MKXJ02	600921	244313411	Single Reads	454	ERX546828	

AWU	A	AWU_AORS_HO8LWZP01	566389	250562733	Single Reads	454	ERX546783
AWU	А	AWU_AORS_HO8LWZP02	576975	250783193	Single Reads	454	ERX546855
AWU	А	AWU_AORS_HOGWG9K01	471512	216412101	Single Reads	454	ERX546798
AWU	А	AWU_AORS_HOGWG9K02	623520	290152549	Single Reads	454	ERX546765
AWU	А	AWU_AORS_HOKMQ7201	415422	194219290	Single Reads	454	ERX546804
AWU	А	AWU_AORS_HOKMQ7202	415767	194112470	Single Reads	454	ERX546810
AWU	А	AWU_AORS_HOTXFWN01	573904	249396146	Single Reads	454	ERX546789
AWU	А	AWU_AORS_HOTXFWN02	616636	251440354	Single Reads	454	ERX546797
AWU	А	AWU_AORS_HOXJLF101	619309	271457362	Single Reads	454	ERX546826
AWU	А	AWU_AORS_HOXJLF102	589373	262356681	Single Reads	454	ERX546833
AWU	А	AWU_AORS_HP9GW5D01	616582	329032439	Single Reads	454	ERX546796
AWU	А	AWU_AORS_HPAKMIN01	604431	247521317	Single Reads	454	ERX546808
AWU	А	AWU_AORS_HPAKMIN02	599433	252223557	Single Reads	454	ERX546799
AWU	А	AWU_AORS_HPDXEDZ01	576140	267133896	Single Reads	454	ERX546781
AWU	А	AWU_AORS_HPDXEDZ02	604043	268649166	Single Reads	454	ERX546786
AWU	А	AWU_AORS_HPJN7LD01	627802	303967961	Single Reads	454	ERX546856
AWU	А	AWU_AORS_HPJN7LD02	632803	301178645	Single Reads	454	ERX546780
AWU	А	AWU_AORS_HPLH4SP01	574419	279919660	Single Reads	454	ERX546839
AWU	А	AWU_AORS_HPLH4SP02	541488	269140415	Single Reads	454	ERX546763
AWU	А	AWU_AORS_HPNH9JK01	566545	277699403	Single Reads	454	ERX546825
AWU	А	AWU_AORS_HPNH9JK02	542232	250634628	Single Reads	454	ERX546835
AWU	А	AWU_AORS_HPPEQTK01	509099	253840880	Single Reads	454	ERX546755
AWU	А	AWU_AORS_HPPEQTK02	397398	183638911	Single Reads	454	ERX546776
AWU	А	AWU_AORS_HPQ6PEM01	324214	140937382	Single Reads	454	ERX546759

AWU	А	AWU_AORS_HPQ6PEM02	604340	258006924	Single Reads	454	ERX546792
AWU	А	AWU_AORS_HPWTB8401	632065	303346757	Single Reads	454	ERX546782
AWU	А	AWU_AORS_HPWTB8402	628856	283953031	Single Reads	454	ERX546850
AWU	А	AWU_AORS_HPYQNEV01	541025	254463651	Single Reads	454	ERX546785
AWU	А	AWU_AORS_HPYQNEV02	595941	281903473	Single Reads	454	ERX546775
AWU	А	AWU_AORS_HQ3AIIJ01	645852	273877444	Single Reads	454	ERX546812
AWU	А	AWU_AORS_HQ3AIIJ02	635565	261155309	Single Reads	454	ERX546853
AWU	А	AWU_AORS_HQ6XQKF01	602974	279814892	Single Reads	454	ERX546854
AWU	А	AWU_AORS_HQ6XQKF02	450682	192755614	Single Reads	454	ERX546857
AWU	А	AWU_AORS_HQBAEKW02	638393	315269575	Single Reads	454	ERX546779
AWU	А	AWU_AORS_HQC2HZF03	233986	110390642	Single Reads	454	ERX546820
AWU	А	AWU_AORS_HQC2HZF04	252397	132477279	Single Reads	454	ERX546757
AWU	А	AWU_AORS_HQC7JZG01	601648	292351551	Single Reads	454	ERX546829
AWU	А	AWU_AORS_HQC7JZG02	598632	280339696	Single Reads	454	ERX546824
AWU	А	AWU_AORS_HQE6HNV01	575033	294218540	Single Reads	454	ERX546837
AWU	А	AWU_AORS_HQE6HNV02	562072	276129568	Single Reads	454	ERX546771
AWU	А	AWU_AORS_HQG0JXZ01	605366	302641389	Single Reads	454	ERX546843
AWU	А	AWU_AORS_HQG0JXZ02	608598	289064135	Single Reads	454	ERX546774
AWU	А	AWU_AORS_HQR99NJ02	267774	134696910	Single Reads	454	ERX546806
AWU	А	AWU_AORS_HQR99NJ03	268450	136910337	Single Reads	454	ERX546840
AWU	А	AWU_AORS_HQR99NJ04	259009	128977810	Single Reads	454	ERX546813
AWU	А	AWU_AORS_HQVT54K01	670862	357817336	Single Reads	454	ERX546767
AWU	А	AWU_AORS_HQVT54K02	629420	327604262	Single Reads	454	ERX546852
AWU	А	AWU_AORS_HQXPWBU01	575053	311123919	Single Reads	454	ERX546834

AWU	А	AWU_AORS_HQXPWBU02	592747	314220456	Single Reads	454	ERX546814
AWU	А	AWU_AORS_HR0B5WL01	476167	129297327	Single Reads	454	ERX546762
AWU	А	AWU_AORS_HR0B5WL02	522655	143434538	Single Reads	454	ERX546832
AWU	А	AWU_AORS_HR7Y0Z201	494273	151280248	Single Reads	454	ERX546827
AWU	А	AWU_AORS_HR7Y0Z202	451613	137776321	Single Reads	454	ERX546773
AWU	А	AWU_AORS_HRJY4EM01	583444	236195624	Single Reads	454	ERX546772
AWU	А	AWU_AORS_HRJY4EM02	507965	197165750	Single Reads	454	ERX546805
AWU	А	AWU_AORS_HRLT5NJ01	439153	146160122	Single Reads	454	ERX546777
AWU	А	AWU_AORS_HRLT5NJ02	400105	146048898	Single Reads	454	ERX546822
AWU	А	AWU_AORS_HRS9HCN01	472104	238860086	Single Reads	454	ERX546802
AWU	А	AWU_AORS_HRS9HCN02	540831	273328131	Single Reads	454	ERX546800
AWU	А	AWU_AORS_HRWNJ0F01	332429	137572971	Single Reads	454	ERX546819
AWU	А	AWU_AORS_HRWNJ0F02	483017	197840503	Single Reads	454	ERX546836
AWU	А	AWU_AORS_HRYGP8A01	593882	234911883	Single Reads	454	ERX546764
AWU	А	AWU_AORS_HRYGP8A02	568850	225739689	Single Reads	454	ERX546809
AWU	А	AWU_AORS_HS3NGS202	593496	271995164	Single Reads	454	ERX546848
AWU	А	AWU_AORS_HSBPWMC07	91079	36958456	Single Reads	454	ERX546791
AWU	А	AWU_AORS_HSBPWMC08	85328	34801846	Single Reads	454	ERX546801
AWU	А	AWU_AORS_HSDIBFO01	581079	282921477	Single Reads	454	ERX546838
AWU	А	AWU_AORS_HSDIBFO02	592838	266142240	Single Reads	454	ERX546784
AWU	А	AWU_AORS_HSFJ8MK01	607026	254366558	Single Reads	454	ERR588819
AWU	А	AWU_AORS_HSFJ8MK02	598573	217174740	Single Reads	454	ERX546849
AWU	А	AWU_AORS_HT2R6K001	616436	354909746	Single Reads	454	ERX546818
AWU	А	AWU_AORS_HTAXONP01	492955	227121584	Single Reads	454	ERX546770

AWU	А	AWU_AORS_HTAXONP02	605770	266418303	Single Reads	454	ERX546844	
AWU	А	AWU_AORS_HTCPZFN01	586104	289733463	Single Reads	454	ERX546788	
AWU	А	AWU_AORS_HTCPZFN02	620227	293411238	Single Reads	454	ERX546807	
AWU	А	AWU_AORS_HTEFU8101	598533	239492636	Single Reads	454	ERX546769	
AWU	А	AWU_AORS_HTEFU8102	622271	230573314	Single Reads	454	ERX546841	
AWU	А	AWU_AORS_HTNM2OH02	602554	281689812	Single Reads	454	ERX546830	
AWU	А	AWU_AORS_HTRLXNS01	618531	306184837	Single Reads	454	ERX546811	
AWU	А	AWU_AORS_HTRLXNS02	614926	297895268	Single Reads	454	ERX546754	
AWU	А	AWU_AORS_HTRQELM01	582923	257934536	Single Reads	454	ERX546846	
AWU	А	AWU_AORS_HTRQELM02	619410	261141889	Single Reads	454	ERX546768	
AWU	А	AWU_AORS_HTTFXII01	657236	291923817	Single Reads	454	ERX546845	
AWU	А	AWU_AORS_HTTFXII02	642957	282417197	Single Reads	454	ERX546761	
AWU	А	AWU_AORS_HTTJ7TZ01	592087	288519958	Single Reads	454	ERX546831	
AWU	А	AWU_AORS_HTTJ7TZ02	594370	280348006	Single Reads	454	ERX546758	
AWU	А	AWU_AORS_HTVBL6N01	660475	383854941	Single Reads	454	ERX546823	
AWU	А	AWU_AORS_HTVBL6N02	634782	363284614	Single Reads	454	ERX546815	
AWU	А	AWU_AORS_HTVFTYI02	664033	336384976	Single Reads	454	ERX546790	
BBX	А	BBX_AOSW_1_D1D53ACXX.IND5	178225679	34704998557	Paired-end	Illumina	ERX697294	471x
BBX	А	BBX_AOSW_1_H32GMBCXX.IND5	129340482 6	61406442546	Paired-end	Illumina	ERX1886616	
BBX	А	BBX_AOSW_1_H57N7BCXX.IND5	114968046	54158595214	Paired-end	Illumina	ERX1886621	
BBX	А	BBX_AOSW_2_H32GMBCXX.IND5	132162999	63021298371	Paired-end	Illumina	ERX1886622	
BBX	В	BBX_BOSW_2_C1CRDACXX.IND6	185226443	36388653397	Paired-end	Illumina	ERX697299	
BBX	С	BBX_COSW_1_H072TAMXX.IND7	88896954	42521091728	Paired-end	Illumina	ERX697298	
BBX	С	BBX_COSW_2_D1D53ACXX.IND7	187527862	36107158449	Paired-end	Illumina	ERX697297	

					-			
BBX	С	BBX_COSW_2_H072TAMXX.IND7	89768432	42909747910	Paired-end	Illumina	ERX697296	
BBX	С	BBX_COSW_2_H57N7BCXX.IND7	141600651	66340720909	Paired-end	Illumina	ERX1886620	
BBX	D	BBX_DOSW_3_C1CRDACXX.IND8	195620139	38283308256	Paired-end	Illumina	ERX697295	
BBX	Е	BBX_EOSW_1_H55MLBCXX.IND9	127908274	59324914635	Paired-end	Illumina	ERX1886617	
BBX	Е	BBX_EOSW_2_H32GLBCXX.IND9	134000800	55186245541	Paired-end	Illumina	ERX1886619	
BBX	Е	BBX_EOSW_2_H55MLBCXX.IND9	104846648	48165716278	Paired-end	Illumina	ERX1886618	
BBX	Е	BBX_EOSW_3_D1D53ACXX.IND9	185076013	35974428528	Paired-end	Illumina	ERX697292	
BBX	F	BBX_FOSW_4_D1D53ACXX.IND10	173673937	33077225178	Paired-end	Illumina	ERX697293	
AWU	A2	LR6000024-DNA_B02-LRAAD-01	96268	477206869	TruSeq Synthetic Reads	Illumina	ERX1936767	бх
AWU	A2	LR6000024-DNA_B02-LRAAD-02	97511	485762221	TruSeq Synthetic Reads	Illumina	ERX1936768	
AWU	A2	LR6000024-DNA_B02-LRAAD-03	99930	497943141	TruSeq Synthetic Reads	Illumina	ERX1936769	
AWU	A2	LR6000024-DNA_B02-LRAAD-04	96777	481729766	TruSeq Synthetic Reads	Illumina	ERX1936770	
AWU	A2	LR6000024-DNA_B02-LRAAD-05	99331	488240514	TruSeq Synthetic Reads	Illumina	ERX1936771	
AWU	A2	LR6000024-DNA_B02-LRAAD-06	133440	621932686	TruSeq Synthetic Reads	Illumina	ERX1936772	
AWU	A2	LR6000024-DNA_B02-LRAAD-07	212972	890601310	TruSeq Synthetic Reads	Illumina	ERX1936773	
AWU	G1	AWU_msDDZ	166474	741066645	TruSeq Synthetic Reads	Illumina	ERX1936761	
AWU	G1	AWU_msDEA	171897	734825556	TruSeq Synthetic Reads	Illumina	ERX1936762	
AWU	G1	AWU_msDED	121611	589161920	TruSeq Synthetic Reads	Illumina	ERX1936765	
AWU	G1	AWU_msDEF	124691	592617439	TruSeq Synthetic Reads	Illumina	ERX1936766	
AWU	G1	AWU_msDEB	175574	750471312	TruSeq Synthetic Reads	Illumina	ERX1936763	
AWU	G1	AWU_msDEC	167087	731676445	TruSeq Synthetic Reads	Illumina	ERX1936764	
AWU	G1	AWU-msDBX	110436	530240247	TruSeq Synthetic Reads	Illumina	ERX1936760	

RNAseq											
AXF	AA	AXF_AAOSW_8_C0D1LACXX.IND2	59050722 11928245844	Paired-end # RNA	Illumina	ERX332625					
AXF	BA	AXF_BAOSW_8_C0D1LACXX.IND4	63191029 12764587858	Paired-end # RNA	Illumina	ERX332624					
AXF	CA	AXF_CAOSW_8_C0D1LACXX.IND5	68158203 13767957006	Paired-end # RNA	Illumina	ERX332621					
AXF	DA	AXF_DAOSW_7_C0D1LACXX.IND6	72263408 14597208416	Paired-end # RNA	Illumina	ERX332626					
AXF	EA	AXF_EAOSW_7_C0D1LACXX.IND7	57005112 11515032624	Paired-end # RNA	Illumina	ERX332623					
AXF	FA	AXF_FAOSW_7_C0D1LACXX.IND12	65878896 13307536992	Paired-end # RNA	Illumina	ERX332622					
BHC	AF	BHC_AFOSW_8_C4VAEACXX.IND12	29858750 5931520880	Paired-end # RNA	Illumina	ERX1916513					
BHC	AG	BHC_AGOSW_7_C4VBLACXX.IND13	27938013 5587303314	Paired-end # RNA	Illumina	ERX1916512					
BHC	BA	BHC_BAOSW_3_C4VR1ACXX.IND15	32668746 6486518369	Paired-end # RNA	Illumina	ERX1796981					
BHC	BB	BHC_BBOSW_3_C4VR1ACXX.IND16	29920489 5929334824	Paired-end # RNA	Illumina	ERX1796984					
BHC	BC	BHC_BCOSW_3_C4VR1ACXX.IND18	33368451 6625154612	Paired-end # RNA	Illumina	ERX1796982					
BHC	BD	BHC_BDOSW_3_C4VR1ACXX.IND19	33982054 6753975751	Paired-end # RNA	Illumina	ERX1796983					
BHD	AA	BHD_AAOSW_1_C3YEPACXX.IND1	29134976 5808377002	Paired-end # RNA	Illumina	ERX1796974					
BHD	AB	BHD_ABOSW_1_C3YEPACXX.IND3	32355510 6448224478	Paired-end # RNA	Illumina	ERX1796976					
BHD	AC	BHD_ACOSW_1_C3YEPACXX.IND8	43958162 8765695282	Paired-end # RNA	Illumina	ERX1796975					
BHD	AK	BHD_AKOSW_2_C3YEPACXX.IND23	31082716 6204885589	Paired-end # RNA	Illumina	ERX1916511					
BHD	AL	BHD_ALOSW_2_C3YEPACXX.IND25	27615039 5520361779	Paired-end # RNA	Illumina	ERX1916509					
BHD	AM	BHD_AMOSW_2_C3YEPACXX.IND27	30157050 6022460185	Paired-end # RNA	Illumina	ERX1916510					
BIG	G	BIG_GOSW_5_C49VTACXX.IND7	27305012 5442125902	Paired-end # RNA	Illumina	ERX1916514					

Supplementary Table 2 Metrics of final haploid (haploid V2) and diploid (diploid V2) versions of the pedunculate oak genome sequence assembly.

	Diploid V2 (Assembly A5 ^a)	Haploid V2 (Assembly H1 ^b)
Assembly	Diploid	Haploid
No. of sequences	8,827	1,409
Cumulative size	1,455,104,916	814,282,569
N50	821,707	1,342,530
N90	198,501	333,129
L50	537	192
L90	1,880	649
% of N's	4.6	2.94
Completeness using BUSCO	210 (90.4%)	202 (90.8%)
^a from Supplementary Table 10		
^b from Supplementary Table 11		

Supplementary Table 3 Comparison of genome assemblies from available heterozygous trees. Best (green) and worst (red) assembly metrics, excluding *Poplulus trichocarpa*.

Species	Assembly availability	# contigs	Cumulative size of contigs (Mb)	Contigs N50 size	# scaffolds	Cumulative size of scaffolds (Mb)	% of N	Scaffold N50 size	Busco %C	Busco %D
Olea europaea	http://denovo.cnag.cat/genomes/olive/ download/Oe6/Oe6.scaffolds.fa.gz	38,053	1,265	87,946	11,038	1,319	4.09	443,100	277 (91.4%)	126 (41.6%)
Quercus robur	This study	22,615	790	69,349	1,409	814	2.94	1,342,530	269 (88.8%)	49 (16.2%)
Betula pendula	https://genomevolution.org/coge/api/v 1/genomes/35079/sequence	27,580	425	49,342	5,642	435	2.34	239,520	261 (86.1%)	38 (12.5%)
Fraxinus excelsior	http://www.ashgenome.org/assemblies	119,515	718	24,932	89,514	867	17.19	103,995	272 (89.8%)	97 (32.0%)
Castanea mollissim a	https://hardwoodgenomics.org/chinese -chestnut-genome#genomedownloads	70,867	710	22,063	41,260	724	1.86	39,561	264 (87.1%)	50 (16.5%)
Quercus lobata	https://valleyoak.ucla.edu/genomicres ources/	255,152	1,069	17,576	94,394	1,183	9.64	161,656	271 (89.4%)	98 (32.3%)
Populus trichocar pa	https://genomevolution.org/coge/api/v 1/genomes/25127/sequence	8,313	423	552,806	1,446	434	2.57	19,465,461	279 (92.0%)	108 (35.6%)

Supplementary Table 4 Annotation of transposable elements.

						1788
		# TE consensus	# genome copies	Genome coverage (kb)	Genome coverage %	TE content coverage %
Class I	Copia	211	89,447	87,215	11.04	20.71
Retro-elements	Gypsy	276	91,652	107,561	13.61	25.54
LTR	LARD/TRIM/Other	80	38,726	29,058	3.68	6.90
Class I	LINE	408	157,114	66,135	8.37	15.70
Retro-elements non-LTR	SINE	30	4,571	1,216	0.15	0.29
Class I	Other	16	20,970	4,224	0.53	1.00
	TIR	313	141,489	52,207	6.61	12,.39
Class II	MITE	67	28,124	7,760	0.98	1.84
DNA	Helitron	8	3,642	2,006	0.25	0.48
ti ansposons	Other	11	2,987	2,012	0.25	0.48
Unknown		317	134,652	54,428	7.27	13.63
Endovirus		13	2,818	4,385	0.55	1.04
Total		1,750	716,192	420,651	53.30 ^a	100

^a The total 53.3% of genome coverage reported here corresponds to the cumulative sum of coverage for the different orders/familes. Total TE genome coverage is 52%, without redundancy between copies.

Supplementary Table 5 List of control SNPs (C) and somatic mutations (SMs) detected in the "3P" pedunculate oak accession. C: control 1815

	Locus ID		Mutation	Origin of the	f(alt)pool	f(alt)pool
Mutation category		Chromosomal location		mutation		>0.5%
С	Sc0000093_652917	Chr1-41939193	T/A	within species	0.9697	
С	Sc0000158_1024005	Chr2-27130279	T/A	within species	0.9637	
С	Sc0000067_389965	Chr3-27491298	A/T	within species	0.9534	
С	Sc0000033_2516576	Chr4-10285224	A/C	within species	0.9585	
С	Sc0000505_233875	Chr5-39212547	T/G	within species	0.9507	
С	Sc0000170_1375115	Chr6-36333647	C/A	within species	0.9542	
С	Sc0000268_125122	Chr7-23637407	T/C	within species	0.9605	
С	Sc0000187_1162488	Chr8-53746285	T/C	within species	0.9700	
С	Sc0000168_672869	Chr9-31527061	C/T	within species	0.9541	
С	Sc0000447_97317	Chr10-22125827	A/G	within species	0.9526	
С	Sc0000099_1051673	Chr11-7530106	A/G	within species	0.9679	
С	Sc0000425_378736	Chr12-27206974	G/A	within species	0.9691	
SM	Sc0000080_1329750	Chr8-58757192	G/A	3P – between XL1 and XL2	0.0330	У
SM	Sc0000573_185294	Chr7-21324198	A/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000010_1057132	Chr3-13766752	G/A	3P – between XL1 and XL2	0.1154	У
SM	Sc000003_4011526	Chr2-84974261	A/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000010_758473	Chr3-13468093	G/A	3P – between XL1 and XL2	0.0000	
SM	Sc0000015_2644541	Chr3-50836723	G/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000057_1996281	Chr11-19272464	C/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000235_409999	Chr12-5868120	C/T	3P – between XL1 and XL2	0.1384	У
SM	Sc0000122_532208	Chr1-8511503	C/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000139_351870	Chr1-28576871	T/C	3P – between XL1 and XL2	0.0000	
SM	Sc0000233_840676	Chr7-12397418	G/A	3P – between XL1 and XL2	0.0000	
SM	Sc0000588_301268	Chr4-27781193	G/A	3P – between XL1 and XL2	0.0000	
SM	Sc0000065_545730	Chr5-66436424	T/C	3P – L1 branch	0.0000	
SM	Sc0000181_1118667	Chr2-50905345	C/T	3P – L1 branch	0.0782	У
SM	Sc0000200_640712	Chr5-54088853	G/A	3P – L1 branch	0.0000	•
SM	Sc0000667_35498	Chr1-10565982	G/A	3P – L1 branch	0.0133	У
SM	Sc0000444_256472	Chr3-25184616	C/T	3P – L1 branch	0.0049	

SNP, SM: somatic mutation, Mutation: reference /alternative (alt) allele, f(alt)pool: frequency of the alternative allele in the pool-seq data set. 1816

SM	Sc0000277_447345	Chr2-113550344	T/C	3P – L1 branch	0.0000	
SM	Sc0000135_631742	Chr7-25169846	T/C	3P – L1 branch	0.0000	
SM	Sc000001_4448299	Chr6-48948241	T/G	3P – L1 branch	0.0000	
SM	Sc0000219_286289	Chr2-49724208	A/T	3P – L1 branch	0.0000	
SM	Sc0000395_657452	Chr8-17311448	G/T	3P – L1 branch	0.0000	
SM	Sc0000099_1809337	Chr11-6772442	C/T	3P – L2 branch	0.0078	у
SM	Sc0000035_1061781	Chr1-32821443	C/T	3P – L2 branch	0.0043	
SM	Sc0000066_1207928	Chr2-22466474	G/A	3P – L2 branch	0.0000	
SM	Sc0000103_228814	unanchored scaffold	C/T	3P – L2 branch	0.0000	
SM	Sc0000578_47594	Chr2-70260112	C/T	3P – L2 branch	0.0000	
SM	Sc0000031_1042378	Chr9_36922447	C/T	3P – between XL1 and XL2	0.0035	
SM	Sc0000114_1570819	Chr4_39071460	A/G	3P – between XL1 and XL2	0.0000	
SM	Sc0000114_960640	Chr4_39681639	G/A	3P – between XL1 and XL2	0.0030	
SM	Sc0000146_1249018	Chr12_26296592	C/T	3P – between XL1 and XL2	0.0000	
SM	Sc0000975_67191	Chr2_23741692	T/C	3P – between XL1 and XL2	0.0000	
SM	Sc000000_3201322	Chr1_23876533	G/A	3P – L1 branch	0.0000	
SM	Sc0000041_558993	Chr1_2895035	G/A	3P - L1 branch	0.0935	У
SM	Sc0000228_252981	Chr5_34619071	C/T	3P - L1 branch	0.0000	
SM	Sc0000277_395519	Chr2_113498518	G/A	3P - L1 branch	0.0608	У
SM	Sc0000570_213658	Chr8_40730574	C/T	3P - L1 branch	0.0244	у
SM	Sc0001123_18450	unanchored scaffold	G/A	3P - L1 branch	0.0270	У
SM	Sc000002_4278465	Chr2_66564257	T/A	3P - L2 branch	0.0000	
SM	Sc000005_165035	Chr11_33010956	T/G	3P - L2 branch	0.0000	
SM	Sc0000242_170918	Chr2_55024334	C/T	3P - L2 branch	0.0000	
SM	Sc0000312_31989	Chr6_28492838	A/T	3P - L2 branch	0.0000	
SM	Sc0000584_280071	unanchored scaffold	A/T	3P - L2 branch	0.3152	у
SM	Sc0000026_779464	unanchored scaffold	G/A	3P – between XL2 and L3	0.0359	У
SM	Sc0000027_691249	Chr2_98491575	A/G	3P – between XL2 and L3	0.0000	
SM	Sc0000042_876919	Chr4_19144163	T/A	3P – between XL2 and L3	0.0000	
SM	Sc0000051_786541	Chr2_30320278	C/T	3P – between XL2 and L3	0.0000	
SM	Sc0000056_626880	Chr5_18059797	C/T	3P – between XL2 and L3	0.0690	у
SM	Sc0000085_693443	Chr10_14911573	C/T	3P – between XL2 and L3	0.0140	у
SM	Sc0000097_1855202	Chr9_47939472	C/T	3P – between XL2 and L3	0.0000	
SM	Sc0000108_1664655	Chr10_27649124	C/T	3P – between XL2 and L3	0.0000	

SM	Sc0000132_1066473	unanchored scaffold	T/A	3P – between XL2 and L3	0.0000	
SM	Sc0000167_777615	Chr2_81379058	T/C	3P – between XL2 and L3	0.0029	
SM	Sc0000170_850309	Chr6_35808841	G/T	3P – between XL2 and L3	0.0000	
SM	Sc0000210_857733	Chr12_35930326	C/T	3P – between XL2 and L3	0.0000	
SM	Sc0000227_150153	Chr10_8928796	G/A	3P – between XL2 and L3	0.0000	
SM	Sc0000266_244617	Chr11_43001996	G/A	3P – between XL2 and L3	0.0000	
SM	Sc0000266_72243	Chr11_43174370	A/C	3P – between XL2 and L3	0.0000	
SM	Sc0000274_597070	Chr2_35105614	G/A	3P – between XL2 and L3	0.0305	у
SM	Sc0000300_540958	Chr5_55161003	C/T	3P – between XL2 and L3	0.0000	
SM	Sc0000620_260076	Chr11_26408896	C/T	3P – between XL2 and L3	0.0028	

Supplementary Table 6 List of control SNPs (C) and somatic mutations (SMs) in the offspring of accession "3P". Acorns of the reference genotype "3P" were collected from the L1 and L2 branches indicated in Fig. 2b. C: control SNP, SM somatic mutation, N: sample size with accurate genotypic information, H0: observed heterozygosity. f(pool): frequency of the alternative allele. A value of 0 in this last column indicates a mutation detected only in the reference "3P" genotype and transmitted to its offspring.

SNP	Locus ID	Origin of the	Success of	% missing	Ν	HO	
Category		mutation	the assay	data			f(pool)
С	Chr1-41939193	within species	Y	0.440	65	0.400	0.9697
С	Chr2-27130279	within species	Y	0.853	17	0.588	0.9637
С	Chr3-27491298	within species	Y	0.276	84	0.595	0.9534
С	Chr4-10285224	within species	Y	0.819	21	0.619	0.9545
С	Chr5-39212547	within species	Y	0.466	62	0.306	0.9507
С	Chr6-36333647	within species	Y	0.905	11	0.455	0.9542
С	Chr7-23637407	within species	Y	0.138	100	0.820	0.9605
С	Chr8-53746285	within species	Y	0.259	86	0.814	0.9700
С	Chr9-31527061	within species	Y	0.198	93	0.215	0.9541
С	Chr10-22125827	within species	Y	0.888	13	0.231	0.9526
С	Chr11-7530106	within species	Y	0.914	10	0.600	0.9679
С	Chr12-27206974	within species	Y	0.172	96	0.875	0.9691
SM	Sc0000573_185294	3P – between XL1 and XL2	Y	0.172	96	0.000	0.000
SM	Sc000003 4011526	3P – between XL1 and XL2	Y	0.284	83	0.084	0.000
SM	Sc000010 758473	3P - between XL1 and XL2	Ŷ	0.233	89	0.124	0.000
SM	Sc0000015 2644541	3P - between XL1 and XL2	Ŷ	0.595	47	0.191	0.000
SM	Sc0000057 1996281	3P – between XL1 and XL2	Y	0.491	59	0.288	0.000
SM	Sc0000122 532208	3P – between XL1 and XL2	Y	0.871	15	0.000	0.000
SM	Sc0000139_351870	3P – between XL1 and XL2	Y	0.198	93	0.000	0.000
SM	Sc0000233_840676	3P – between XL1 and XL2	Y	0.026	113	0.000	0.000
SM		3P – between XL1 and XL2	Y	0.267	85	0.000	0.000
SM	Sc0000065_545730	3P – L1 branch	Y	0.836	19	0.000	0.000

SM	Sc0000200_640712	3P – L1 branch	Y	0.069	108	0.111	0.000
SM	Sc0000444_256472	3P – L1 branch	Y	0.034	112	0.018	0.005
SM	Sc0000277_447345	3P – L1 branch	Y	0.690	36	0.111	0.000
SM	Sc0000135_631742	3P – L1 branch	Y	0.595	47	0.000	0.000
SM	Sc000001_4448299	3P – L1 branch	Y	0.629	43	0.000	0.000
SM	Sc0000219_286289	3P – L1 branch	Ν	NA	0	NA	0.000
SM	Sc0000395_657452	3P – L1 branch	Ν	NA	0	NA	0.000
SM	Sc0000035_1061781	3P – L2 branch	Y	0.276	84	0.000	0.004
SM	Sc0000066_1207928	3P – L2 branch	Y	0.190	94	0.000	0.000
SM	Sc0000103_228814	3P – L2 branch	Y	0.897	12	0.000	0.000
SM	Sc0000578_47594	3P – L2 branch	Y	0.026	113	0.000	0.000

Species acronym*	#genes	#orthogroups	#genes in orthogroups	#singletons	% Genes in orthogroups	#genes shared with at least one other species	%genes shared with at least one other species	#species specific orthogroups	#genes in species specific orthogroups
Al	32,657	17,186	27,260	5,397	0.83	24,449	0.75	813	2,811
At	27,416	16,716	24,733	2,683	0.90	24,334	0.89	141	399
Wa	23,440	12,775	20,192	3,248	0.86	17,402	0.74	364	2,790
Fv	32,831	14,304	25,093	7,738	0.76	19,953	0.61	1,275	5,140
Gm	56,044	15,235	46,400	9,644	0.83	41,978	0.75	1,552	4,422
Rc	31,220	14,658	21,088	10,132	0.68	18,913	0.61	754	2,175
St	35,119	13,041	28,897	6,222	0.82	21,825	0.62	1,060	7,072
Ср	27,584	13,483	20,285	7,299	0.74	18,334	0.66	520	1,951
Сс	24,533	13,916	21,425	3,108	0.87	20,497	0.84	316	928
Eg	36,376	13,615	29,063	7,313	0.80	26,402	0.73	722	2,661
Md	63,514	17,217	46,524	16,990	0.73	37,225	0.59	3,324	9,299
Pt	41,335	14,921	33,604	7,731	0.81	31,412	0.76	728	2,192
Рр	27,864	14,545	24,651	3,213	0.88	23,230	0.83	311	1,421
Qr	25,808	11,813	22,498	3,310	0.87	20,761	0.80	479	1,737
Тс	29,452	14,591	23,608	5,844	0.8	21,722	0.74	465	1,886
$V \nu$	26,346	12,951	19,774	6,572	0.75	18,135	0.69	589	1,639
Total #genes	541,539		435,095	106,444		386,572			48,523

1824 Supplementary Table 7 Result of the OrthoMCL analysis and comparison between the 16 eudicot species used in this study.

1825 1826

* Al Arabidopsis lyrata, At Arabidopsis thaliana, Wa Citrullus lanatus, Fv Fragaria vesca, Gm Glycine max, Rc Ricinus communis, St Solanum tuberosum, Cp Carica papaya, Cc Citrus climentina, Eg Eucalyptus grandis, Md Malus domestica, Pt Populus trichocarpa, Pp Prunus persica, Qr Quercus robur, Tc Theobroma cacao, Vv Vitis vinifera.

Supplementary Table 8 Repertoire of NB-LRR-related disease resistance genes in oak.
Genes are classified in different categories according to the presence of the canonical NBARC (NB), leucine-rich repeat (LRR) domains and/or the N-terminal domains typically
associated with disease resistance NB-LRR genes in plant genomes, namely Toll interleukin
receptor-like (TIR), coiled-coil (CC) and resistance to powdery mildew protein RPW8 (R)
domains. X indicates the presence of a putative integrated domain (ID).

			Integrated domains
Category	Acronym	Total	(X)
CC-NB-LRR (X)	CNL	258	16
CC-NB (X)	CN	47	3
CC-LRR	CL	3	
CC (X)	С	14	1
NB-LRR-CC-NB-LRR	NLCNL	1	
CC(3x)-NB-LRR	C(3x)NL	1	
RPW8-NB-LRR (X)	RNL	15	3
RPW8	R	1	
TIR-NB-LRR (X)	TNL	186	11
TIR-NB (X)	TN	25	3
TIR-LRR (X)	TL	3	1
TIR	Т	151	
NB-LRR (X)	NL	240	11
NB (X)	Ν	61	4
LRR (X)	L	85	1
Total		1,091	54

Supplementary Table 9 Genetic diversity (π) at 0-fold, 4-fold degeneracy and π_0/π_4 ratio. **Estimates were averaged over 1,000** randomly picked genes in each category and repeated 1838 100 times. Mean and 95% confidence intervals (in parentheses) are reported. Values should 1839 be multiplied by 10⁻³.

	Total	Expanded	Contracted	Unchanged
"3P" Ger	nome sequence			
π_0	5.0 (4.6, 5.5)	7.0 (6.6, 7.4)	3.2 (3.0, 3.4)	3.2 (3.0, 3.5)
π_4	11.4 (10.6, 12)	12.5 (11.7, 13.3)	10.8 (10.2, 11.4)	10 (9.2, 10.9)
π_0/π_4	0.44 (0.39, 0.49)	0.56 (0.52, 0.61)	0.3 (0.27, 0.32)	0.32 (0.29, 0.37)
Pool-seq	uencing			
π_0	5.4 (5.0, 5.7)	7.6 (7.2, 7.9)	2.9 (2.7, 3.0)	3 (2.8, 3.2)
π_4	10.8 (10.3, 11.3)	12.5 (12.0, 13.0)	9.5 (9.2, 9,9)	9.3 (8.9, 9.8)
π_0/π_4	0.5 (0.46, 0.53)	0.6 (0.57, 0.65)	0.3 (0.28, 0.32)	0.32 (0.3, 0.34)
		(0.20, 0.22)	0.02 (0.0, 0.01)

Supplementary Table 10 Metrics of the pedunculate oak assembly at each step of the Newbler process.

	Newbler A1	Newbler A2	Newbler A3	Newbler A4	Newbler A5
Assembly step	Raw output	Graph simplification	Scaffolding	Gap closing	Contamination removal
# sequences	296,255	198,695	9,025	9,025	8,827
Cumulative size	1,313,577,586	1,330,866,990	1,455,541,024	1,458,028,538	1,455,104,916
N50	9,499	16,207	818,147	821,283	821,707
N90	1,800	3,322	538	194,343	198,501
L50	38,579	23,591	193,405	7538	537
L90	158,717	89,893	1,892	1,893	1,880
% of N's	0	1.3	11.19	4.63	4.6

1849 Supplementary Table 11 Metrics of pedunculate oak assembly at each step of the Celera 1850 process.

	Celera C1	Celera C2
Assembly step	Raw output	Scaffolding
No. of sequences	296,255	14,088
Cumulative size	1,313,577,586	1,273,117,594
N50	9,499	266,385
N90	1,800	55,257
L50	38,579	1,418
L90	158,717	5,257
% of N's	0	9.24

Supplementary Table 12 Structural manual curation of mRNAs indicating the type of protein coding structure (CDS) curation.

1,714 genes	
1,347 (79%)	
367 (21%)	
233	
93	
0	
41	
	1,714 genes 1,347 (79%) 367 (21%) 233 93 0 41

1859 Supplementary Table 13 Description of the RNAseq libraries used to annotate non 1860 coding RNA.

RNAseq library file	Tissues/environmental conditions	NCBI accessions
AXF_AAOSW_8_1.fastq.gz AXF_AAOSW_8_2.fastq.gz	Ecodormant buds harvested from two adult trees in 2005 (2005.12.01)	ERP004204
AXF_BAOSW_8_1.fastq.gz AXF_BAOSW_8_2.fastq.gz	Swelling buds harvested from two adult trees in 2006 (2006.24.03)	ERP004204
AXF_CAOSW_8_1.fastq.gz AXF_CAOSW_8_2.fastq.gz	Differentiating xylem sampled in April 2004 from adult trees.	ERP004204
AXF_DAOSW_7_1.fastq.gz AXF_DAOSW_7_2.fastq.gz	Roots harvested from 6-month-old seedlings after exposure to cold, heat, high CO_2 concentration, water stress and hypoxia.	ERP004204
AXF_EAOSW_7_1.fastq.gz AXF_EAOSW_7_2.fastq.gz	Leaves harvested on 6 month old seedlings after exposure to cold, heat, high CO_2 concentration, water stress and hypoxia.	ERP004204
AXF_FAOSW_7_1.fastq.gz AXF_FAOSW_7_2.fastq.gz	Dedifferentiated <i>in vitro</i> callus from genotype # DF 159	ERP004204
BHD_AAOSW_1_1.fastq.gz BHD_AAOSW_1_2.fastq.gz	White roots harvested from five-week-old sessile oak seedlings. Pool of 10 seedlings.	ERA763633
BHD_ABOSW_1_1.fastq.gz BHD_ABOSW_1_2.fastq.gz	White roots harvested from five-week-old sessile oak seedlings. Pool of 10 seedlings.	ERA763633
BHD_ACOSW_1_1.fastq.gz BHD_ACOSW_1_2.fastq.gz	White roots harvested from five-week-old sessile oak seedlings. Pool of 10 seedlings.	ERA763633
BHC_BAOSW_3_1_C4VR1ACXX.IN D15_noribo_clean.fastq.gz BHC_BAOSW_3_2_C4VR1ACXX.IN D15_noribo_clean.fastq.gz	Endodormant buds (sampled Oct. 2 nd 2013: pool of 5 sessile oak genotypes from the Laveyron population in the Pyrenees)	ERA763635
BHC_BBOSW_3_1_C4VR1ACXX.IN D16_noribo_clean.fastq.gz BHC_BBOSW_3_2_C4VR1ACXX.IN D16_noribo_clean.fastq.gz	Endodormant buds : (sampled Oct. 2 nd 2013: pool of 5 other sessile oak genotypes from the Laveyron population in the Pyrenees)	ERA763635
BHC_BCOSW_3_1_C4VR1ACXX.IN D18_noribo_clean.fastq.gz BHC_BCOSW_3_2_C4VR1ACXX.IN D18_noribo_clean.fastq.gz	Ecodormant buds : (sampled March 10 th 2014:pool of 5 sessile oak genotypes from the Laveyron population in the Pyrenees)	ERA763635
BHC_BDOSW_3_1_C4VR1ACXX.IN D19_noribo_clean.fastq.gz BHC_BDOSW_3_2_C4VR1ACXX.IN D19_noribo_clean.fastq.gz	Ecodormant buds : (sampled March 10 th 2014: pool of 5 other sessile oak genotypes from the Laveyron population in the Pyrenees)	ERA763635

Supplementary Table 14 Description of the miRNAseq libraries used to identify and validate miRNAs (NCBI bioproject accession: PRJNA361225).

miRNAseq library file (2 replicates)		Sample type		NCBI accessions
	Elevation (m)	Location/ valley/sampling date	Bud dormancy stage /genotypes pooled for library construction	
A1-Endo.fastq.gz A2-Endo.fastq.gz	1,600	Artouste/ Ossau/Oct. 7th 2013	Endodormancy A1 A2	SRR5181470 SRR5181469
A1-Eco.fastq.gz A2-Eco.fastq.gz	1,600	Artouste/ Ossau/April 7th 2014	Ecodormancy A1 A2	SRR5181464 SRR5181463
LH1-Endo.fastq.gz LH2-Endo.fastq.gz	800	Le Hourque/ Ossau/ Oct. 6th 2013	Endodormancy LH1 LH2	SRR5181472 SRR5181471
LH1-Eco.fastq.gz LH2-Eco.fastq.gz	800	Le Hourque/ Ossau/ March 16th 2014	Ecodormancy LH1 LH2	SRR5181466 SRR5181465
J1-Endo.fastq.gz J2-Endo.fastq.gz	100	Josbaig/ Ossau/ Oct. 5th 2013	Endodormancy J1 J2	SRR5181474 SRR5181473
J1-Eco.fastq.gz J2-Eco.fastq.gz	100	Josbaig/ Ossau/ March 10th 2014	Ecodormancy J1 J2	SRR5181468 SRR5181467
PR1-Endo.fastq.gz PR2-Endo.fastq.gz	1,600	Péguère/ Luz/ Oct. 4th 2013	Endodormancy PR1 PR2	SRR5181458 SRR5181457
PR1-Eco.fastq.gz PR2-Eco.fastq.gz	1,600	Péguère/ Luz/April 8th 2014	Ecodormancy PR1 PR2	SRR5181452 SRR5181451
P1-Endo.fastq.gz P2-Endo.fastq.gz	800	Papillon/ Luz/ Oct. 3rd 2013	Endodormancy P1 P2	SRR5181460 SRR5181459
P1-Eco.fastq.gz P2-Eco.fastq.gz	800	Papillon/ Luz/March 17th 2014	Ecodormancy P1 P2	SRR5181454 SRR5181453
L1-Endo.fastq.gz L2-Endo.fastq.gz	100	Laveyron/ Luz/ Oct. 2 nd 2013	Endodormancy L1 L2	SRR5181462 SRR5181461
L1-Eco.fastq.gz L2-Eco.fastq.gz	100	Laveyron/ Luz/March 13th 2014	Ecodormancy L1 L2	SRR5181456 SRR5181455

1868 Supplementary Table 15 Number of predicted and annotated ncRNA loci.

Family/sub- family		Software		#Unique
rRNA	#RNAmmer predictions	#cmsearch predictions	#Overlapping loci	#Unique
rRNA				136
5S	49	65	44	70
LSU/5.8S	13	14	7	22
SSU	20	52	20	44
tRNA	#tRNAscan-SE predictions	#cmsearch predictions	#Overlapping loci	
tRNA	827	815	790	852
miRNA	#sRNA-PIAn predictions annotated as miRNA	#cmsearch predictions	#Overlapping loci	
miRNA	1508	204	59	1594
Others	-	#cmsearch predictions		
SnoRNA				486
C/D	-	412	-	412
H/ACA	-	74	-	74
SnRNA	-		-	225
U1	-	34	-	34
U11	-	1	-	1
U2	-	55	-	55
U12	-	1	-	1
U4	-	33	-	33
U5	-	24	-	24
U6	-	64	-	64
U6atac		13	-	13
RnaseMRP	-	2	-	2
RNaseSRP	-	31	-	31

Supplementary Table 16 Distribution of the various non-coding element categories for small RNAseq data.

	1874
Non-coding elements	% aligned reads
Predicted ncRNA (P)	41.0%
LncRNA(L)	25.5%
Transposon elements (T)	38.3%
Total (P+L+T)	72.4%
	1880

1883 Supplementary Table 17 Geographical location of the natural stands from which the

1884 pedunculate oak genotypes were sampled for pool sequencing.

Site name:	ISS Landes								
Country:	France								
Latitude/Longitude:	001°05' W / 44°13' N								
Elevation:	46m								
Total area:	25,600ha								
Ecosystem:	Intensively managed								
Tree species:	Alnus, Betula, Castanea, Corylus, Crataegus, Fagus, Fraxinus, Pinus, Prunus, Quercus, Salix, Sorbus								
Land ownership:	Mainly private								
Protection:	Includes Natura 2000 sites								

1889 Supplementary Table 18 List of selected pedunculate oak genotypes used for pool 1890 sequencing.

Tree ID	Circumference (in cm at breast height)	Longitude (degrees, minutes seconds)	Latitude (degrees, minutes seconds)	Longitude (decimal format)	Latitude (decimal format)
74	139	-1.03129337	44.1717826	-1.053592703	44.288285056
352	83	-1.10264801	44.1348514	-1.174022236	44.230142753
357	156	-1.10195197	44.1347728	-1.172088818	44.229924535
358	162	-1.10178383	44.1347546	-1.171621740	44.229873982
501	137	-1.07406943	44.1248027	-1.127970645	44.213340929
521	92	-1.04406161	44.1252738	-1.077948924	44.214649407
523	137	-1.04439618	44.1252668	-1.078878276	44.214629927
602	59	-1.09019065	44.1139846	-1.150529574	44.194401571
607	206	-1.09013664	44.1145476	-1.150379563	44.195965590
1106	190	-1.05490144	44.1353328	-1.096948441	44.231480098
1108	310	-1.05472739	44.1349324	-1.096464983	44.230367702
1135	169	-1.02262027	44.1347846	-1.040611864	44.229957358
1136	138	-1.02280321	44.1346933	-1.041120023	44.229703499
1152	96	-1.00381904	44.1334381	-1.010608455	44.226216872
1153	210	-1.00406239	44.1334619	-1.011284422	44.226283131
1345	264	-1.07068438	44.1436749	-1.118567729	44.243541419
1361	255	-1.053003	44.134922	-1.091675153	44.2303391
1366	269	-1.053368	44.134774	-1.092688243	44.22992846
1410	260	-1.06359865	44.1032791	-1.109996257	44.175775217
1415	413	-1.063471	44.103621	-1.109641898	44.17672361

Supplementary Table 19 List of libraries for each of the three levels (L1, L2, L3) and number of sequences used for somatic mutation detection.

Tree		NCBI	Read length	#Raw	Total length	Mean read length
Level	Libray ID _ run ID	Accession	(before trimming)	reads	(after trimming)	(after trimming)
L1	BBX_AOSW_1_1_D1D53ACXX.IND5		101	178225679	17503027812	98.20710411
L1	BBX_AOSW_1_2_D1D53ACXX.IND5	ERX697294	101	178225679	17201970745	96.51791393
L1	BBX_AOSW_1_1_H32GMBCXX.IND5		251	129340482	31344486178	242.3408796
L1	BBX_AOSW_1_2_H32GMBCXX.IND5	ERX1886616	251	129340482	30061956368	232.4249601
L1	BBX_AOSW_2_1_H32GMBCXX.IND5		251	132162999	32027294956	242.3317812
L1	BBX_AOSW_2_2_H32GMBCXX.IND5	ERX1886622	251	132162999	30994003415	234.5134693
L1	BBX_AOSW_1_2_H57N7BCXX.IND5		251	114968046	26394294031	229.5793914
L1	BBX_AOSW_1_1_H57N7BCXX.IND5	ERX1886621	251	114968046	27764301183	241.4958082
L2	BBX_COSW_2_1_D1D53ACXX.IND7		101	187527862	18297743615	97.57346679
L2	BBX_COSW_2_2_D1D53ACXX.IND7	ERX697297	101	187527862	17809414834	94.96943358
L2	BBX_COSW_1_1_H072TAMXX.IND7		251	88896954	21647068232	243.5074236
L2	BBX_COSW_1_2_H072TAMXX.IND7	ERX697298	251	88896954	20874023496	234.8114593
L2	BBX_COSW_2_1_H072TAMXX.IND7		251	89768432	21850638929	243.4111685
L2	BBX_COSW_2_2_H072TAMXX.IND7	ERX697296	251	89768432	21059108981	234.5937042
L2	BBX_COSW_2_1_H57N7BCXX.IND7		251	141600651	34063995611	240.5638348
L2	BBX_COSW_2_2_H57N7BCXX.IND7	ERX1886620	251	141600651	32276725298	227.9419273
L3	BBX_EOSW_3_1_D1D53ACXX.IND9		101	185076013	18128331412	97.9507345
L3	BBX_EOSW_3_2_D1D53ACXX.IND9	ERX697292	101	185076013	17846097116	96.42577029
L3	BBX_EOSW_2_1_H32GLBCXX.IND9		251	134000800	28711601308	214.2644022
L3	BBX_EOSW_2_2_H32GLBCXX.IND9	ERX1886619	251	134000800	26474644233	197.5707924
L3	BBX_EOSW_1_1_H55MLBCXX.IND9		251	127908274	30449513248	238.0574164
L3	BBX_EOSW_1_2_H55MLBCXX.IND9	ERX1886617	251	127908274	28875401387	225.7508485
L3	BBX_EOSW_2_1_H55MLBCXX.IND9		251	104846648	24974555304	238.2007988
L3	BBX_EOSW_2_2_H55MLBCXX.IND9	ERX1886618	251	104846648	23191160974	221.1912485

1901 Supplementary Table 20 MuTect comparisons indicating whether candidate SNPs are expected to be detected or not, depending on the

1902 age of the mutation. L1, L2, L3 = end of selected branches; X_{L1} and X_{L2} = L1-branch and L2-branch initiation sites (see also Fig. 2b).

			MuTect comparisons (reference vs. potentially mutated libraries)								
		Colored tree section									
		in Fig. 2b	L1 vs. L2	L1 vs. L3	L2 vs. L1	L2 vs. L3	L3 vs. L1	L3 vs. L2			
Mutations	$X_{L1}-X_{L2} \\$	blue	Х	Х	Ø	Ø	Ø	Ø			
occurring	$X_{L2} - L3$	pink	Ø	Х	Ø	Х	Ø	Ø			
between	$X_{L1} - L1$	green	Ø	Ø	Х	Ø	Х	Ø			
levels:	$X_{L2} - L2$	yellow	Х	Ø	Ø	Ø	Ø	Х			

1906 Supplementary Table 21 List of the 15 eudicot plant genomes selected for the evolutionary analysis. Growth habit or lifespan (W: woody perennials vs. H: annual herbaceous species) is indicated in the last column.

Scientific name	Common name	# of genes	Assembly version	Order	Family	Genus	Growth habit
Arabidopsis lyrata	Lyrate rockcress	32,657	v1.0	Brassicales	Brassicaceae	Arabidopsis	Н
Arabidopsis thaliana	Thale cress	27,416	TAIR10	Brassicales	Brassicaceae	Arabidopsis	Н
Citrullus lanatus	Watermelon	23,440	v1	Cucurbitales	Cucurbitaceae	Citrullus	Н
Fragaria vesca	Strawberry	32,831	v1.1	Rosales	Rosaceae	Fragaria	Н
Glycine max	Soybean	56,044	Wm82.a2.v1	Fabales	Fabaceae	Glycine	Н
Ricinus communis	Castorbean	31,221	v0.1	Malpighiales	Euphorbiaceae	Ricinus	Н
Solanum tuberosum	Potato	35,119	v3.4	Solanales	Solanaceae	Solanum	Н
Carica papaya	Papaya	27,584	ASGPBv0.4	Brassicales	Caricaceae	Carica	W
Citrus clementina	Clementine	24,533	v1.0	Sapindales	Rutaceae	Citrus	W
Eucalyptus grandis	Eucalyptus	36,376	v2.0	Myrtales	Myrtaceae	Eucalyptus	W
Malus domestica	Apple	63,514	v1.0	Rosales	Rosaceae	Malus	W
Populus trichocarpa	Poplar	41,335	v3.0	Malpighiales	Salicaceae	Populus	W
Prunus persica	Peach	27,864	v2.1	Rosales	Rosaceae	Prunus	W
Theobroma cacao	Cocoa	29,452	v1.1	Malvales	Malvaceae	Theobroma	W
Vitis vinifera	Grape	26,346	Genoscope_12X	Vitales	Vitaceae	Vitis	W

Supplementary Table 22 Contribution of gene models for the 16 studied species to orthoMCL orthogroups. 1912

Growth	Snecies	Abbreviation -	Genes in orthogroups				# orthogroups		#species- specific	
habit	species	ADDICVICTION	Total	Mean	SD	Max	(%)		orthogroups (%)	
	Arabidopsis lyrata	Al	27,260	0.74	1.67	96	19,658	(49.8)	813	(2.1)
	Arabidopsis thaliana	At	24,733	0.67	1.56	124	20,128	(51.0)	141	(0.4)
	Citrullus lanatus	Wa	20,192	0.55	2.66	363	24,069	(61.0)	364	(0.9)
Herbaceous species	Fragaria vesca	Fv	25,093	0.68	2.31	167	22,540	(57.2)	1,275	(3.2)
T	Glycine max	Gm	46,400	1.26	3.68	295	21,609	(54.8)	1,552	(3.9)
	Ricinus communis	Rc	21,088	0.57	1.23	79	22,186	(56.3)	754	(1.9)
	Solanum tuberosum	St	28,897	0.78	7.01	1062	23,803	(60.4)	1,060	(2.7)
	Carica papaya	Ср	20,285	0.55	2.63	395	23,361	(59.2)	520	(1.3)
	Citrus climentina	Cc	21,425	0.58	1.95	130	22,928	(58.1)	316	(0.8)
	Eucalyptus grandis	Eg	29,063	0.79	3.59	228	23,229	(58.9)	722	(1.8)
	Malus domestica	Md	46,524	1.26	4.04	378	19,627	(49.8)	3,324	(8.4)
Woody perennials	Populus trichocarpa	Pt	33,604	0.91	2.75	183	21,923	(55.6)	728	(1.8)
pereninais	Prunus persica	Рр	24,651	0.67	5.37	907	22,299	(56.5)	311	(0.8)
	Quercus robur	Qr	22,498	0.61	3.31	359	25,031	(63.5)	479	(1.2)
	Theobroma cacao	Tc	23,608	0.64	2.46	208	22,253	(56.4)	465	(1.2)
	Vitis vinifera	Vv	19,774	0.54	1.49	105	23,893	(60.6)	589	(1.5)

Supplementary Table 23 Major family of repetitive elements identified by RepeatMasker within the sequenced BAC clones.

	Length	
Family	(bp)	%
DNA transposon	289,344	36
hAT	54,917	6
EnSpm/CACTA	38,913	4
MuDR	36,662	4
Helitron	35,283	4
Harbinger	17,583	2
Polinton	17,078	2
Mariner/Tc1	12,792	1
Retrotransposon	504,226	63
-LTR Retrotransposon	361,480	45
Gypsy	222,020	27
Copia	126,984	15
-Non-LTR Retrotransposon	142,746	17
Total length of repeats	794,208	

Supplementary Table 24 Oak BAC sequence statistics.

Total sequence length	4,344,182 bp
Sequence length excluding stretches of Ns	4,282,332 bp (number of Ns: 61,850)
GC content %	35.9
Number of predicted protein coding genes	$198^1, 50^2, 30^3$
Number of predicted protein coding genes	198
with homology to oak unigene ⁴	
tRNA genes	4
Gene density	6 genes/100 kb
Mean gene length	$4,028 \text{ bp}^5$
Mean number of exons per gene	5.4
Mean exon length	232 bp
% of genes with introns	83.5
Average intron length	615 bp
¹ Approved: gene structure was modified or validated a ² Problematic: gene structure remains after manual cura ³ deleted ⁴ from Lesur et al. ²³ ⁵ UTRs were not considered.	fter manual curation. ation.

Supplementary Table 25 Summary of overlapping regions between allelic BACs. BAC1 and BAC2 referred to pairs of allelic BACs.

BAC 1	BAC 2	% of BAC 1 covered	% identity	E-value	Range of overlap BAC 1 (bp)	Length of overlapping region_BAC 1 (bp)	Range of overlap BAC 2	Length of overlapping region_BAC 2 (bp)
50E24	177A20	44	98	0.0	74,218-140,871	66,655	21,871-75,811	53,940
5E10	107I07	43	97	0.0	34-52,488	52,454	11,315-86,163	74,848
12J1	121F17	50	97	0.0	3,328-69,071	65,743	1-107,378	107,378
27L03	48K1	27	97	0.0	72-22,592	22,520	94,896-110,662	25,766
64H3	30P1	55	99	0.0	11,197-105,888	94,691	1-87,454	87,454
Supplementary Table 26 Results of BLAST-n alignment (Evalue=0 and identity >95%) between overlapping BAC regions. BAC1 and BAC2 are pairs of allelic BAC.

BAC 1 (start-end)	BAC 2 (start-end)	% identity	E-value
50E24 (82472-97166)	177A20 (101617-86956)	98.313	0.0
50E24 (97342-98711)	177A20 (86962-85573)	97.557	0.0
50E24 (99445-102142)	177A20 (85526-82785)	95.796	0.0
50E24 (102139-102632)	177A20 (77962-77469)	98.178	0.0
50E24 (102630-104811)	177A20 (72280-70100)	99.313	0.0
50E24 (104964-113458)	177A20 (67500-59008)	97.533	0.0
50E24 (115554-119919)	177A20 (59015-54685)	93.276	0.0
50E24 (120139-121173)	177A20 (54688-53673)	88.509	0.0
50E24 (121653-122102)	177A20 (43131-42688)	95.778	0.0
50E24 (122088-130991)	177A20 (42426-33477)	96.247	0.0
50E24 (130983-137845)	177A20 (31276-24398)	95.750	0.0
50E24 (138678-140871)	177A20 (24080-21871)	97.473	0.0
5E10 (34-13493)	107I07 (86163-72751)	96.690	0.0
5E10 (6111-8673)	107I07 (96063-93431)	89.234	0.0
5E10 (15109-16637)	107I07 (72758-71221)	95.596	0.0
5E10 (16633-19460)	107I07 (60951-58146)	95.046	0.0
5E10 (19562-20049)	107I07 (58152-57640)	93.177	0.0
5E10 (20042-25226)	107I07 (57568-52375)	97.546	0.0
5E10 (20099-30015)	107I07 (48048-38134)	95.357	0.0
5E10 (30710-44385)	107I07 (37408-23814)	95.014	0.0
5E10 (44376-47570)	107I07 (18423-15229)	97.444	0.0
5E10 (47654-51561)	107I07 (15245-11315)	95.392	0.0
5E10 (51556-52488)	107I07 (929-1)	98.178	0.0
12J1 (3328-9386)	121F17 (107378-101370)	97.776	0.0
12J1 (10865-15722)	121F17 (101374-96618)	95.163	0.0
12J1 (15714-16429)	121F17 (96088-95351)	92.473	0.0
12J1 (17136-17988)	121F17 (94236-93385)	97.541	0.0
12J1 (17555-25498)	121F17 (92576-84615)	97.074	0.0
12J1 (25929-36992)	121F17 (67374-56308)	97.545	0.0
12J1 (27681-28891)	121F17 (111037-109862)	94.403	0.0
12J1 (40309-55775)	121F17 (51089-35673)	97.417	0.0
12J1 (57057-65797)	121F17 (12263-3515)	97.174	0.0
12J1 (65796-69071)	121F17 (3253-1)	97.063	0.0
27L03 (72-1944)	48K1 (94896-96762)	97.340	0.0
27L03 (1937-6237)	48K1 (97857-102122)	97.846	0.0
27L03 (6914-13853)	48K1 (102111-109054)	97.113	0.0
27L03 (14881-16542)	48K1 (109051-110662)	92.123	0.0
27L03 (15395-16915)	48K1 (70312-71817)	90.582	0.0
27L03 (20326-20919)	48K1 (95799-96389)	93.311	0.0
27L03 (22071-22592)	48K1 (102497-103027)	90.038	0.0
64H3 (11197-22071)	30P1 (87454-76608)	98.232	0.0
64H3 (27384-29525)	30P1 (71127-68965)	98.661	0.0

64H3 (29522-35146)	30P1 (61773-67407)	96.079	0.0
64H3 (35355-37869)	30P1 (61236-58733)	98.648	0.0
64H3 (46894-49574)	30P1 (50014-47328)	97.993	0.0
64H3 (60365-80565)	30P1 (46359-26158)	99.975	0.0
64H3 (83318-94716)	30P1 (17599-6207)	99.073	0.0
64H3 (94715-97846)	30P1 (4554-1423)	100.000	0.0
64H3 (99674-100979)	30P1 (1322-1)	95.925	0.0

1942	Supplementary Table 27 Metrics of the previous release (V1) and current release (V2) of
1943	the oak diploid genome assembly.

	Diploid V1 ^b	Diploid V2
Assembly	454 + Illumina	454 + Illumina + Synthetic Long Reads
No. of sequences	17,910	8,827
Cumulative size	1,354,311,717	1,455,104,916
N50	256,640	821,707
N90	35,065	198,501
L50	1,468	537
L90	6,626	1,880
% of N's	11.56	4.6
Completeness using BUSCO	274 (90.4%)	275 (90.8%)
Oak RNA-seq genes (90,786 contigs) ^a	86,457 (95.2%)	86,488 (95.3%)

^a from Lesur et al. ²³, ^b from Plomion et al. ¹⁹

1946 Supplementary Table 28 Classification of marker-scaffold relationships into four
1947 categories. The number of markers and the number of scaffolds within each category are
1948 provided.

Scaffold- marker relationship	Comment	Number of markers/2,615	Number of scaffolds (cumulative size Mb)
Category#1	Scaffold anchored with a single marker	165	165 (90.8)
Category#2	Scaffold anchored with at least 2 markers from the same LG	1412	331 (320)
Category#3	Scaffold anchored with more than 50% of the markers from the same LG	898	116 (214)
Category#4	Scaffolds (unassigned) with less than 50% of the markers from the same LG	140	116 (46.7)

1952 Supplementary Table 29 Rank correlations (rho) between genetic and physical positions along the 12 chromosomes. LG: linkage group.
1953

		No. of	#markers on	Chr_start	Chr_end	LG_start	LG_end	
LG	size (cM)	markers	chromosomes	(bp)	(bp)	(cM)	(cM)	rho
1	66.43	421	320	223,913	55,067,536	0.64	66.43	0.998
2	103.92	922	676	79,368	115,173,360	0.03	103.92	0.999
3	75.98	400	281	244,065	57,437,871	6.3	75.98	0.998
4	75.7	291	171	339,968	44,508,357	3.42	75.42	0.994
5	85.84	398	263	90,378	70,598,779	0.61	85.41	0.998
6	74.87	537	409	201,326	55,995,377	0.64	74.87	0.998
7	65.26	419	321	75,105	51,549,230	1.8	63.5	0.998
8	70.8	572	459	105,078	71,279,127	1.86	70.2	0.998
9	68.7	400	273	118,866	50,074,090	0.94	68.7	0.996
10	66.8	381	284	332,011	50,211,705	0.62	66.8	0.998
11	66.46	391	316	451,420	51,991,272	1.08	66.46	0.991
12	66.82	457	297	677	39,751,979	0.29	66.82	0.996
Total	887.58	5,589	4,070		711,376,508		866.28	0.997

1957 Supplementary Table 30 Transposable element annotation: comparison between the 16 eudicot species used in this study.

Scientific name	Woody/ Herbaceous	Common name	Ref.	Assembly length annotated (Mb)	TE (Mb)	TE %	LTR % TE	Non-LTR % TE	Other class I % TE	Class I % TE	Class II % TE	Other % TE	
Arabidopsis lyrata	Н	Arabidopsis lyrata	38	207	61	29.7	64	8	0	72	25	3	
Arabidopsis thaliana	Н	Thale cress	38	135	32	23.7	50	28	0	78	6	16	
Citrullus lanatus	Н	Watermelon	153	354	160	45.2	Major	NA	NA	Major	NA	NA	
Fragaria vesca	Н	Strawberry	154	209	48	22.81	69.8	2	0	71.8	28.2	0	
Glycine max	Н	Soybean	155	955	561	58.7	71.5	0.4	0	71.9	28.1	0	
Ricinus communis	Н	Castor bean	156	350	176	50.3	32.2	0.3	3.6	36.1	1.8	62.1	
Solanum tuberosum	Н	Potato	157	727	452	62.2	45.93	4.51	0	50.4	6.2	43.4	
Carica papaya	W	Papaya	42	815	423	51.9		77	0	77	0.2	22.8	
Citrus clementina	W	Clementine	43	816	347	42.5	47	2.9	0	49.9	6.3	43.8	
Eucalyptus grandis	W	Eucalyptus	41	817	409	50	73	7	5	85	11	4	
Malus domestica	W	Apple	40	818	347	42.4	73.4	15.3	0	88.7	2.1	9.2	
Populus trichocarpa	W	Poplar	158	820	362	44.2	18.6	1.4	0.1	20.1	6.1	73.8	
Prunus persica	W	Peach	39	226	67	29.6	66.1	2.1	1.2	69.4	30.6	0	
Theobroma cacao	W	Cocoa bean	159	346	144	41.5	77.9	0.4	0	78.3	21.7	0	
Vitis vinifera	W	Grape	160	467	193	41.4	55.9	9.5	1.3	66.7	2	31.3	
Quercus robur	W	Oak	this study	814	421	52	53.1	16	1	70.1	15.2	14.7	

1960 Supplementary Table 31 Oak gene structure statistics.

Total protein coding genes	25,808
Gene space (Mb)	75
Gene density (# genes / 10 kb)	0.32
Gene mean / median (bp)	2,907
Gene median (bp)	2,137
CDS mean (bp)	1,174
CDS median (bp)	942
#CDS < 500 bp	4,367
#CDS > 3 kb	1,162
Genes with introns (%)	79%
#Introns/gene (mean)	3.3

Supplementary Table 32 Horizontal transfers of LTR retrotransposons between oak and other plant species.

Name of the LTR-retrotransposon family	Species involved in the transfer
RLX-incomp_Qrob_v2_More29k-B-R2774-Map5_reversed	oak / grapevine
RLX-incomp-chim_Qrob_v2_More29k-B-R25479-Map5_reversed	oak / grapevine
RLX-incomp-chim_Qrob_v2_More29k-B-R32795-Map5_reversed	oak / grapevine
RLX-comp_Qrob_v2_More29k-B-G5453-Map6_reversed	oak / grapevine
RLX-comp_Qrob_v2_More29k-B-P1015.803-Map7	oak / grapevine
RLX-incomp_Qrob_v2_More29k-B-R289-Map20_reversed	oak / grapevine
RLX-comp_Qrob_v2_More29k-B-R13571-Map19	oak / poplar
RLX-incomp_Qrob_v2_More29k-B-R2774-Map5_reversed	oak / grapevine / peach tree

1970 Supplementary Table 33 List of putative aquaporins identified in the pedunculate oak genome (haplome assembly). Number of exons and 1971 protein length (in amino-acids) are given. The aromatic/arginine selectivity filter (H, transmembrane helix and LE, loop E), the NPA motifs (LB, 1972 loop B and LE, loop E) and the five Froger's positions were identified from multiple sequence alignments.

				Ar/l	R filter	•	NPA	NPA motif		Froger's position					
gene model ID	N° exon	protein (AA)	H2	Н5	LE1	LE2	LB	LE	Р 1	P 2	P3	P4	P5	subclass	remarks
Qrob_T0687390.2	5	277	W	V	А	R	NPA	NPA	F	S	А	Y	М	NIP1	
Qrob_T0687410.2	5	289	W	V	А	R	NPS	NPA	F	Т	А	Y	М	NIP1	
Qrob_T0405130.2	5	261	W	V	А	R	NPA	NPA	F	S	А	Y	Ι	NIP4	GC at ex/intron boundary
Qrob_T0144140.2	5	273	W	V	А	R	NPA	NPA	F	S	А	Y	V	NIP4	
Qrob_T0275880.2 (¹)	5	268	W	V	А	R	NPA	NPA	F	S	А	Y	V	NIP4	
Qrob_T0748200.2 (²)	4	298	А	Ι	G	R	NPS	NPV	F	Т	А	F	L	NIP5	
Qrob_T0118430.2	5	304	Т	Ι	G	R	NPA	NPV	F	Т	А	Y	Μ	NIP6	
Qrob_T0697150.2	5	282	А	V	G	R	NPA	NPA	Y	S	А	Y	V	NIP7	
Qrob_T0345370.2 (*)	4	285	F	Н	Т	R	NPA	NPA	G	S	А	F	W	PIP1	
Qrob_T0236650.2	4	289	F	Н	Т	R	NPA	NPA	E	S	А	F	W	PIP1	
Qrob_T0705530.2	4	286	F	Н	Т	R	NPA	NPA	E	S	А	F	W	PIP1	
Qrob_T0348530.2	4	287	F	Н	Т	R	NPA	NPA	Q	S	А	F	W	PIP1	
Qrob_T0373060.2	4	278	F	Н	Т	R	NPA	NPA	Μ	S	А	F	W	PIP2	
Qrob_T0438960.2	4	262	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	GC at ex/intron boundary

Qrob_T0438980.2	4	262	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	GC at ex/intron boundary
Qrob_T0438970.2	4	262	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	GC at ex/intron boundary
Qrob_T0438950.2	4	262	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	GC at ex/intron boundary
Qrob_T0438990.2	4	286	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	
Qrob_T0602100.2	4	287	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	
Qrob_T0530060.2	4	281	F	Н	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	
Qrob_T0131450.2	4	281	F	Η	Т	R	NPA	NPA	А	S	А	F	W	PIP2	
Qrob_T0602110.2	4	285	F	Η	Т	R	NPA	NPA	Q	S	А	F	W	PIP2	
Qrob_T0237440.2	1	239	А	V	Р	Ν	NPS	NPA	Р	А	А	Y	W	SIP	
Qrob_T0714870.2	3	241	Ι	М	Р	Ν	NPT	NPA	Р	А	А	Y	W	SIP	
Qrob_T0098460.2	3	237	S	Η	G	S	NPL	NPA	Р	V	А	Y	W	SIP	
Qrob_T0108440.2	3	252	Н	Ι	А	V	NPA	NPA	Т	S	А	Y	W	TIP1	
Qrob_T0398210.2	3	252	Η	Ι	А	V	NPA	NPA	Т	S	А	Y	W	TIP1	
Qrob_T0119780.2	2	251	Η	Ι	А	V	NPA	NPA	Т	S	А	Y	W	TIP1	
Qrob_T0656320.2	2	253	Η	Ι	А	V	NPA	NPA	Т	S	А	Y	W	TIP1	
Qrob_T0412470.2	3	248	Η	Ι	G	R	NPA	NPA	Т	S	А	Y	W	TIP2	
Qrob_T0538460.2	3	250	Η	Ι	G	R	NPA	NPA	Т	S	А	Y	W	TIP2	
Qrob_T0264600.2	3	246	Н	Ι	А	R	NPA	NPA	Т	S	А	Y	W	TIP4	
Orob T0082220.2 ([*])	3	234	Н	Ι	А	R	NPA	NPA	Т	S	А	Y	W	TIP4	T deletion -> variant protein
Orob T0375680.2	3	254	N	V	G	С	NPA	NPA	v	S	А	Y	W	TIP5	L
 — 		-			-	-				-				-	

Qrob_T0158140.2	1	236	V	V	А	R	NPM	NPA	Μ	С	А	F	W	XIP2	
Qrob_T0158150.2 (*)	1	262	V	Ι	V	G	SPE	NPA	Μ	С	А	F	W	XIP2	
Qrob_T0158180.2	2	307	Ι	Ι	А	Κ	SPI	NPA	Μ	С	А	F	W	XIP2	
Qrob_T0158190.2	2	295	Ι	Ι	V	Κ	SPI	NPA	Μ	С	А	F	W	XIP2	
Qrob_T0158200.2	3	334	Ι	Т	V	R	NPA	NPA	V	С	А	F	W	XIP1	
Qrob_T0656330.2	2	214						NPA						Invalid	unreliable reading frame

(¹) Sequence analysis was performed after the manual merging of Qrob_T0275880.2 and Qrob_T0275890.2.

(²) Due to poor sequence quality, sequence analysis was performed on its allelic version Qrob_T0751510.2 (qrob_v2_scaffold_2295:14651-1620 following manual curation). (*) Sequence analysis was performed after the manual curation of intron/exon prediction.

1974

1976 Supplementary Table 34 List of R2R3-MYB, MYB-3R and MYB-4R identified in the 1977 pedunculate oak genome (haplome assembly). MYB predicted proteins were retrieved by three different approaches: those containing the MYB domain with the Pfam signature 1978 1979 PF00249, those automatically annotated as MYB proteins, and those with homology to one of the Arabidopsis R2R3-MYB proteins after a BLAST-p search with an e-value of 10e⁻¹⁰ as the 1980 1981 threshold. The predicted proteins identified were inspected and manually curated. R2R3-MYB genes were named with consecutive numbers starting from the first gene on the first 1982 chromosome scaffold (QroMYB1 - QroMYB129) to the genes not assigned to any 1983 1984 chromosome (QroMYB130 - QroMYB139). 3R-MYB and 4R-MYB genes are named with 1985 letters in alphabetical order, also starting from the first gene on the first chromosome scaffold (OroMYB3R-A to OroMYB3R-E, and OroMYB4R-A). 1986

1987

MYB ID	Transcript id	MYB Subgroup	Scaffold_ID on H2.3	Pseudomo lecule	Gene start	Gene end	Gene length (in bp UTR + CDS + introns)
	Transer pr_1a	SusBroup	Orob H2 3 Sc000	loculo	5	othe the	(((((((((((((((((((((((((((((((((((((((
QrobMYB1	Qrob_T0404990.2	WPS-III	0317 Orob H2 3 Se000	Chr1	414608	415980	1373
QrobMYB2	Qrob_T0371530.2	SAtMYB71	0141	Chr1	5600909	5602170	1262
QrobMYB3	Qrob_T0371540.2	SAtMYB71	Qrob_H2.5_Scooo 0141	Chr1	5606927	5608155	1229
QrobMYB4	Qrob_T0371920.2	SAtM5	Qrob_H2.3_Sc000 0122	Chr1	9166327	9164971	1357
QrobMYB5	Qrob_T0731180.2	S14	Qrob_H2.3_Sc000 0439	Chr1	17362712	17364185	1474
QrobMYB6	Qrob_T0252590.2	SAtM80	Qrob_H2.3_Sc000 0299	Chr1	17427162	17425846	1317
QrobMYB7	Qrob_T0252570.2	SAtM80	Qrob_H2.3_Sc000 0299	Chr1	17472705	17471387	1319
QrobMYB8	Qrob_T0252550.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17515334	17513768	1567
QrobMYB9	Qrob_T0252540.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17534040	17532476	1565
QrobMYB10	Qrob_T0252530.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17555342	17553771	1572
QrobMYB11	Qrob_T0252520.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17595722	17593968	1755
QrobMYB12	Qrob_T0252500.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17665018	17663242	1777
QrobMYB13	Qrob_T0252490.2	WPS-V	Qrob_H2.3_Sc000 0299	Chr1	17716131	17714566	1566
QrobMYB14	Qrob_T0595370.2	SAtM91	Qrob_H2.3_Sc000 0542	Chr1	28175476	28174268	1209
OrobMYB15	Orob T0660350.2	S5	Qrob_H2.3_Sc000 0038	Chr1	39742573	39743584	1012
OrobMYB16	Orob T0402380.2	S2 & S3	Qrob_H2.3_Sc000 0332	Chr1	45752069	45750448	1622
OrobMYB17	Orob T0307500.2	S25	Qrob_H2.3_Sc000 0054	Chr1	49689933	49692324	2392
C	C		Qrob_H2.3_Sc000				
QrobMYB18	Qrob_T0722940.2	S14	0511 Qrob_H2.3_Sc000	Chr2	4595691	4597686	1996
QrobMYB19	Qrob_T0059750.2	S22	0025 Qrob_H2.3_Sc000	Chr2	5496656	5495769	888
QrobMYB20	Qrob_T0022470.2	S18	0016 Orob H2.3 Sc000	Chr2	10726752	10728232	1481
QrobMYB21	Qrob_T0270990.2	S19	0040 Orob H2 3 Sc000	Chr2	18266538	18263431	3108
QrobMYB22	Qrob_T0304630.2	S5	0158 Orob H2 3 Sc000	Chr2	26868539	26867340	1200
QrobMYB23	Qrob_T0304650.2	S5	0158 Orob H2 3 Sc000	Chr2	26893414	26894979	1566
QrobMYB24	Qrob_T0304670.2	S5	0158 Oreh 112.2 S-000	Chr2	26920641	26921682	1042
QrobMYB25	Qrob_T0304700.2	S5	Q100_H2.5_SC000 0158	Chr2	26949901	26950860	960

			Orob H2 3 Sc000				
QrobMYB26	Qrob_T0304800.2	WPS-I	0158	Chr2	27068410	27067516	895
QrobMYB27	Qrob_T0351500.2	SAtM35	Qrob_H2.3_Scooo 0145	Chr2	32194944	32197138	2195
QrobMYB28	Qrob_T0418840.2	S14	Qrob_H2.3_Sc000 0192	Chr2	39581531	39580189	1343
QrobMYB29	Qrob_T0178010.2	WPS-II	Qrob_H2.3_Sc000 0043	Chr2	41309398	41310479	1082
QrobMYB30	Qrob_T0121400.2	SAtM40	Qrob_H2.3_Sc000 0083	Chr2	47295425	47296526	1102
QrobMYB31	Qrob_T0203360.2	WPS-II	Qrob_H2.3_Sc000 0076	Chr2	53133326	53134402	1077
QrobMYB32	Qrob_T0562460.2	WPS-II	Qrob_H2.3_Sc000 0524	Chr2	55606964	55605873	1092
QrobMYB33	Qrob_T0395770.2	S14	Qrob_H2.3_Sc000 0314	Chr2	55965127	55966278	1152
QrobMYB34	Qrob_T0398080.2	S6	Qrob_H2.3_Sc000 0207	Chr2	69744225	69747170	2946
QrobMYB35	Qrob_T0459570.2	WPS-V	Qrob_H2.3_Sc000 0287	Chr2	71044675	71046349	1675
QrobMYB36	Qrob_T0459610.2	WPS-V	Qrob_H2.3_Sc000 0287	Chr2	71143342	71141657	1686
OrobMYB37	Orob T0324610.2	S5	Qrob_H2.3_Sc000 0127	Chr2	72547948	72549947	2000
OrobMYB38	Orob T0324630.2	\$5	Qrob_H2.3_Sc000 0127	Chr2	72611633	72614513	2881
OrobMVB39	Qrob_T0324680.2	\$5	Qrob_H2.3_Sc000	Chr2	72858478	72860109	1632
OrobMVB40	Qrob_T0365070.2	S1	Qrob_H2.3_Sc000	Chr2	85503235	85594741	1507
OrobMVB41	Qrob_T0102440.2	51 50a	Qrob_H2.3_Sc000	Chr2	87682086	87684183	2008
OrobMVB42	Qrob_T0105940.2	S15	Qrob_H2.3_Sc000	Chr2	80155341	80154041	1301
QrobMVD42	Qrob_T0195940.2	S15 SA4M92	Qrob_H2.3_Sc000	Chr2	06414206	06415092	1679
QrodM Y B43	QF05_10245380.2	SAUM82	Qrob_H2.3_Sc000	Chr2	10658008	96415985 10658160	10/8
QrobM Y B44	Qrob_102/8/40.2	514	Qrob_H2.3_Sc000	Chr2	11126528	1 11126345	1516
QrobMYB45	Qrob_10018660.2	S5	0022 Qrob_H2.3_Sc000	Chr2	5	8	1828
QrobMYB46	Qrob_T0170360.2	S15	0015 Qrob_H2.3_Sc000	Chr3	49517391	49516013	1379
QrobMYB47	Qrob_T0202260.2	S9a	0176 Qrob_H2.3_Sc000	Chr3	54676627	54673958	2670
QrobMYB48	Qrob_T0038250.2	SAtM46	0070 Qrob_H2.3_Sc000	Chr4	3427089	3424578	2512
QrobMYB49	Qrob_T0642710.2	S18	0629 Orob H2.3 Sc000	Chr4	24187005	24190552	3548
QrobMYB50	Qrob_T0641860.2	WPS-I	0468 Orob H2.3 Sc000	Chr5	4569573	4570457	885
QrobMYB51	Qrob_T0641880.2	WPS-I	0468 Orob H2.3 Sc000	Chr5	4694801	4695685	885
QrobMYB52	Qrob_T0641900.2	WPS-I	0468 Orob H2 3 Sc000	Chr5	4777403	4778287	885
QrobMYB53	Qrob_T0641910.2	S5	0468 Orob H2 3 Sc000	Chr5	4820012	4818765	1248
QrobMYB54	Qrob_T0070380.2	SAtM35	0056 Orob H2 3 Sc000	Chr5	18540876	18538781	2096
QrobMYB55	Qrob_T0523450.2	SAtMYB26	0464 Oreb H2 3 Se000	Chr5	23855006	23856326	1321
QrobMYB56	Qrob_T0221460.2	WPS-V	0055 Orab 112.3 Se000	Chr5	26538372	26536935	1438
QrobMYB57	Qrob_T0108420.2	SAtMYB27	0198 Oreb 112.3 Se000	Chr5	33148198	33147172	1027
QrobMYB58	Qrob_T0072890.2	S11	0072 Orab 112.2 Se000	Chr5	40907218	40905536	1683
QrobMYB59	Qrob_T0653450.2	SAtMYB26	QF0B_H2.3_SC000 0325	Chr5	45794689	45793005	1685
QrobMYB60	Qrob_T0697670.2	SAtMYB26	0325 0rsh 112.2 S 000	Chr5	45851239	45849785	1455
QrobMYB61	Qrob_T0697680.2	SAtMYB26	Qrob_H2.3_SC000 0325 Orab_U2.2_5_000	Chr5	45875202	45873379	1824
QrobMYB62	Qrob_T0697730.2	SAtMYB26	Qrob_H2.3_SC000 0325	Chr5	45929859	45928201	1659
QrobMYB63	Qrob_T0701860.2	S11	Qrob_H2.3_Sc000 0424	Chr5	65885040	65883694	1347

QrobMYB64	Qrob_T0058400.2	S5	Qrob_H2.3_Sc000 0065	Chr5	67078130	67081148	3019
QrobMYB65	Qrob_T0577750.2	S 7	Qrob_H2.3_Sc000 0053	Chr6	11404917	11400280	4638
QrobMYB66	Qrob_T0005810.2	S15	Qrob_H2.3_Sc000 0006	Chr6	21152052	21153310	1259
QrobMYB67	Qrob_T0047220.2	S25	Qrob_H2.3_Sc000 0006	Chr6	21502446	21504146	1701
QrobMYB68	Qrob_T0379650.2	SAtMYB71	Qrob_H2.3_Sc000 0566	Chr6	25445570	25447050	1481
QrobMYB69	Qrob_T0690000.2	SAtM46	Qrob_H2.3_Sc000 0565	Chr6	27337413	27334918	2496
QrobMYB70	Qrob_T0199530.2	WPS-I	Qrob_H2.3_Sc000 0179	Chr6	43171185	43172069	885
QrobMYB71	Qrob_T0199740.2	S22	Qrob_H2.3_Sc000 0179	Chr6	43505358	43506296	939
QrobMYB72	Qrob_T0346410.2	S4	Q100_H2.5_SC000 0011 Orob_H2.3_Sc000	Chr6	53280743	53279865	879
QrobMYB73	Qrob_T0418350.2	S10 & S24	0204 0rob H2 3 Sc000	Chr6	56246814	56245513	1302
QrobMYB74	Qrob_T0738480.2	S 1	0662 0rob H2 3 Sc000	Chr7	9584571	9583099	1473
QrobMYB75	Qrob_T0119930.2	SAtMYB71	0123 Orob H2 3 Se000	Chr7	10575199	10573946	1254
QrobMYB76	Qrob_T0388360.2	S14	0367 Orob H2 3 Se000	Chr7	47727255	47728879	1625
QrobMYB77	Qrob_T0657180.2	SAtM91	0478 Orob H2 3 Sc000	Chr7	51905061	51903988	1074
QrobMYB78	Qrob_T0033520.2	S9b	0007 0rob H2 3 Sc000	Chr8	19882307	19880196	2112
QrobMYB79	Qrob_T0626620.2	S15	0156 Orob H2 3 Sc000	Chr8	26379626	26380864	1239
QrobMYB80	Qrob_T0647280.2	S18	0283 Orob H2 3 Sc000	Chr8	36441820	36438939	2882
QrobMYB81	Qrob_T0654710.2	WPS-V	0642 Orob H2 3 Sc000	Chr8	38444539	38445951	1413
QrobMYB82	Qrob_T0668650.2	WPS-V	0642 Orob H2 3 Sc000	Chr8	38539863	38541297	1435
QrobMYB83	Qrob_T0668640.2	WPS-V	0642 Orob H2 3 Sc000	Chr8	38597083	38598848	1766
QrobMYB84	Qrob_T0466400.2	S11	0570 Orob H2 3 Sc000	Chr8	40545518	40544193	1326
QrobMYB85	Qrob_T0436030.2	S22	0008 0rob H2 3 Sc000	Chr8	47999625	48000389	765
QrobMYB86	Qrob_T0437590.2	WPS-II	0334 Orob H2 3 Sc000	Chr8	51157045	51155572	1474
QrobMYB87	Qrob_T0303790.2	S23	0251 Orob H2 3 Sc000	Chr8	66214685	66211087	3599
QrobMYB88	Qrob_T0411100.2	S21	0251 Orob H2 3 Sc000	Chr8	66330876	66332389	1514
QrobMYB89	Qrob_T0277200.2	S21	0457 Orob H2 3 Sc000	Chr9	4255611	4253903	1709
QrobMYB90	Qrob_T0344270.2	S5	0600 Orob H2.3 Sc000	Chr9	5365424	5366566	1143
QrobMYB91	Qrob_T0344260.2	S5	0600 Orob H2.3 Sc000	Chr9	5379324	5380656	1333
QrobMYB92	Qrob_T0344200.2	S 4	0600 Orob H2.3 Sc000	Chr9	5457882	5456407	1476
QrobMYB93	Qrob_T0191710.2	S5	0155 Orob H2.3 Sc000	Chr9	7921886	7920786	1101
QrobMYB94	Qrob_T0612820.2	S16	0047 Orob H2.3 Sc000	Chr9	14775500	14778940	3441
QrobMYB95	Qrob_T0612850.2	S9b	0047 Orob H2.3 Sc000	Chr9	14807889	14810821	2933
QrobMYB96	Qrob_T0575950.2	S2 & S3	0017 Qrob_H2.3_Sc000	Chr9	17357797	17359638	1842
QrobMYB97	Qrob_T0246270.2	SAtMYB88	0017 Qrob_H2.3_Sc000	Chr9	19640731	19634180	6552
QrobMYB98	Qrob_T0464600.2	SAtMYB85	0446 Qrob_H2.3_Sc000	Chr9	20541214	20543134	1921
QrobMYB99 QrobMYB10	Qrob_T0123830.2	S10 & S24	0112 Qrob_H2.3_Sc000	Chr9	24701012	24702528	1517
0 QrobMYB10	Qrob_T0460820.2	S21	0152 Qrob_H2.3_Sc000	Chr9	29184636	29183320	1317
1	Qrob_T0369530.2	S2 & S3	0031	Chr9	36883092	36884669	1578

Out NVD 10			On-1 112 2 8-000				
QrobMYB10 2	Qrob_T0612260.2	S14	Qrob_H2.3_Sc000 0097	Chr9	48108561	48107270	1292
QrobMYB10 3	Qrob_T0381310.2	SAtMYB27	Qrob_H2.3_Sc000 0278	Chr10	8229312	8230354	1043
QrobMYB10 4	Qrob_T0451690.2	S13	Qrob_H2.3_Sc000 0669	Chr10	10999232	11000574	1343
QrobMYB10 5	Qrob_T0452180.2	S 1	Qrob_H2.3_Sc000 0085	Chr10	14501122	14502632	1511
QrobMYB10 6	Qrob_T0555330.2	no subgroup	Qrob_H2.3_Sc000 0085	Chr10	14962232	14966952	4721
QrobMYB10 7	Qrob_T0555340.2	no subgroup	Qrob_H2.3_Sc000 0085	Chr10	14980766	14983349	2584
QrobMYB10 8	Qrob_T0179590.2	SAtMYB26	Qrob_H2.3_Sc000 0951	Chr10	17804921	17803483	1439
QrobMYB10 9	Qrob_T0286880.2	S2 & S3	Qrob_H2.3_Sc000 0009	Chr10	18999043	18995821	3223
QrobMYB11 0	Qrob_T0361180.2	S20	Qrob_H2.3_Sc000 0450 Outh_H2.3_Sc000	Chr10	24976888	24978282	1395
QrobMYB11 1	Qrob_T0352280.2	WPS-III	Qrob_H2.3_Sc000 0347	Chr10	27861855	27860633	1223
2 Orah MVD 11	Qrob_T0439830.2	WPS-III	Qrob_H2.3_Sc000 0347	Chr10	27882273	27881283	991
QrobMYB11 3	Qrob_T0309480.2	S21	QF0B_H2.3_SC000 0263	Chr10	33715578	33714210	1369
QrobMYB11 4	Qrob_T0416480.2	S21	Qrob_H2.3_Sc000 0394	Chr10	40481766	40480595	1172
QrobMYB11 5	Qrob_T0189820.2	S5	Qrob_H2.3_Sc000 0253	Chr11	588966	587658	1309
QrobMYB11 6	Qrob_T0189800.2	S5	Qrob_H2.3_Sc000 0253	Chr11	618203	616862	1342
QrobMYB11 7	Qrob_T0189780.2	S5	Qrob_H2.3_Sc000 0253	Chr11	640279	638759	1521
QrobMYB11 8	Qrob_T0275930.2	WPS-III	Qrob_H2.3_Sc000 0230	Chr11	3182332	3181117	1216
9 OrehMVD12	Qrob_T0409250.2	SAtMYB26	0166 Orah 112.3_Sc000	Chr11	10466232	10465025	1208
0 OrobMVP12	Qrob_T0203780.2	S22	0089 Orah H2 3 Sa000	Chr11	23625175	23624288	888
1 OrobMVP12	Qrob_T0011720.2	S4	Q100_H2.3_Sc000 0005 Orab_H2.3_Sc000	Chr11	35738109	35739384	1276
2 OrobMVB12	Qrob_T0291490.2	S13	0266 Orob H2 3 Sc000	Chr11	43032216	43033889	1674
3 OrobMVB12	Qrob_T0387070.2	S4	0389 Orob H2 3 Sc000	Chr11	45113992	45112970	1023
4 OrobMYB12	Qrob_T0099630.2	S5	0037 Orob H2 3 Sc000	Chr11	51663438	51665216	1779
5 OrobMVB12	Qrob_T0302600.2	S10 & S24	0235 Orob H2 3 Sc000	Chr12	6308619	6309765	1147
6 OrobMVB12	Qrob_T0541420.2	SAtMYB85	0023 0rob H2 3 Sc000	Chr12	9014270	9012796	1475
7 OrobMVB12	Qrob_T0082290.2	S20	0412 Orob H2 3 Sc000	Chr12	16016899	16018134	1236
8 OrobMYB12	Qrob_T0115630.2	SAtM5	0146 Orob H2 3 Sc000	Chr12	25663954	25662876	1079
9 OrobMYB13	Qrob_T0388890.2	SAtM103	0652 Orob H2 3 Sc000	Chr12	30990081	30992496	2416
0 OrobMYB13	Qrob_T0026160.2	S20	0044 Orob H2 3 Sc000	UA	1660115	1661281	1167
1 OrobMYB13	Qrob_T0468060.2	S22	0161 Orob H2.3 Sc000	UA	1159605	1158892	714
2 OrobMYB13	Qrob_T0675440.2	SAtMYB26	0165 Orob H2.3 Sc000	UA	1378120	1376578	1543
3 OrobMYB13	Qrob_T0675450.2	SAtMYB26	0165 Orob H2.3 Sc000	UA	1418714	1417338	1377
4 OrobMYB13	Qrob_T0122500.2	S18	0234 Orob H2.3 Sc000	UA	176552	171484	5069
5 OrobMYB13	Qrob_T0411820.2	SAtM5	0354 Orob H2.3 Sc000	UA	342530	343605	1076
6 QrobMYB13	Qrob_T0412190.2	WPS-IV	0358 Qrob_H2.3 Sc000	UA	390490	388344	2147
7 QrobMYB13	Qrob_T0728290.2	S13	0840 Qrob_H2.3_Sc000	UA	172040	170522	1519
8 QrobMYB13	Qrob_T0653500.2	SAtMYB26	0945 Qrob_H2.3 Sc000	UA	11717	13539	1823
9	Qrob_T0769430.2	WPS-III	1151	UA	32240	33455	1216

QrobMYB3R			Qrob_H2.3_Sc000				
-A	Qrob_T0010150.2	MYB-3R	0013	Chr2	60246757	60253755	6999
QrobMYB3R			Qrob_H2.3_Sc000				
-B	Qrob_T0576750.2	MYB-3R	0478	Chr7	51811899	51816721	4823
QrobMYB3R			Qrob_H2.3_Sc000				
-C	Qrob_T0033120.2	MYB-3R	0007	Chr8	20517700	20521806	4107
QrobMYB3R			Qrob_H2.3_Sc000				
-D	Qrob_T0129360.2	MYB-3R	0474	Chr12	14949147	14953664	4518
QrobMYB3R			Qrob_H2.3_Sc000				
-E	Qrob_T0264880.2	MYB-3R	0046	Chr12	22045628	22052297	6670
QrobMYB4R			Qrob_H2.3_Sc000				
-A	Qrob_T0439780.2	MYB-4R	0347	Chr10	28525076	28534063	8988

Supplementary Table 35 Gene content of the glutaredoxin, thioredoxin and glutathione transferase families in the pedunculate oak genome (haplome assembly) and selected embryophytes. In addition to oak sequences, other sequences were retrieved from genomic data available from thr Phytozome V11 portal by BLAST-p and tBLAST-n analyses using *P. trichocarpa* and *A. thaliana* sequences as references. The different classes in the grx, trx and gst families were defined according to^{83–85}, respectively.

	Q. robur	P. trichocarpa	A. thaliana	V. vinifera	P. persica	C. clementina	F. vesca	E. grandis	R. communis	C. papaya	T. cacao	O. sativa	S. bicolor	P. patens
GLUTAREDOXINS	25	38	33	25	24	23	24	32	22	18	17	29	32	15
Class I	5	6	6	5	5	5	4	7	6	4	5	5	5	5
C1	1	2	1	1	1	1	1	2	1	0	1	0	0	0
C2	1	1	1	1	1	1	1	2	1	1	1	2	2	3
C3	1	1	1	1	1	1	1	1	1	1	1	1	1	1
C4	1	1	1	1	1	1	1	1	1	1	1	1	1	0
C5	0	0	1	0	0	0	0	0	0	0	0	0	0	0
S12	1	1	1	1	1	1	0	1	2	1	1	1	1	1
Class II	4	5	4	5	5	4	3	5	4	3	4	5	6	8
S14	1	1	1	2	1	1	1	1	1	0	1	1	1	2
\$15	1	1	1	1	2	1	0	2	1	1	1	2	2	2
S16	1	1	1	1	1	1	1	1	1	1	1	1	2	1
S17	1	2	1	1	1	1	1	1	1	1	1	1	1	3
Class III	14	24	21	13	12	12	15	18	10	9	11	17	19	2
Class IV	2	3	2	2	2	2	2	2	2	2	2	2	2	0
THIOREDOXINS	41	49	41	35	39	31	34	48	31	30	36	34	33	34
CDSP32	1	1	1	1	1	1	1	1	1	1	1	1	1	1
Clot	1	1	1	1	1	1	1	1	1	0	1	1	1	1
HCF164	1	1	1	1	1	1	1	1	1	1	1	1	1	1
Trx f	1	1	2	1	1	1	1	2	1	1	1	1	1	3
Trx h	6	10	11	6	8	6	8	10	6	8	8	7	6	5
Trx m	4	8	4	3	5	3	4	4	4	3	4	4	3	6
Trx o	1	1	2	1	1	1	1	1	1	1	1	1	1	1
Trx x	1	1	1	1	1	1	1	1	0	1	1	1	1	2

Trx y	1	2	2	1	2	1	1	2	1	1	1	1	1	1
Trx z	1	1	1	1	1	1	1	1	1	1	1	1	1	2
Trx like 1	1	1	1	1	1	1	1	1	1	0	1	1	1	1
Trx like 2	2	3	2	2	2	2	2	2	1	1	2	1	2	2
Trx lilium 1	2	2	3	2	2	2	3	3	2	2	2	2	3	0
Trx lilium 2	1	2	1	1	1	1	1	2	1	1	2	1	1	1
Trx lilium 3	1	1	1	1	1	1	1	1	1	1	1	1	1	1
TDX	1	2	1	1	1	0	1	1	1	0	1	1	0	0
NRX1	5	5	1	5	4	2	0	7	2	3	1	2	2	0
NRX2	1	1	0	1	1	1	1	2	1	1	1	1	1	0
NRX3	1	1	1	1	1	1	1	1	0	1	1	1	1	0
NTRa/b	6	2	2	1	1	1	1	2	2	1	2	2	2	2
NTRc	1	1	1	1	1	1	1	1	1	1	1	1	1	2
FTR-b	1	1	1	1	1	1	1	1	1	0	1	1	1	2
GLUTATHIONE TRANSFERASES	88	83	61	59	71	68	50	110	52	36	60	78	90	39
DHAR	1	3	3	2	2	2	2	3	3	2	2	2	3	3
GHR	2	2	4	1	2	2	2	3	2	0	3	2	2	2
GSTL	2	3	3	4	2	3	3	8	3	2	2	3	4	1
mPGES2	2	3	1	2	2	2	2	3	1	1	2	1	1	2
GSTI	0	0	0	0	0	0	0	0	0	0	0	0	0	1
GSTH	0	0	0	0	0	0	0	0	0	0	0	0	0	7
GSTF	12	8	13	8	9	8	5	19	4	5	9	16	17	9
GSTT	1	2	3	1	1	2	1	1	3	1	1	1	2	2
GSTZ	2	2	2	3	2	3	2	2	2	1	2	4	4	1
ΕF1Βγ	2	3	2	?	2	2	3	1	1	1	1	2	2	3
Ure2p	0	0	0	0	0	0	0	0	0	0	0	0	0	1
Metaxin	1	2	1	1	1	1	1	1	1	1	1	1	1	2
GSTU	62	54	28	36	47	42	28	62	31	21	36	45	53	0
TCHQD	1	1	1	1	1	1	1	7	1	1	1	1	1	5

Supplementary Table 36 List of MLO genes identified in the pedunculate oak genome (haplome assembly). MLO predicted proteins were retrieved by three different approaches: those automatically annotated as MLO proteins, those containing the MLO domain with the Pfam signature PF03094, and those with homology to one of the *Arabidopsis* MLO proteins after BLAST-p search using an e-value of 10e⁻¹⁰ as the threshold. The predicted proteins identified were inspected and manually curated. The number of predicted transmembrane domains was analyzed with Phobius (http://phobius.sbc.su.se/).

2000

MLO gene ID	Complete/partial	MLO	Chrom.	#residues	exons	#Trans
	protein	Clade				membrane
	-					domains
Qrob_T0355750.2	complete	VI	1	561	15	7
Qrob_T0355780.2	complete	V	1	584	15	7
Qrob_T0355790.2	complete	V	1	585	15	7
Qrob_T0173130.2	complete	Ι	2	519	13	7
Qrob_T0222290.2	complete	II	2	510	15	7
Qrob_T0482700.2	complete	III	2	522	15	7
Qrob_T0725960.2	partial		2			
Qrob_T0725970.2	partial		2			
Qrob_T0562700.2	complete	II	5	504	15	8
Qrob_T0346330.2	complete	Ι	6	564	15	7
Qrob_T0603780.2	complete	III	6	573	15	7
Qrob_T0327490.2	partial		7			
Qrob_T0455210.2	complete	V	8	565	15	7
Qrob_T0468620.2	complete	Ι	8	558	15	7
Qrob_T0572110.2	complete	V	8	539	14	7
Qrob_T0572120.2	complete	V	8	533	14	7
Qrob_T0572170.2	complete	V	8	575	15	7
Qrob_T0032420.2	complete	II	10	516	14	7
Qrob_T0032520.2	complete	II	10	520	14	7
Qrob_T0032530.2	complete	II	10	521	14	7
Qrob_T0254270.2	complete	V	10	557	15	7
Qrob_T0254250.2	partial		10			
Qrob_T0254260.2	partial		10			
Qrob_T0032510.2	partial		10			
Qrob_T0254290.2	partial		11			
Qrob_T0523040.2	complete	II	Scaf. 408	505	15	7

Supplementary Table 37 Mildew resistance locus o (MLO) family members from selected plant species and their phylogenetic classification^a. Clade V (in bold) corresponds to the clade for which a function in powdery mildew susceptibility/resistance has been demonstrated. We completed the table provided by Acevedo-Garcia et al.⁹¹ with recently published data and data obtained from the pedunculate oak genome (haplome assembly).

2008

Scientific	Common	#MLO			Cla	de ID				
name	name	genes								
			Ι	II	III	IV	V	VI	VII	Reference*
Arabidopsis thaliana	Thale cress	15	3	3	5	0	3	1	0	161
Cucumis lanatus	Cucumber	14	4	1	3	0	3	1	2	162
Cucumis sativus	Cucumber	14	4	2	3	0	3	1	1	163
Fragaria vesca	Strawberry	18	3	6	1	1	3	2	1	92
Glycine max	Soybean	39 °	7	5	8	2	11	6	0	164
Gossypium hirsutum	Cotton	38 °	8	11	3	2	8	2	4	165
Malus domestica	Apple	21 ^c	5	5	3	0	4	2	1	92
Oryza sativa	Rice	12	2	6	1	3	0	0	0	166
Prunus persica	Peach	16	3	6	2	0	3	2	0	167
Prunus persica	Peach	19	3	5	2	1	3	3	1	92
Solanum lycopersicum	Tomato	17 ^b	3	4	3	0	4	1	1	168
Solanum tuberosum	Potato	13	3	4	3	0	3	0	0	169
Triticum aestivum	Wheat	8	1	3	1	3	0	0	0	170
Vitis vinifera	Grapevine	17	3	3	2	1	6	2	0	171
Quercus robur	Oak	19	3	6	2	0	7	1	0	this study

2009 ^a Only fully characterized MLO families are shown. Classification is based on previous publications.

2010 ^b One truncated MLO family member (SIMLO14) was excluded from phylogenetic analysis

2011 ^c Species with recent whole-genome duplication

2012 * from Acevedo-Garcia et al.⁹¹ until 2014

2013

Supplementary Table 38 Annotation of the *Populus trichocarpa* laccase genes. Poplar laccase gene annotations were updated according to the most recent annotation (v3) available in Phytozome. Synonyms of the gene annotations used in this study are presented, together with previous annotations based on Phytozome v2 annotation.

Annotation ¹³²	Annotation ¹²⁹	Synonym (Annotation v2)	P trichocarpa Alias	Gene name (v3)	Transcript name (v3)
PtrLAC1	PtrLAC1	POPTR_0001s14010	PtrLAC1	Potri.001G054600	Potri.001G054600.1
PtrLAC2	PtrLAC2	POPTR_0001s18500	PtrLAC2	Potri.001G184300	Potri.001G184300.1
PtrLAC3	PtrLAC3	POPTR_0001s21380	PtrLAC3	Potri.001G206200	Potri.001G206200.1
PtrLAC4	PtrLAC4	POPTR_0001s25580	PtrLAC4	Potri.001G248700	Potri.001G248700.1
PtrLAC5	PtrLAC5	POPTR_0001s35740	PtrLAC5	Potri.001G341600	Potri.001G341600.1
PtrLAC6	PtrLAC6	POPTR_0001s41160	PtrLAC6	Potri.001G401100	Potri.001G401100.1
PtrLAC7	PtrLAC7	POPTR_0001s41170	PtrLAC7	Potri.001G401300	Potri.001G401300.1
PtrLAC8	PtrLAC8	POPTR_0004s16370	PtrLAC8	Potri.004G156400	Potri.004G156400.1
	PtrLAC9	POPTR_0005s22230	PtrLAC9	Potri.005G200500	Potri.005G200500.1
PtrLAC9	PtrLAC10	POPTR_0005s22240	PtrLAC10	Potri.005G200600	Potri.005G200600.1
PtrLAC10	PtrLAC11	POPTR_0005s22250	PtrLAC11	Potri.005G200700	Potri.005G200700.1
PtrLAC11	PtrLAC12	POPTR_0006s08740	PtrLAC12	Potri.006G087100	Potri.006G087100.1
PtrLAC12	PtrLAC13	POPTR_0006s08780	PtrLAC13	Potri.006G087500	Potri.006G087500.1
PtrLAC13	PtrLAC14	POPTR_0006s09520	PtrLAC14	Potri.006G094100	Potri.006G094100.1
PtrLAC14	PtrLAC15	POPTR_0006s09830	PtrLAC15	Potri.006G096900	Potri.006G096900.1
PtrLAC15	PtrLAC16	POPTR_0006s09840	PtrLAC16	Potri.006G097000	Potri.006G097000.1
PtrLAC49	PtrLAC51	POPTR_0958s00200	PtrLAC17	Potri.006G097100	Potri.006G097100.1

PtrLAC16	PtrLAC17	POPTR_0007s13050	PtrLAC18	Potri.007G023300	Potri.007G023300.1
PtrLAC17	PtrLAC18	POPTR_0008s06430	PtrLAC19	Potri.008G064000	Potri.008G064000.1
PtrLAC18	PtrLAC19	POPTR_0008s07370	PtrLAC20	Potri.008G073700	Potri.008G073700.1
PtrLAC19	PtrLAC20	POPTR_0008s07380	PtrLAC21	Potri.008G073800	Potri.008G073800.1
PtrLAC20	PtrLAC21	POPTR_0009s03940	PtrLAC22	Potri.009G034500	Potri.009G034500.1
PtrLAC21	PtrLAC22	POPTR_0009s04720	PtrLAC23	Potri.009G042500	Potri.009G042500.1
PtrLAC22	PtrLAC23	POPTR_0009s10550	PtrLAC24	Potri.009G102700	Potri.009G102700.1
PtrLAC23	PtrLAC24	POPTR_0009s15840	PtrLAC25	Potri.009G156600	Potri.009G156600.1
PtrLAC24	PtrLAC25	POPTR_0009s15860	PtrLAC26	Potri.009G156800	Potri.009G156800.1
PtrLAC25	PtrLAC26	POPTR_0010s19080	PtrLAC27	Potri.010G183500	Potri.010G183500.1
PtrLAC26	PtrLAC27	POPTR_0010s19090	PtrLAC28	Potri.010G183600	Potri.010G183600.1
PtrLAC27	PtrLAC28	POPTR_0010s20050	PtrLAC29	Potri.010G193100	Potri.010G193100.1
PtrLAC28	PtrLAC29	POPTR_0011s06880	PtrLAC30	Potri.011G071100	Potri.011G071100.1
PtrLAC29	PtrLAC30	POPTR_0011s12090	PtrLAC31	Potri.011G120200	Potri.011G120200.1
PtrLAC30	PtrLAC31	POPTR_0011s12100	PtrLAC32	Potri.011G120300	Potri.011G120300.1
PtrLAC31	PtrLAC32	POPTR_0012s04620	PtrLAC33	Potri.012G048900	Potri.012G048900.1
PtrLAC32	PtrLAC33	POPTR_0013s14890	PtrLAC34	Potri.013G152700	Potri.013G152700.1
PtrLAC33	PtrLAC34	POPTR_0014s09610	PtrLAC35	Potri.014G100600	Potri.014G100600.1
PtrLAC36	PtrLAC38	POPTR_0015s04370	PtrLAC36	Potri.015G040400	Potri.015G040400.1
PtrLAC35	PtrLAC37	POPTR_0015s04350	PtrLAC37	Potri.015G040600	Potri.015G040600.1
PtrLAC34	PtrLAC36	POPTR_0015s04340	PtrLAC38	Potri.015G040700	Potri.015G040700.1
	PtrLAC35	POPTR_0015s04330	PtrLAC39	Potri.015G040800	Potri.015G040800.1
PtrLAC37	PtrLAC39	POPTR_0016s11500	PtrLAC40	Potri.016G106000	Potri.016G106000.1

PtrLAC38	PtrLAC40	POPTR_0016s11520	PtrLAC41	Potri.016G106100	Potri.016G106100.1
PtrLAC39	PtrLAC41	POPTR_0016s11540	PtrLAC42	Potri.016G106300	Potri.016G106300.1
PtrLAC48	PtrLAC50	POPTR_0091s00270	PtrLAC43	Potri.016G107900	Potri.016G107900.1
PtrLAC40	PtrLAC42	POPTR_0016s11950	PtrLAC44	Potri.016G112000	Potri.016G112000.1
PtrLAC41	PtrLAC43	POPTR_0016s11960	PtrLAC45	Potri.016G112100	Potri.016G112100.1
PtrLAC42	PtrLAC44	POPTR_0019s11810	PtrLAC46	Potri.019G088500	Potri.019G088500.1
PtrLAC43	PtrLAC45	POPTR_0019s11820	PtrLAC47	Potri.019G088600	Potri.019G088600.1
PtrLAC44	PtrLAC46	POPTR_0019s11830	PtrLAC48	Potri.019G088700	Potri.019G088700.1
PtrLAC45	PtrLAC47	POPTR_0019s11850	PtrLAC49	Potri.019G088800	Potri.019G088800.1
PtrLAC46	PtrLAC48	POPTR_0019s11860	PtrLAC50	Potri.019G088900	Potri.019G088900.1
PtrLAC47	PtrLAC49	POPTR_0019s14530	PtrLAC51	Potri.019G121700	Potri.019G121700.1

2018Supplementary Table 39 Number of laccase genes per phylogenetic group for2019Arabidopsis, poplar and oak.

	A. thaliana	Q. robur	P. trichocarpa
Group 1	1	1	5
Group 2	6	14	24
Group 3	1	1	1
Group 4	3	1	5
Group 5	4	1	6
Group 6	1	7	6
Group 7	1	2	4
Total	17	27	51

2023Supplementary Table 40 Available or soon to be released eudicot whole-genome2024sequences(asreportedin2025https://genomevolution.org/wiki/index.php/Sequenced_plant_genomes as of 18 December20262016), with growth habit from Zanne et al.¹⁵² (H = herbaceous, W = woody), supplemented2027with Google searches, and % woody based on the strong prior from Fitzjohn et al.¹⁴⁹ at the2028genus, family and order levels. The names of the species were updated with taxize R-package,2029using Plant List version 1.1.

				% woody		
Order	Family	Sequenced species	Growth form	Genus	Family	Order
Brassicales	Brassicaceae	Arabidopsis halleri	Н	0	5	15
Brassicales	Brassicaceae	Arabidopsis lyrata	Н	0	5	15
Brassicales	Brassicaceae	Arabidopsis thaliana	Н	0	5	15
Fabales	Fabaceae	Arachis hypogaea	Н	0	63	62
Caryophyllales	Amaranthaceae	Beta vulgaris	Н	0	38	42
Brassicales	Brassicaceae	Brassica napus	Н	0	5	15
Brassicales	Brassicaceae	Brassica oleracea	Н	0	5	15
Brassicales	Brassicaceae	Brassica rapa	Н	0	5	15
Brassicales	Brassicaceae	Camelina sativa	Н	0	5	15
Rosales	Cannabaceae	Cannabis sativa	Н	0	96	75
Solanales	Solanaceae	Capsicum annuum	Н	0	62	50
Fabales	Fabaceae	Cicer arietinum	Н	0	63	62
Cucurbitales	Cucurbitaceae	Citrullus lanatus	Н	0	13	11
Cucurbitales	Cucurbitaceae	Cucumis melo	Н	0	13	11
Cucurbitales	Cucurbitaceae	Cucumis sativus	Н	0	13	11
Asterales	Asteraceae	Erigeron canadensis	Н	8	25	26
Rosales	Rosaceae	Fragaria vesca	Н	0	76	75
Fabales	Fabaceae	Glycine max	Н	13	63	62
Rosales	Cannabaceae	Humulus lupulus	Н	0	96	75
Fabales	Fabaceae	Lotus corniculatus	Н	25	63	62
Fabales	Fabaceae	Lupinus angustifolius	Н	22	63	62
Fabales	Fabaceae	Medicago truncatula	Н	8	63	62
Lamiales	Phrymaceae	Mimulus guttatus	Н	27	24	45
Proteales	Nelumbonaceae	Nelumbo nucifera	Н	0	0	100
Solanales	Solanaceae	Nicotiana benthamian	a H	33	62	50
Fabales	Fabaceae	Phaseolus vulgaris	Н	0	63	62
Brassicales	Brassicaceae	Raphanus raphanistru	m H	0	5	15
Brassicales	Brassicaceae	Sisymbrium irio	Н	0	5	15

Solanales	Solanaceae	Solanum lycopersicum	Н	65	62	50
Solanales	Solanaceae	Solanum melongena	Н	65	62	50
Solanales	Solanaceae	Solanum tuberosum	Н	65	62	50
Lamiales	Lentibulariaceae	Utricularia gibba	Н	0	3	45
Brassicales	Brassicaceae	Aethionema arabicum	Н	0	5	15
Ranunculales	Ranunculaceae	Aquilegia formosa	Н	0	16	31
Brassicales	Brassicaceae	Capsella rubella	Н	0	5	15
Sapindales	Rutaceae	Citrus clementina	W	100	100	100
Brassicales	Cleomaceae	Cleome houtteana	Н	15	15	15
Brassicales	Brassicaceae	Eutrema parvulum	Н	0	5	15
Brassicales	Brassicaceae	Eutrema salsugineum	Н	0	5	15
Lamiales	Lentibulariaceae	Genlisea aurea	Н	47	3	45
Brassicales	Brassicaceae	Leavenworthia alabamica	Н	0	5	15
Malpighiales	Linaceae	Linum usitatissimum	Н	33	54	79
Solanales	Solanaceae	Lycopersicon pennellii	Н	0	62	50
Rosales	Rosaceae	Prunus mume	W	100	76	75
Rosales	Rosaceae	Pyrus bretschneideri	W	100	76	75
Solanales	Solanaceae	Solanum arcanum	Н	65	62	50
Solanales	Solanaceae	Solanum habrochaites	Н	65	62	50
Solanales	Solanaceae	Solanum pimpinellifolium	Н	65	62	50
Fabales	Fabaceae	Vigna radiata	Н	0	63	62
Ericales	Actinidiaceae	Actinidia chinensis	W	100	100	84
Malvales	Thymelaeaceae	Aquilaria agallocha	W	100	94	86
Sapindales	Meliaceae	Azadirachta indica	W	100	100	100
Fagales	Betulaceae	Betula nana	W	100	100	100
Fabales	Fabaceae	Cajanus cajan	W	100	63	62
Brassicales	Caricaceae	Carica papaya	W	100	93	15
Sapindales	Rutaceae	Citrus sinensis	W	100	100	100
Gentianales	Rubiaceae	Coffea canephora	W	100	82	76
Myrtales	Myrtaceae	Eucalyptus grandis	W	100	100	92
Lamiales	Oleaceae	Fraxinus excelsior	W	100	99	45
Malvales	Malvaceae	Gossypium raimondii	W	93	83	86
Malpighiales	Euphorbiaceae	Hevea brasiliensis	W	100	72	79
Rosales	Rosaceae	Malus domestica	W	100	76	75
Malpighiales	Euphorbiaceae	Manihot esculenta	W	100	72	79

Malpighiales	Salicaceae	Populus trichocarpa	W	100	100	79
Rosales	Rosaceae	Prunus persica	W	100	76	75
Fagales	Fagaceae	Quercus robur	W	100	100	100
Malpighiales	Euphorbiaceae	Ricinus communis	W	100	72	79
Malpighiales	Salicaceae	Salix purpurea	W	100	100	79
Malvales	Malvaceae	Theobroma cacao	W	100	83	86
Ericales	Ericaceae	Vaccinium corymbosum	W	100	98	84
Ericales	Ericaceae	Vaccinium macrocarpon	W	100	98	84
Vitales	Vitaceae	Vitis vinifera	W	100	98	98
Rosales	Rhamnaceae	Ziziphus jujuba	W	100	99	75

Supplementary Table 41 Summary of the data used for GO term enrichment analysis
for three gene categories: TDG (tamdemly duplicated genes), LDG (long distanceduplicated genes) and SG (singleton genes). Abbreviations are as follows: MF (Molecular
Function), BP (Biological Process), CC (Cellular Component).

	Total	MF	BP	CC
No. of genes with GO terms in the reference	16,820	15,413	10,073	3,604
Total No. of GO terms	3,433	1,179	1,867	387
No. of TDGs with GO terms	6,686	6,280	4,103	1,086
No. of LDGs with GO terms	6,230	5,680	3,844	1,536
No. of SGs with GO terms	3,904	3,453	2,126	982
Significant GO terms: TDGs	97	55	32	10
Significant GO terms: LDGs	144	65	62	17
Significant GO terms: SGs	240	80	130	30

Supplementary Table 42 Summary of the data used for GO term enrichment analysis in the orthogroups expanded in pedunculate oak.

	Total	MF	BP	CC
No. of genes with GO terms in the reference	16,820	15,413	10,073	3,604
No. of genes with GO terms in orthogroups expanded in oak	4,217	4,032	2,267	445
Total No. of GO terms in orthogroups expanded in oak	3,433	1,179	1,867	387
Significant GO terms in orthogroups expanded in oak	58	33	17	8

Supplementary Table 43 Summary of gene ontology (GO) enrichment analysis in woody perennials (A) and herbaceous species (B). Abbreviations are as follows: Molecular function (MF), biological process (BP) and cellular component (CC).

A. Woody perennials				
	Total	MF	BP	CC
No. of orthogroups with GO terms in the reference	36,844	16,703	11,495	5,073
Total number of GO terms in the reference	3,936	1,341	2,131	464
No. of significant expanded orthogroups with GO terms	108	104	84	39
No. of significant GO terms in expanded orthogroups	61	38	19	4
B. Herbaceous species				
	Total	MF	BP	CC
No. of orthogroups with GO terms in the reference	36,844	16,703	11,495	5,073
Total No. of GO terms in the reference	3,936	1,341	2,131	464
No. of significant expanded orthogroups with GO terms	23	16	12	4
No. of significant GO terms in expanded orthogroups	7	5	2	0

2051 11.Supplementary Figures

Supplementary Fig. 1 Genome coverage distribution of the V1 (diploid, black) and V2
(diploid, red) assemblies, showing fewer regions with twice the expected coverage in the V2
assembly, i.e. better resolved haplotypes in the V2 assembly.



Supplementary Fig. 2 Illustration of the syntenome approach between *Prunus persica* and
 Quercus robur. Pedunculate oak scaffolds are anchored by oak markers (green scaffolds
 matching red dots of the oak genetic linkage map) or by peach gene models (purple scaffolds)
 or by both (combined green-purple scaffolds).



Supplementary Fig. 3 Distribution of TE families. Distribution of TE families according to:
(a) their main order or superfamily (Gypsy/Copia) in the consensus library (1,750 consensus)
and (b) their genome coverage (716,192 copies, 52% of the genome).





Supplementary Fig. 4 Chromosomal variations of genetic diversity. (A) Proportion of heterozygous sites within the "3P" reference genome sequence. (B) estimation of Tajima's π at the population level. Both metrics were calculated in 1 Mb windows, sliding by steps of 250 kb. Colors correspond to Tukey's Honestly Significant Difference criterion at the α = 0.05 significance level. Box plots were drawn with R with default parameters. The bottom and top of the box are the 25th and 75th percentile (the lower and upper quartiles, respectively), thus delineating the interquartile range (IQR). The band near the middle of the box is the 50th percentile (i.e. the median). For the ends of the whiskers we used the default box plot parameter for statistical dispersion in R (1.5*IQR). Below the figures the sample size (number of windows) and quantile values are provided for each chromosome and each metric.



2089 Sample sizes and qualile values.

${f A}$ chrom	#windows	1 stcentile	5thcentile	10thcentile	1 stquartile	median	3rdquartile	90thcentile	95thcentile	99thcentile
1 Chr01	130	0.002363013	0.004393948	0.005261262	0.007251443	3 0.008672124	0.010039868	0.011519258	3 0.012145932	0.016840459
2 Chr02	266	0.002100697	0.004298015	0.005754484	0.007370706	5 0.008988398	3 0.010277284	0.011669246	5 0.012467453	0.014217985
3 Chr03	137	0.001081677	0.002477953	0.004194525	0.007676995	5 0.009552104	0.011155354	0.012308122	2 0.012993091	0.013777254
4 Chr04	99	0.001895541	0.004651434	0.006457263	0.009059405	5 0.011011622	2 0.012866448	0.013945719	0.014408945	0.015519873
5 Chr05	160	0.002897904	0.004659509	0.005936392	0.00798619	0.009694981	0.011206569	0.012567117	0.013375598	0.015656735
6 Chr06	131	0.001494684	0.002640328	0.003820639	0.006321167	0.009001377	0.010412181	0.011392821	0.01235098	0.013901824
7 Chr07	107	0.002069173	0.004260246	0.005199583	0.007680156	5 0.009023406	5 0.0102929	0.011745387	0.012485407	0.014034843
8 Chr08	177	0.002690201	0.004800595	0.006018946	0.007705742	2 0.009324316	5 0.010612334	0.011557058	3 0.012273925	0.014481952
9 Chr09	115	0.001977248	0.00323089	0.004647822	0.007088226	5 0.009489949	0.011178917	0.012862547	0.014187532	0.015566817
10 Chr10	119	0.001263906	5 0.003383373	0.004767478	0.007632744	0.00916589	0.010998188	0.012691446	5 0.013874577	0.014762114
11 Chr11	108	0.002044077	0.004747339	0.006210409	0.008371558	3 0.010282012	2 0.011518549	0.012360799	0.013290434	0.014556657
12 Chr12	90	0.002745973	0.003568476	0.005181146	0.007615457	7 0.009013432	2 0.01060477	0.011471683	8 0.011952998	0.012647347

${f B}$ chrom	#windows	1stcentile	5thcentile	10thcentile	1 stquartile	median	3rdquartile	90thcentile	95thcentile	99thcentile
1 Chr01	145	0.006036747	0.007824487	0.008626952	0.0098874	4 0.010948342	2 0.011802318	3 0.01257944	1 0.013850033	0.015472398
2 Chr02	315	0.007005252	0.008245111	0.00893641	0.009946993	3 0.01079225	5 0.011729033	3 0.013046817	7 0.013603244	0.014903649
3 Chr03	144	0.008221722	0.008864512	0.00936807	0.010093188	3 0.011332564	4 0.01192446	7 0.013027595	5 0.013648772	0.014547199
4 Chr04	119	0.007066025	0.009430628	0.009896027	0.011095004	4 0.012225594	4 0.01336722	0.01396720	1 0.014483707	0.015332212
5 Chr05	180	0.008182576	0.008781872	0.009904223	0.010659378	8 0.011487088	8 0.01247425	5 0.01338964	7 0.013723549	0.015578464
6 Chr06	165	0.005597463	0.007781344	0.008438614	0.009589428	8 0.010516228	8 0.01122136	5 0.012670056	5 0.013657986	0.015023686
7 Chr07	126	0.007055785	0.008025906	0.009054159	0.00982444	4 0.010753628	8 0.011695085	5 0.012823905	5 0.013157198	0.015324665
8 Chr08	180	0.00576823	0.008757703	0.009330757	0.010481886	5 0.011365739	9 0.01227747	7 0.013089583	3 0.013717592	0.015245776
9 Chr09	139	0.005964687	0.007555009	0.008606873	0.009947773	3 0.011206606	6 0.01206739	0.012986125	5 0.0136571	0.014526484
10 Chr10	119	0.008395567	0.009141206	0.009512838	0.010143959	0.011079144	4 0.012335854	4 0.013440607	7 0.014436443	0.016871645
11 Chr11	135	0.007864851	0.008833194	0.009467378	0.010663593	3 0.011324205	5 0.012517949	0.013841680	5 0.014949459	0.016224063
12 Chr12	101	0.005913755	0.007754588	0.008757966	0.009844794	4 0.010519347	7 0.011761386	5 0.012460929	9 0.012820039	0.014536021

2094 Supplementary Fig. 5 Dating of branch insertion and bud sampling for DNA extraction. (a)
2095 Coring of a lateral branch at its insertion into the main trunk. (b) Wood core used to estimate
2096 the age of the lateral branch. (c) Bud sampling at the extremity of a lateral branch.



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2102 Supplementary Fig. 6 Maximum likelihood phylogenetic analysis of TIR-NB-LRR-related genes. The NB domain of TNL-related genes (i.e., TNL, TNLX, NL, NLX, TN, TNX, TL, N) 2103 2104 corresponding to orthogroup #1000 was used to study the relationship betwen selected tree 2105 (Cc, Citrus clementina; Cp, Carica papaya; Eg, Eucalyptus grandis; Md, Malus domestica; 2106 Pp, Prunus persica; Pt, Populus trichocarpa; Qr, Quercus robur; Tc, Theobroma cacao; Vv, 2107 *Vitis vinifera*) and herbaceous plant species (At, Arabidopsis thaliana; Al, Arabidopsis lyrata; 2108 Cl, Citrullus lanatus; Fv, Fragaria vesca; Gm, Glycine max; Rc, Ricinus communis; St, 2109 Solanum tuberosum), represented by brown and green branches, respectively. The oak TNL-2110 related genes clades are shown as light brown squares. The red arrows in these blocks 2111 correspond to physical clusters along the genome, and chromosomes (Chr_x) or scaffolds (Scaf x) are indicated (only the major clusters are indicated). The number of genes from 2112 2113 different species falling into the major clades are shown (only the dominant species were 2114 counted for each clade). Bootstrap values over 70% and 80% (of 1000 replicates) are 2115 indicated by gray and black dots, respectively. Supported terminal nodes are not shown, to make the tree easier to read. The NB domain of APAF-1 was used as an outgroup to root the 2116 tree. Clades containing the NB domains of the TNL genes Rpp4, Rpp5, Rps4 reported in 2117 A. thaliana are indicated. 2118



2119

2121 **Supplementary Fig. 7** Fold-enrichment over background level (*x*-axis) of the significant 2122 gene ontology (GO) terms (P<0.01) of the orthogroups expanded in woody perennials. GO 2123 terms representing biological processes are shown as red lines, cellular components are shown 2124 in blue and molecular functions are shown in green. Sample sizes are provided in 2125 **Supplementary Data Set 8** sheet #5).

2126



2128

Supplementary Fig. 8 Number of amino acids under positive selection for each of the 24 positions of the LRR domain unit. L: Leu, x: variable, N: Asn, G: Gly, I: Ile, P: Pro.



Supplementary Fig. 9 NUCmer alignment and dotplots of Cabog (A) and Newbler (B)
scaffolds against two pedunculate oak BACs (177A20 and 107I07).



Supplementary Fig. 10 Generation of the haploid assembly (1n) of pedunculate oak with haplomerger. In general, both haplotypes are well separated in the 2n assembly (blue and orange haplotypes). For each aligned block (pink polygons), we retained only the longest sequence (haplotype blue or orange) as recommended by the creators of haplomerger, to maximize gene content. Scaffolds merging the two haplotypes (as Scaffold F) in the 2n assembly were retained, without modifications, in the 1n assembly.

2149



Supplementary Fig. 11 Alignment of two allelic BACs (#50E24 and #177A20) against the
V2 diploid assembly (scaffold #0721 and #0436) and the V2 haploid assembly (Sc00000330).
Gray boxes represent NUCmer alignments, blue rectangles correspond to SNPs specific to
BAC #177A20 and red rectangles to SNPs specific to BAC #50E24. Green boxes correspond
to flanking transposable elements.



Supplementary Fig. 12 Alignment of two allelic BACs (#12J1 and #121F17) against the V2
diploid assembly (scaffold #0397 and #01822) and the V2 haploid assembly (Sc00000226).
Gray boxes represent NUCmer alignments, blue rectangles correspond to SNPs specific to
BAC #121F17 and red rectangles to SNPs specific to BAC #12J1. Green boxes correspond to
flanking transposable elements.



- Supplementary Fig. 13 Alignment of the two allelic BACs (#107I07 and #5E10) against the
 V2 diploid assembly (scaffold #1218 and #0333) and the V2 haploid assembly (Sc00000189).
 Gray boxes represent NUCmer alignments, blue rectangles correspond to SNPs specific to
 BAC #107I07 and red rectangles to SNPs specific to BAC #5E10. Green boxes correspond to
 flanking transposable elements.



2176 Supplementary Fig. 14 Region of the Orob H2.3 Sc0000378 scaffold containing a snoRNA cluster. Intergenic regions and snoRNAs are shown on separate lines. The name of the 2177 2178 predicted snoRNA is given at the beginning of the predicted gene. For C/D box-predicted 2179 ncRNAs, the C and D motifs are colored in red, the terminal hairpin is colored in blue. For H/ACA box-predicted ncRNAs, the hairpins are colored in blue and green. The secondary 2180 2181 structure is also indicated in parentheses. The ACA box is shown in red. Pink and blue shaded 2182 motifs at the beginning of the cluster represent putative RNA pol II promoter motifs (site II 2183 and TEF, respectively).



2185 **Supplementary Fig. 15** Genome-wide proportion of heterozygous sites (warm colors) and 2186 Tajima's π (cold colors) estimates. The proportion of heterozygous sites was calculated after 2187 calling heterozygous SNPs within the "3P" reference genome sequence by remapping all 2188 Illumina reads of this genotype. Tajima's π was estimated by a whole-genome sequencing 2189 strategy based on a pool of 20 pedunculate oaks. Both metrics were calculated in 1 Mb sliding 2190 windows moving in 250 kb steps.

2191



Supplementary Fig. 16 Dotplot comparison of oak-grape-peach-cocoa genomes. Dot illustration of grape-cocoa, grape-peach and grape-oak genome comparisons. Considering grape to be the closest modern representative of the n=21 rosid ancestor (derived from a post- γ ancestor with 7 protochromosomes shown in color on the *y*-axis of the dotplots), clear relationships are observed between the grape-cocoa (1:1), grape-peach (1:1) and grape-oak (1:1) genomes (see dotplot diagonals in each chart, shown with green circles), supporting the absence of lineage-specific polyploidization events in the considered species.

2201



2204

Supplementary Fig. 17 List of 16 plant species used for gene expansion contraction analysis
 in pedunculate oak. (A) Phylogenetic tree format of the 16 species used in orthoMCL/CAFE
 software. (B) Phylogenetic tree representation of the 16 species. Red dots correspond to
 branch specific whole genome duplication events. Species initials refer to Supplementary
 Table 7.

2212 A

2213	[(St:120,(Vv:117,(((Wa:101,((Md:53,(Pp:52,Fv:52):1):47,(Gm:99,Qr:99):1):1):1,(Rc:81,Pt:81
2214):21):8,(Eg:109,(Cc:95,(Tc:87,(Cp:72,(At:5,Al:5):67):15):8):14):1):7):3)]



Supplementary Fig. 18 Distribution of the 524 orthogroups expanded in pedunculate oak across 15 plant species.



Supplementary Fig. 19 Proportion of genes classified as singleton genes (SGs, blue), tandem
 duplicated genes (TDGs, green) and long distance-duplicated genes (LDGs, red) in the 524
 significant expanded orthogroups in pedunculate oak (PO).



2232 Supplementary Fig. 20 Identification of speciation and duplication events in the pedunculate 2233 oak genome. Illustration of the K_s distribution (x-axis) of gene pairs (y-axis) observed for oak 2234 (Quercus robur)/peach (Prunus persica) orthologs (green) and for grape (blue), peach (red), 2235 cocoa (brown) and oak (purple) paralogs. The oak/peach ortholog K_s distribution defines the 2236 position of the speciation event between these two species, with a single ancestral triplication 2237 event (γ) common to grape, peach, cocoa and oak and predating the speciation event. The burst of tandem duplicates highlighted by the purple K_s peak occurred after oak/peach 2238 2239 speciation and appears to be an oak-specific event. Ks values for grape, peach and cocoa 2240 paralogous gene pairs were restricted to the γ triplication as a matter of comparison to the 2241 corresponding ancestral polyploidization event in oak.



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Supplementary Fig. 21 Dot plot representation of duplicates and extracted tandemly
duplicated genes (TDGs). Dot plot representation of the pedunculate oak genome against
itself for the complete set of paralogous pairs (left) and extracted TDGs (right).



2250 Supplementary Fig. 22 Validation of tandemly duplicated genes in pedunculate oak. (A) Distribution of the proportion of gap characters in the alignments. (B) Distribution of the 2251 2252 proportion of variable sites (SNPs) in the alignments, expressed as a ratio of the number of 2253 variable sites to total alignment length, after the exclusion of gap positions. Below each plot, a 2254 box plot shows the 2.5th, 25th, 50th, 75th and 97.5th percentiles. Summary statistics for the 2255 11,695 tandem duplicate pairs (black curve), and the 12,603 allelic pairs (light gray curve) identified by comparing the sets of genes in the diploid and haploid versions of the peduculate 2256 oak reference genome. Pairwise nucleotide sequence alignments were performed with 2257 2258 MUSCLE.



Supplementary Fig. 23 Genome coverage distribution of the longest scaffold (black) and coverage distribution of tandemly duplicated genes (red).





Supplementary Fig. 24 Box plot of the number of genes per orthoMCL cluster for each of the 16 species studied, including pedunculate oak. Species initials refer to Supplementary

 Table 7. Sample size for each species is indicated in Supplementary Table 22. A Tukey box

 plot was used. The bottom and top of the box are the 25^{th} and 75^{th} percentile (the lower and upper quartiles, respectively), thus delineating the interquartile range (IQR). The band near the middle of the box is the 50th percentile (i.e. the median). For the ends of the whiskers we used the default box plot parameter for statistical dispersion in R (1.5*IQR).



Supplementary Fig. 25 GenomeScope output generated from the 31-mers distribution. The size of the pedunculate oak haploid genome was at 736 Mb.



GenomeScope Profile len:735,931,021bp uniq:67.1% het:1.52% kcov:27.4 err:0.24% dup:0.604

Supplementary Fig. 26 Comparison of allelic BAC structures of the reference Pedunculate
oak genotype "3P". Manually curated genes are represented as green arrows with the head
indicating the direction of transcription. Repetitive elements are represented as purple boxes.
(A) 5E10_107I07, (B) 27L03_48K01, (C) 12J01_121F17, (D) 50E24_177A20, (E)
64H03_30P01.





Supplementary Fig. 27 Alignment of two allelic BACs (#50E24 and #177A20) against the
 V1 diploid assembly (scaffold #1466). Gray boxes represent nucmer alignments, blue
 rectangles correspond to SNPs specific to BAC #177A20 and red rectangles to SNPs specific
 to BAC #50E24. Green boxes correspond to flanking transposable elements.



Supplementary Fig. 28 Alignment of two allelic BACs (#12J1 and #121F17) against the V1
 diploid assembly (scaffold #3597). Gray boxes represent NUCmer alignments, blue rectangles
 correspond to SNPs specific to BAC #121F17 and red rectangles to SNPs specific to BAC
 #12J1. Green boxes correspond to flanking transposable elements.



Supplementary Fig. 29 Alignment of two allelic BACs (#107I07 and #5E10) against the V1
 diploid assembly (scaffolds #282 and #1030). Gray boxes represent NUCmer alignments, blue
 rectangles correspond to SNPs specific to BAC #107I07 and red rectangles to SNPs specific
 to BAC #5E10. Green boxes correspond to flanking transposable elements.



2312 Supplementary Fig. 30 Flow chart indicating the procedure leading to the identification of

- the 4,070 mapped markers (2,127+1,943) of the oak genome browser "marker" track.
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Supplementary Fig. 31 High-density genetic linkage map of the pedunculate oak genome
(5,589 markers) showing the map positions of the 4,070 markers aligned on the 12
chromosomes, with possible inversion tolerated within a 5 cM interval.



Supplementary Fig. 32 Physical – genetic relationships. Left panels- Physical position (in
Mb on the haplome) and genetic location (in cM on the composite linkage map) for 4,070
markers used to populate the "marker" track of the pedunculate oak genome browser.
Inversions between marker assignments on the genetic and physical maps are tolerated within
a 5 cM window. Set#1 and set#2 markers from Supplementary Data Set 3 sheet #1 are
indicated by blue and red dots, respectively. Right panels- recombination rate along the 12
chromosomes (chr 1-12).



2328 Supplementary Fig. 33 TE vs non-TE content in the 16 sequenced genomes considered in 2329 this study. The total in Mb (x-axis) corresponds to the fraction of the genome annotated. The 2330 generated with Taxonomy tree on the left was the NCBI Browser 2331 (www.ncbi.nlm.nih.gov/Taxonomy/CommonTree/wwwcmt.cgi). Only the topology is shown 2332 and the branch lengths are not proportional to evolutionary divergence time.

2333



Supplementary Fig. 34 Endogenous viruses in the pedunculate oak genome. Phylogenetic reconstruction from the multiple sequence alignment of 58 reverse transcriptase domains from representative member from endogenous *Caulimoviridae* RTs found in the oak genome (blue branches, n=8), reference RT sequences from eight *Caulimoviridae* genera (n=41), Gypsy LTR retrotransposons (n=5), and from mammalian endogenous retroviruses (ERV, n=4).



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Supplementary Fig. 35 Highly repeated fragments of viruses. Overview of the multiple
 sequences alignment of 762 highly similar fragments (raw data) from Caulimoviridae found
 in the pedunculate oak genome.



Supplementary Fig. 36 Distribution of Caulimoviridae along the 12 pedunculate oak
chromosomes (Qrob_Chr01-12), determined with sliding windows of 300 kb and an overlap
of 200 kb.



- Supplementary Fig. 37 Evidence of protein functions according to annotation category:
 BLAST/rpsBLAST (red), domain/motifs (green), and localization/targeting-based analysis
 (blue).

protein function evidences 25516 genes (99%)



Supplementary Fig. 38 Comparison of observed divergence of TE copies from their
respective consensus sequences, for different TE orders and superfamilies. DTX: Class II
(DNA) TIR, DHX: Class II Helitron, RLC: Class I LTR Copia, RLG: Class I LTR Gypsy,
RLX: Class I LTR other, RIX: Class I LINE, noCat: unclassified TE

Distribution of TE copies divergence



Supplementary Fig. 39 Transpositional dynamics of nine highly repeated LTRretrotransposon families in the oak genome. Histograms represent the age distribution of the retrotransposons, showing the asynchronism of retrotranspositional activity in pedunculate oak over the last six million years. The magenta curves represent local density estimates. The title of each histogram indicates the family name and its number of copies.

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Supplementary Fig. 40 Distribution of TE (red area), genes (green area) and GC content
(blue line) along the 12 chromosomes (Qrob_Chr01-12) of the pedunculate oak genome.

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Supplementary Fig. 41 Distribution of the Gypsy (light-blue) and Copia (dark-blue)
 superfamily of ClassI-LTR retrotransposons along the 12 chromosomes (Qrob_Chr01-12) of
 the pedunculate oak genome sequence.


2397 **Supplementary Fig. 42** Comparison of gene-to-closest TE distance between genes from 2398 expanded gene families (n=5,433 genes) and genes from unchanged gene families (n=15,166 2399 genes). (A) Two classes of distance [1-500bp], [501-5000bp] Pearson's Chi-squared test with 2400 Yates' continuity correction: *P*-value = $2.2e^{-16}$. (B) 10 classes of distance [1-500 bp], [501-2401 1000 bp]...[4501-5000 bp] Pearson's Chi-squared test: *P*-value = $2.2e^{-16}$.



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2406 **Supplementary Fig. 43**Comparison of gene-to-closest TE distance between sets of tandemly 2407 duplicated genes (TDG; n=8,532 genes) and single copy genes (SG; n=6,325 genes). (A) Two 2408 classes of distance [1-500 bp], [501-5000 bp] Pearson's Chi-squared test with Yates' 2409 continuity correction: *P*-value = $2.2e^{-16}$. (B) 10 classes of distance [1-500 bp], [501-1000 2410 bp]...[4501-5000 bp] Pearson's Chi-squared test: *P*-value = $1.5e10^{-14}$.

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Supplementary Fig. 44 Phylogenetic analysis of aquaporins. (A) Proteins are from *Quercus robur* (Qrob, dot), *Arabidopsis thaliana* (At⁵⁸) and *Populus trichocarpa* (Pt⁶¹). Protein sequences were compared in ClustalW analyses, and a consensus Neighbor-Joining tree was generated in MEGA6 (bootstrap: 500 replicates, distance based on number of differences method excluding gaps). (B) Exon-intron structure of *Quercus robur* aquaporins as displayed by GSDS2.0 (http://gsds.cbi.pku.edu.cn/).





Supplementary Fig. 45 Phylogenetic analysis of R2R3-MYB. R2R3-MYB proteins are from 2423 2424 Quercus robur, Populus trichocarpa, Eucalyptus grandis, Vitis vinifera, Arabidopsis thaliana 2425 and Oriza sativa. The R2R3-MYB proteins selected for E. grandis, V. vinifera, A. thaliana and O. sativa are the same as those used by Soler et al.⁶³, except for LOC_Os01g62410, which 2426 2427 was not used. For P. trichocarpa, the MYB proteins were selected as described by Chai et al.¹⁷², Potri.013G046300.1. 2428 except for Potri.003G1238001, Potri.015G143400.1, 2429 Potri.019G018400.2, Potri.006G097300.1, Potri.016G112300.1, and Potri.008G064200.1, which were not included. These sequences were discarded after manual inspection. R2R3-2430 MYBs were aligned using MAFFT with the FFT-NS-i algorithm¹⁷³, and a neighbour-joining 2431 phylogenetic tree with 1000 bootstrap replicates was constructed with MEGA5¹⁷⁴, with the 2432 2433 Jones-Taylor-Thornton substitution model used to calculate the evolutionary distances, a rate 2434 of variation between sites with a gamma distribution of 1, and the comparison of sequences 2435 with the complete deletion method. Bootstrap values are shown next to the branches. The tree 2436 is drawn to scale, with branch lengths calculated on the basis of the number of amino-acid 2437 substitutions per site. Each triangle represents a R2R3-MYB subgroup. Subgroup names are 2438 included next to each clade, together with a short name to simplify nomenclature. The total 2439 number of R2R3-MYB genes of each species, as a whole and for each subgroup, is also 2440 included. Subgroups expanded in woody plants are highlighted in light orange or red.

	Q. ro	P. tr	E. gr	V. vi	A. th	O. sa
¹⁷ Subgroup 6 (S6)	1	5	11	10	4	0
23 93 Woody Preferential Subgroup I (WPS-I)	5	4	6	3	0	0
94 ✓ Subgroup AtMYB82 (SAtM82)	1	2	1	2	1	0
¹⁶ 3 ¹ _93 Subgroup 15 (S15)	4	3	4	3	3	0
12 86 Subgroup AtMYB5 (SAtM5)	3	6	2	2	1	1
39 23 Subgroup 5 (S5)	18	6	16	11	1	2
19 70 Woody Preferential Subgroup II (WPS-II)	4	5	3	1	0	0
31 51 Subgroup 4 + AtMYB6 & AtMYB8 (S4)	4	7	4	5	6	5
⁵² [™] Woody Preferential Subgroup III (WPS-III)	5	5	4	4	0	0
Subgroup 7 (S7)	1	3	1	2	3	2
25 55 Woody Preferential Subgroup IV (WPS-IV)	1	4	6	1	0	0
36 Woody Preferential Subgroup V (WPS-V)	12	5	2	2	0	0
2 48 98 Subgroup AtMYB35 (SAtM35)	2	3	1	2	1	1
2 38 Subgroup 14 (S14)	7	10	10	7	6	9
Subgroup 2 & Subgroup 3 + AtMYB10 & AtMYB72 (S2 & S3)	4	6	4	5	7	5
Subgroup 10 & Subgroup 24 (S10 & S24)	3	7	5	6	6	6
Subgroup 1 (S1)	3	4	4	3	5	7
1 16 Subgroup 9a (S9a)	2	4	1	3	2	3
Crob_P0555330.2 45 Qrob_P0555340.2 41g18710.1/atMYB47 13 99At1g18710.1/atMYB95						
Subgroup 12 (S12)	0	0	0	0	6	0
	2	4	2	2		2
	3	0	5	3	4	
95 Subgroup AtMYB40 (SAtM40)	1	2	2	1	1	0
75 AtMYB43, AtMYB85 & AtMYB99 (SAtM85)	2	6	4	3	5	5
92 Subgroup 16 (S16)	1	1	2	1	3	3
15 Subgroup AtMYB103 (SAtM103)	1	2	1	1	1	1
32 g/ Subgroup AtMYB26 & AtMYB67 (SAtM26)	10	5	1	3	2	2
18 Subgroup 13 (\$13)	3	7	7	4	4	2
						0
Subgroup AtMYB71, AtMYB79 & AtMYB121 (SAtM71)	4	5	1	2	3	3
Subgroup AtMYB27, AtMYB48 & AtMYB59 (SAtM27)	2	3	1	2	3	3
Subgroup 19 + AtMYB57 (S19)	1 3	2 8	2 4	1	3 6	2
LOC_Os05g41166.1						
19 19 Subgroup 18 + SAtMYB125 (S18 & SAtM125)	4	7	3	5	7	5
At2g26950.1/AtMYB104 (Subgroup 18)	2	3	1	2	1	1
Subgroup 21 + AtMYB89 (SAtM21)	5	13	6	6	8	4
15 45 Subgroup 22 (S22)	5	9	5	2	4	5
Subgroup 23 (S23) 62 LOC_0s03g13310.1	1	1	2	2	3	1
60 Subgroup 25 + AtMYB98 (S25)	2	6	2	3	7	5
61 99 Subgroup AtMYB88 & AtMYB124 (SAtM88)	1	2	2	1	2	1
0.1	120	197	1/1	122	126	105
	139	107	141	123	120	105

Supplementary Fig. 46 Classification and percentage of pedunculate oak R2R3-MYB genes as a function of their mode of duplication and expansion in woody perennials. R2R3-MYB genes were first classified into three categories on the basis of duplication mode (see online methods): tandem duplicated genes (TDGs), long distance-duplicated genes (LDGs), and singleton genes (SGs). The TDGs and LDGs were further classified into genes belonging or not belonging to subgroups expanded in woody perennials.

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Supplementary Fig. 47 Phylogenetic analysis of SWEET. Sequences were aligned by 2456 2457 ClustalW and a tree was constructed with the neighbor-joining method. The different clades of SWEET genes defined by Chen et al.⁷¹ are color-coded. *Qrob* indicates predicted 2458 2459 pedunculate oak (Quercus robur) polypeptides, and the reference species are abbreviated as 2460 follows: Arabidopsis thaliana (At); Solanum tuberosum (St); Eucalyptus grandis (Eg); Malus 2461 domestica (Md). The tree is rooted on SWEET homologs from Chlamydomonas reinhardtii 2462 (Cr). The symbols indicate pedunculate oak genes differentially expressed during interactions with the ectomycorrhizal fungi Piloderma croceum and Tuber magnatum, the ectomycorrhiza 2463 2464 helper bacterium Streptomyces sp. AcH 505, and the causal agent of oak powdery mildew 2465 Erysiphe alphitoides. For phylogenetic analysis, protein-coding sequences from Arabidopsis thaliana, Solanum tuberosum, Eucalyptus grandis, Malus domestica and Chlamvdomonas 2466 2467 (non-plant reference) obtained reinhardtii were from the NCBI 2468 (http://www.ncbi.nlm.nih.gov), and protein-coding sequences from oak were extracted from 2469 the haplome. Phylogenetic distances between the SWEET proteins were calculated from a 2470 multiple sequence alignment (ClustalW), by the neighbor-joining method (MEGA6), with 2471 bootstrapping (1000 replicates).



Supplementary Fig. 48 Phylogenetic analysis of GSTUs. Sequences encoding GSTUs from Quercus robur, Populus trichocarpa, Arabidopsis thaliana, Oryza sativa and Sorghum *bicolor* were retrieved from Phytozome (https://phytozome.jgi.doe.gov/pz/portal.html). Sequences were then aligned with Clustal-Omega¹⁷⁵. The alignment was manually adjusted with SeaView software ¹⁰⁷ and curated with GBlocks¹⁷⁶. The unrooted phylogenetic tree was constructed with BioNJ¹⁷⁷ in Seaview and further edited with FigTree software (http://tree.bio.ed.ac.uk/software/figtree/). The robustness of the branches was assessed by the bootstrap method with 1000 replications (not shown). Sequences corresponding to Quercus robur, Populus trichocarpa, Arabidopsis thaliana, Oryza sativa and Sorghum bicolor GSTUs are shown in red, blue, green, cyan and pink, respectively. The expanded clusters identified in orthoMCL analysis are highlighted on red branches. Given its considerable divergence, the Qrob_P0196930.2 protein annotated as GSTU was removed from the analysis.



2489 Supplementary Fig. 49 Phylogenetic analysis of MLO. The proteins used are from *Quercus robur* (n=19 regular genes and 7 unreliable genes), *Prunus persica* $(n=18)^{92}$ and *Arabidopsis* 2490 thaliana (n=7, TAIR - https://www.arabidopsis.org/). The analysis was performed on the 2491 Phylogeny.fr platform¹⁷⁸, as follows: alignment with MUSCLE (v3.8.31) configured for 2492 2493 highest accuracy; removal of ambiguous regions (i.e. containing gaps and/or poorly aligned) 2494 with Gblocks (v0.91b) using the following parameters (minimum length of a block after gap 2495 cleaning: 10; no gap positions were allowed in the final alignment; all segments with 2496 contiguous nonconserved positions of more than eight residues were rejected; minimum 2497 number of sequences for a flanking position: 85%); reconstruction of the phylogenetic tree 2498 with the maximum likelihood method implemented in PhyML 3.0. The WAG substitution 2499 model was selected, assuming an estimated proportion of invariant sites of 0.003 and four 2500 gamma-distributed rate categories to account for rate heterogeneity across sites. The gamma shape parameter was estimated directly from the data (gamma=1.340). The reliability of 2501 2502 internal branches was assessed with the aLRT test (SH-Like). Graphical representation and 2503 phylogenetic tree generation were achieved with TreeDyn (v198.3). Branch boostrap support values are displayed in blue. Clades were named according to the presence of Arabidopsis 2504 thaliana and Prunus persica proteins ⁹². 2505





Supplementary Fig. 50 Biosynthesis of hydrolyzable tannins from gallic acid, via the β glucogallin intermediate.



Supplementary Fig. 51 Phylogeny of oak genes potentially involved in hydrolyzable tannin biosynthesis. (A) Phylogeny of annotated oak genes and Arabidopsis genes involved in the chorismate pathway. Genes encoding 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase (DHS), 3-dehydroquinate synthase (DHQS), 3-dehydroquinate dehydratase/shikimate 5-dehydrogenase (DHQ-SDH), shikimate kinase (SK), 5-enolpyruvylshikimate 3-phosphate synthase (EPSPS) and chorismate synthase (CS) are presented. (B) Phylogeny of the Arabidopsis members and annotated oak members of the UGT 74, 75, 83 and 84 families. Protein sequences were aligned using ClustalW and the UPGMA tree was drawn based on Jukes-Cantor distances (with Geneious 6.1.8).



Supplementary Fig. 52 Comparative phylogenetic analysis of the laccase protein sequences from Quercus robur, Populus trichocarpa, Eucalyptus grandis, Vitis vinifera, Arabidopsis *thaliana* and *Oriza sativa*. Sequences were aligned with Clustal-Omega¹⁷⁵. The alignment was manually adjusted with SeaView software¹⁰⁷ and curated with GBlocks¹⁷⁶. *Arabidopsis* ascorbate oxidases (AO1, AO2, and AO3) were added and used as an outgroup. The phylogenetic tree was calculated with SeaView, using PhyML with the LG model and the aLRT method for branch support, NNI heuristic for optimal tree structure search and BioNJ for optimizing tree topology. The phylogenetic tree was further edited with FigTree software (http://tree.bio.ed.ac.uk/software/figtree/).



Supplementary Fig. 53 Example of a pedunculate oak specimen sampled for genetic
diversity analysis by the pool-seq approach.



2543 © Christophe Plomion

Supplementary Fig. 54 Distribution of π_4 , π_0 and the π_0/π_4 ratio for gene families.



Supplementary Fig. 55 A phylogenetic tree for eudicots, based on the tree generated by 2549 Zanne et al.¹³⁸ collapsed to the order level with clade sizes denoted by triangle size. For each 2550 order, we show the number of woody and herbaceous species for which whole-genome 2551 will 2552 sequences are already or soon be available (as reported in 2553 https://genomevolution.org/wiki/index.php/Sequenced_plant_genomes as of 18 December 2016). Variable and diverse species, according to Fitzjohn et al.¹⁴⁹, for which genome 2554 sequences are available are highlighted in green. 2555





Supplementary Fig. 56 Number of GO terms per gene for the 16,820 pedunculate oak genemodels with a GO.



Supplementary Fig. 57 Number of genes per Gene Ontology (GO) term for the 1,722 unique

2564 pedunculate oak GO terms.



Supplementary Fig. 58 Number of genes within orthoMCL orthogroups expanded in pedunculate oak.



Supplementary Fig. 59 Number of gene ontology (GO) terms per orthogroup expanded in
pedunculate oak, for the 1,722 unique GO terms.





Supplementary Fig. 60 Fold-enrichment (*x*-axis) of significant gene ontology (GO) terms (P<0.01) of the orthogroups expanded in pedunculate oak relative to the background of the whole genome. GO representing biological processes are shown as red lines, cellular components are shown in blue and molecular functions are shown in green. The vertical dashed line represents the 1.5 threshold from which we considered interesting biologically relevant enrichments. Sample sizes are provided in **Supplementary Data Set 8** sheet #4.

