Supplementary Data 1: Extended Methods and Results

Introduction into Quercus Section Cerris

Quercus section Cerris Dumort. (Denk et al., 2017a) is a small group of oaks of the Eurasian subgenus Cerris Oerst., which includes the iconic cork oak, Q. suber L., the source of commercial cork. Well-developed cork layers are present not only in the western Mediterranean Q. suber but also in the East Asian Q. variabilis Blume ("Chinese cork oak"). Both are used industrially (Huang et al., 1999). Clearly developed but much thinner cork layers occur in Q. crenata Lam., Q. afares Pomel, and some varieties and forms of Q. cerris L. (for example var. gussonei Borzì in Sicily; "subsp./var. pseudocerris" in Turkey and Lebanon; Maire, 1961; Schirone et al., 2015; Pignatti, 2017–2019). Distinctly corky bark does not occur in any other oak section, but it is present in at least five out of the 15 species included in section Cerris and justifies the name "cork oaks" for the section.

The cork oak section is remarkable biogeographically and ecologically. First, it is the only group of oaks that is most diverse in western Eurasia, despite also being widespread in East Asia (Supplementary Data 3). Second, it is strictly deciduous but comprises several semievergreen species adapted to summer-dry and winter-cold, continental climates. Third, a single species in this group, *Q. cerris*, the most widespread western Eurasian cork oak, independently evolved lobed leaves reminiscent of many species in *Quercus* section *Quercus*, the white oaks. Finally, the cork oaks provide well-documented examples of intrasectional and intersectional introgressive hybridization that provide insight into the timing of diversification and gene flow in the genus. In a recent study, Simeone et al. (2018) investigated western Eurasian members of section Cerris using a broad-sampled chloroplast (trnH-psbA) and a multi-copy nuclear (5S-IGS) marker. Chloroplast haplotypes reflected shared biogeographic histories of all western Eurasian Cerris species and its sister section Ilex, indicating several phases of "chloroplast capture" and secondary, species-decoupled, geographic sorting. Simeone et al. (2018) suggested that the precursors of western Eurasian Cerris species came into contact with Ilex only after the initial range expansion of *Cerris* from north-eastern Asia during the early Oligocene. In addition, four main nuclear 5S-IGS lineages were identified within Cerris, three of which were private to potential sibling species: (i) root-proximal 'Oriental' lineage in Q. afares, Q. libani Olivier, Q. trojana Webb; (ii) 'Occidental' lineage found in the widespread Q. cerris and the narrow-endemics *Q. castaneifolia* C.A.Mey. (northern Iran), *Q. look* Kotschy (Levant), and *Q. euboica* (Papioann.) K.I.Chr. (Euboea), and as intra-individual dimorphism in western *Q. trojana*; (*iii*) 'Vallonea' lineage exclusive to the eastern Mediterranean *Q. brantii* Lindl., *Q. ithaburensis* Decne., and *Q. macrolepis* Kotschy; and (*iv*) a derived lineage characterizing the western-most species pair *Q. crenata-Q. suber* (Fig. S4-1).

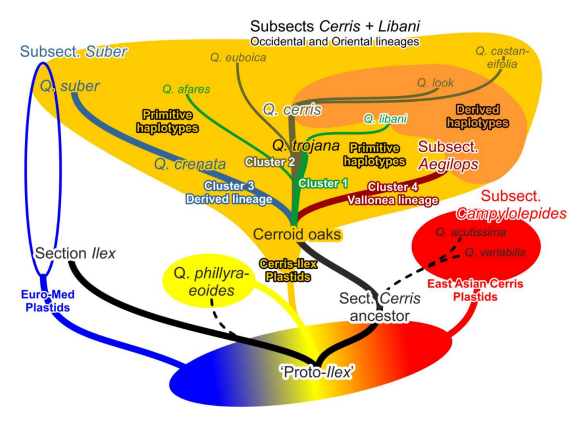


Figure S1-1. Cactus-type tree metaphor summarizing putative reticulate-evolutionary pathways in western Eurasian species of section *Cerris* (modified after Simeone *et al.*, 2018, fig. 6). Lines indicate major nuclear (5S-IGS) lineages, coloured fields shared plastid (*trnH-psbA*) haplotypes.

Stratigraphic note

We follow the latest version (v. 2022/10) of the International Chronostratigraphic Chart (https://stratigraphy.org/; Cohen *et al.*, 2013, updated). The major change has been the re-recognition of lower/early, middle, and upper/late Neogene and Quaternary strata as formally accepted and named units, in some cases between series/epochs and stages/ages (Aquitanian + Burdigalian = Early Miocene; Langhian + Serravallian = Middle Miocene; Tortonian + Messinian = Late Miocene; Gelasian + Calabrian = Early Pleistocene).

Methods: Biogeography and Climatic Niche

Fossilized Birth-Death analysis

For fossilized birth-death (FBD) analyses, the RAD-seq matrix was reduced to 29 tips, a single tip per species within both sects *Cerris* and *Ilex*, with the exception of *Q. cerris*, for which two individuals were kept that did not group together in any analyses and might represent cryptic species. Loci were retained if they were present in at least 10 individuals. A NEXUS file was exported using the RADAMI package, including 47 additional lines of undetermined positions (coded as "?"), one per fossil included in the FBD analyses.

FBD analyses were conducted in BEAST2 (Bouckaert et al., 2014). Markov chain Monte Carlo (MCMC) runs of 50 million generations each were run from ten independent random starting points on each of three random draws from the uniform distribution of the fossil age ranges. Analyses were conducted using a nucleotide substitution model that allows for rate variation and invariant sites $(\Gamma + I)$, with the shape parameter (α) and proportion of invariant positions estimated, and four gamma categories. The relaxed log normal clock was used, with the clock rate estimated. Priors were specified using defaults with these exceptions: origin of the clade was set at an initial position of 58 million years (Ma), with a range of 60 to 49 Ma, based on previous results (Hipp et al., 2020) that suggest the origin of subgenus Cerris to be close to the origin of the genus; and sampling proportion was estimated with bounds between 0.4 and 0.6, with a starting proportion of 0.45. Clade priors were specified by assigning each fossil in section Cerris or Ilex to one of thirteen pre-defined clades identified in the maximum likelihood analysis as MRCA priors, enforcing monophyly of the clade including the fossils and extant taxa (Supplementary Table 3). The position of the three Cyclobalanopsis fossils was designated by assigning all taxa except the Cyclobalanopsis fossils to a single "not Cyclobalanopsis" clade. Scripts for exporting data, BEAST2 XML configuration files, and RAD-seq data matrices are all archived in the code repository for this paper (release v0.99-1: https://github.com/andrew-hipp/cerris-fbd; https://doi.org/10.5281/zenodo.7369341), along with output files and instructions for executing the scripts.

Köppen profiles and biome profiles

We used grid-weighted 'Köppen signatures' (Denk *et al.*, 2013; Bouchal *et al.*, 2018; Grímsson *et al.*, 2018), henceforth 'Köppen profiles', to summarize the climate niches occupied by species of *Cerris* and to investigate climate niche evolution within and among subsections of *Cerris* (*main-text Table 1*; *Supplementary Data 3*). A Köppen profile (**Fig. S1-2**) reflects the proportional coverage of the various Köppen-Geiger climate zones (cf. Kottek *et al.*, 2006; Peel *et al.*, 2007) by a modern species based on gridded distribution data.

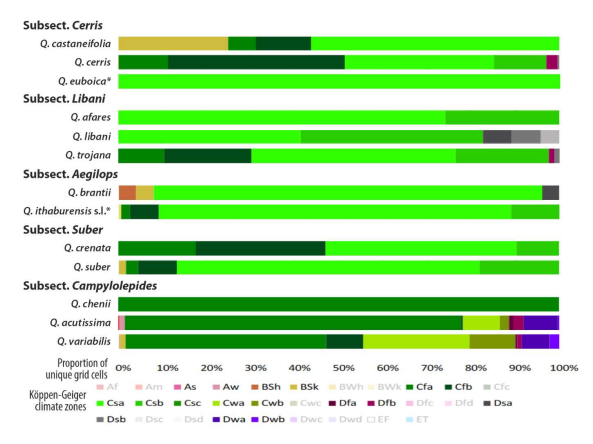


Figure S1-2. Köppen profiles for species of section *Cerris*, sorted by phylogeny. Bottom: earliest diverging subsectional lineage; top: *Cerris* core clade comprising sects *Cerris* and *Libani* (see *Supplementary Data 3* for full details). * GBIF data doesn't distinguish between *Q. macrolepis* and *Q. ithaburensis* (s.str.) and the geographic overlap between the two species has so far not been assessed *in situ*; no SNP data are available for *Q. euboica*, placed based on Simeone *et al.* (2018).

Today, *Cerris* oaks thrive in three main climate types: Subtropical to mild-temperate "warm temperate" C climates with average temperatures of the coldest month $T_{min} > -3$ °C and < +18 °C, (cool-)temperate "snow climates" with pronounced winter cold, D, $T_{min} < -3$ °C, and

tropical "equatorial" A climates (marginally, Indochina) with $T_{min} > +18$ °C. The second letters in the two-/three-letter climate formula used in the Köppen-Geiger classification indicate fully humid (f) summer-dry (s), or winter-dry (w) conditions. In addition, for equatorial climates, the second letter m indicates monsoon climates. For details of the three-letter code, see Kottek et al. (2006) and Peel et al. (2007). Some species further occur in arid B climates (main-text Table 1; Supplementary Data 3). We opted for using the Köppen-Geiger system, which does not recognize a subtropical zone, instead of the Köppen-Trewartha system because there are no ready-to-use high-resolution Köppen-Trewartha (Trewartha and Horn, 1980) climate grids. Furthermore, the Scotese et al. (2014) palaeo-climate zones used to cross-check climate preference of fossil-taxa (see below) correspond to the Köppen-Geiger system, distinguishing five major zones: 'tropical' (= equatorial A-climates), 'arid' (= B-climates), 'warm (temperate)' (\approx warm temperate C-climates), 'cool (temperate)' ($\approx Dxa/Dxb$, snow climates with summer) and 'cold' (continental-dry D and E climates without summer).

Modern species distributions were connected to fossil distributions by using georeferenced occurrence data for each species, downloaded from the GBIF database (www.gbif.org; Supplementary Data 3). Each data set was checked for natural distribution outliers (e.g. specimens from botanical gardens outside the natural distribution range of a species). Published chorological data were used to detect these outliers (e.g. Browicz and Zieliński, 1982; Fang et al., 2009; San-Miguel-Ayanz et al., 2016; Caudullo et al., 2017). The cleaned georeferenced occurrence data were then plotted onto 5 arc minutes Köppen-Geiger grid (1986–2010 data; Rubel et al., 2017) to establish Köppen profiles for all species of section Cerris (Fig. S1-2); and on major terrestrial biome maps (Olson et al., 2001; Fig. S1-3) to assess species' forest biome preferences (full details in Supplementary Data 3). For the Köppen-Geiger plots, the georeferenced dataset was filtered so that multiple occurrences in a single grid cell were only counted once (labelled 'unique grid cells' in the diagrams). Likewise, for the biome plots, georeferenced data were filtered so that multiple occurrences with the same coordinates were treated as single occurrences (labelled 'unique localities' in the diagrams).

The georeferenced data and the Köppen-Geiger map with resolution of 5 arc minutes were processed using the 'Sample Raster Values Toolbox' in QGIS Version 3.16.4-Hannover (http://www.qgis.org). The biome shape files were processed using the 'Geoprocessing Tool' in QGIS. The biomes and Köppen-Geiger climates occupied by extant members of section *Cerris* are shown as maps generated with QGIS and as frequency (proportional distribution)

diagrams. The raw coordinate data are archived in EXCEL spreadsheet format (Supplementary Data 3: Tabulated Data S3-1).

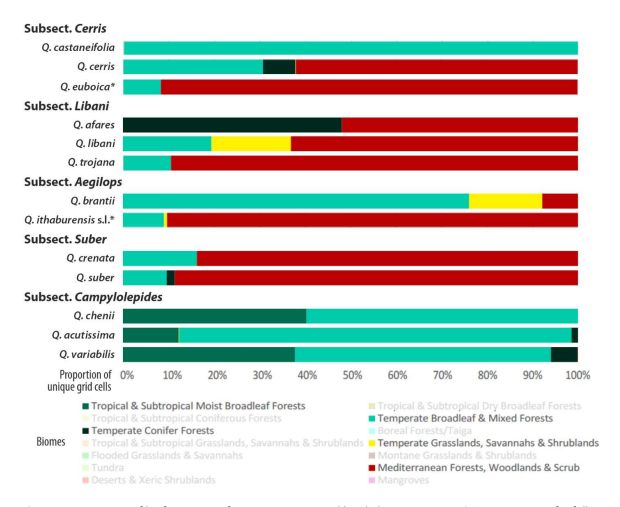


Figure S1-3. Biome profiles for species of section *Cerris*, sorted by phylogeny. See *Supplementary Data 3* for full details. * GBIF data doesn't distinguish between *Q. macrolepis* and *Q. ithaburensis* (s.str.); *Q. euboica* placed after Simeone *et al.* (2018).

In addition, historical climate data (1970–2000) were compiled for 2,779 georeferenced occurrences for members of *Quercus* sect. *Cerris* using WORLDCLIM vers. 2.1 (https://www.worldclim.org/data/worldclim21.html) at a resolution of 30 seconds (ca. 1 km², Fick *et al.*, 2017; *Supplementary Data 3*: Tabulated Data S3-3). To characterize the climate envelope of modern-day species, we plotted the mean temperature of the coldest month (MTCM) against precipitation during the winter quarter (PCQ; WORLDCLIM variable BIO19), as well as monthly temperature and precipitation averages (MMT, MMP) and monthly minimal temperatures (MTmin; *Supplementary Data 3*).

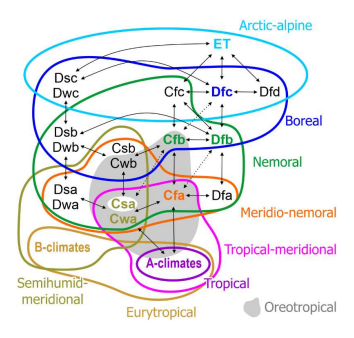


Figure S1-4. Categorisation of plants as vegetation elements based on their 'Köppen signatures' (after Denk *et al.*, 2013; emended by Bouchal *et al.*, 2018, and Grímsson *et al.*, 2018). Arrows indicate modern-day latitudinal/longitudinal and altitudinal transitions (neighbouring climates). Oaks of sect. *Cerris* do not extend into climates inhabited by 'arctic-alpine' vegetation elements; using this scheme they represent 'semihumid-meridional', 'meridio-nemoral' and 'nemoral' vegetation elements.

Maximum likelihood reconstructions of major climate niches and main biomes

Based on the quantitative assessment of biome and climate zone preferences of the modern-day species, we binned extant and fossil species into five basic categories (Supplementary Table 2), accounting either for biome or climate zone preferences (main-text Table 1; Supplementary Data 3). Our generalisation and categorisation make use of the terminology and concepts introduced by Schroeder (1998) and Denk et al. (2013; see Fig. S1-4) and allow (i) direct comparison of biome and climate zones preferences, which are commonly correlated but not synonymous, and (ii) relation of quantitative modern-day categorisation qualitatively to our fossil-taxon set. Towards that end, we first defined for each fossil-taxon of section Cerris (columns Biome and Major Köppen climate type in Supplementary Table 2) the putative covered biomes: 'Tropical & Subtropical Coniferous Forest', 'Tropical & Subtropical Moist Broadleaf Forest', 'Mediterranean Forests, Woodlands, and Scrubs', and 'Temperate Conifer Forest'. We also specified climate zones for each fossil taxon, distinguishable in the fossil record as five principal combinations: (i) Cf + Cw (may include Aw for oldest fossils); (ii)

chiefly Cf; (iii) Cf/Cw extending into Df/Dw; (iv) Cf + Df; and (v) Cf + Cs. The scoring is based on our knowledge about the fossil floras (place, time, plant association) and general background information about the climatic history of the northern hemisphere (Scotese *et al.*, 2014). We then identified common patterns and assigned the following five categories to the modern-day and fossil-taxa (**Fig. S1-5**; columns *Category according biome*; *Category according climate zone* in *Supplementary Table 2*; matrix *MainClimates* in **Data File S1-1**; characters #10 – biome class, and #11 – climate class).

0–Moist-Subtropical: Associated exclusively with the Tropical and Subtropical Moist Broadleaf Forests biome; fossil species that can be associated with Cf climates during global greenhouse phases or at low latitudes, i.e. exclusively within the southern half of the 'warm' zones (\approx modern-day humid subtropics), possibly extending in the extinct 'boreal tropical' palaeo-climate zones in Scotese *et al.* (2014). In case of modern-day Cerris oaks, species are associated almost exclusively with summer-moist climates with hot summers (Cfa, Cwa climates). The only modern species with an accordingly characteristic climate niche is the East Asian Q. Chenii, firmly restricted to the Cfa climate of central-eastern China.

1–Meridio-Nemoral (modified from Denk *et al.*, 2013): Associated with the ecotone between Tropical and Subtropical Moist Broadleaf Forests and Temperate Broadleaf and Mixed Forests biomes. Fossil-species are linked exclusively to *Cf* climates and are placed within Scotese *et al.*'s (2014) 'warm(-temperate)' palaeo-climate zones. A modern-day Meridio-Nemoral species has its main distribution in subtropical to temperate climates with ample precipitation in the hot summer (typically *Cfa* + *Cwa* extending into warm *Cfb/Cwb* variants).

2–Nemoral (cf. Schroeder 1998; Denk *et al.*, 2013): Either restricted to Temperate Broadleaf and Mixed Forests or extending into both Tropical and Subtropical Moist Broadleaf Forests and Temperate Coniferous Forests biomes; fossil species associated with fully humid *Cf* and *Df* climates ranging from 'warm' into 'cool(-temperate)' palaeo-climate zones. Nemoral species are fully temperate species preferring mild climates with ample precipitation during the typically long (~8 months with MTCM > 10 °C) growing season. Key taxon for this category is *Fagus* (beech), which has its climax distribution in fully temperate *Cfb* climates, extending into subtropical *Cfa* lowland and, latitudinally and altitudinally, into cool-temperate *Dfa/Dfb* climates. Fossil assemblages of this category would typically contain broadleaved oaks of subgenus *Quercus* associated with beech and maples with palmate leaves (as found in modern-day species of *Acer* sects *Acer*, *Macrophylla*, *Palmata* p.p., and *Platanoidea*). The only modern-

day *Cerris* oak that qualifies for this category is *Q. variabilis*, being the most widespread species of section *Cerris* with a strong preference for the Temperate Broadleaf and Mixed Forests biome and the full range of summer-humid warm temperate (*Cf*, *Cw*) climates.

3–Meridional (cf. Schroeder, 1998): Generalists tolerating summer-drought, otherwise similar to the Nemoral category. Fossil-species of this category can be associated with Tropical and Subtropical Moist Broadleaf Forests *and* biomes with seasonal drought/ water stress (Mediterranean Forests, Woodlands, and Scrub and Temperate Coniferous Forests biomes); they thrive in fully humid *Cf* as well as summer-dry *Cs* climates. The natural distribution of modern-day Meridional species is restricted to the subtropical belt but characterized by a split preference for both fully Mediterranean lowland climates (unlike in species of the previous categories), summer-dry with hot summers (*Csa*), and their altitudinal, potentially moister successions: summer-dry (*Csb*) or perhumid (*Cfb*) climates with warm summers. In contrast to the species categorized as Full-Mediterranean (next category), they can endure substantial winter cold and frost phases, with some species extending into the boreal, continental, summer-dry snow climates found at mid-latitudes in Western Eurasia. They do not usually form part of coastal vegetation but prefer higher altitudes and the interior.

4–Full-Mediterranean: Summer-drought tolerant specialists restricted to summer-hot and winter-mild biomes and climates. Modern-day species of this category are restricted to Mediterranean Forests, Woodlands, and Scrub biome (or Temperate Coniferous Forests biome: *Q. afares*) and hot, summer-dry *Csa* climates, while only marginally extending into moister biomes and moister (Mediterranean-*Cfa*) or colder (*Csb*) climates. Further, Full-Mediterranean species can be found in hot or cold "steppe" climates (*BSh*, *BSk*).

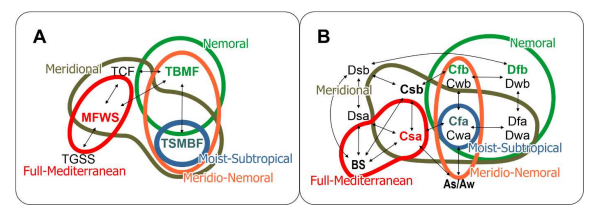


Figure S1-5. Schematic illustrating the biomes (A) and climates (B) covered by the here used general categories.

Ancestral states of the unordered five-state categorical climate character were reconstructed under the *Mk1* model in MESQUITE v 2.75 (Maddison and Maddison, 2011). We used two different input trees: (i) the original dated tree for standard top-down reconstruction of ancestral states, i.e. using only the information scored for the modern-day species; (ii) the dated tree with nodes and tips added to account for states of fossil-taxa. Fossil-taxa that could be associated with a distinct branch (lineage) were treated as sister lineages and used to break down the according branch. We used the oldest possible age of the fossil-taxon as age of the putative MRCA, and the youngest possible age to define the MRCA-added-tip distance. The resultant Mesquite-NEXUS file is included in the Github repository/ Zenodo submission (Supplementary Data 4: Data File S4-1).

Tests for Inter-Species Gene Flow

Evidence of introgressive hybridisation was investigated using Patterson's *D*-statistic test (Durand *et al.*, 2011) as implemented in IPYRAD (Eaton and Overcast, 2020). The following text describes the rational (background), experimental set-up and summarizes the main results.

Background: Quercus afares hybrid origins

The only non-morphological evidence for a hybrid origin of *Q. afares* was the study of Mir et al. (2006), which included data from the two postulated (based on morphology and geography) donors of O. × afares: the Cerris oak O. suber, molecularly, a distant relative, and the white oak O. canariensis, sym-/parapatric with O. afares, a species from a different subgenus and much different evolutionary and geographic origin (see Denk and Grimm, 2010; see also the Fagaceae phylogenies and data compiled by Zhou et al., 2022). Without including an additional tip as outgroup not involved in the tested hybrid scenario, e.g. Q. ilex, Mir et al. (2006) were unable to perform a topology test, so their evidence is based on isozyme allele frequencies. The test is particularly interesting because a section Cerris × section Quercus hybrid is unlikely, at least in natural populations. While the occurrence of (±ancient) intersectional hybrids can be assumed for sections of the same subgenus, there has been so far no evidence for mixing between the two main lineages, the exclusively Old World (Eurasian) subgenus Cerris and the originally New World (American) subgenus Ouercus in areas (Eurasia) where they grow sympatrically today or in the past (Denk and Grimm, 2010; Simeone et al., 2018). In contrast, species groups and lineages characterized by sharing of plastid haplotypes such as Mediterranean Cerris and Ilex oaks and subtropical-tropical East Asian cycle

cup (sect. *Cyclobalanopsis*) and Ilex oaks (Simeone *et al.*, 2016, 2018; Zhou *et al.*, 2022), or for which past introgression has been inferred using nuclear data (sects *Ponticae* and *Quercus*, McVay *et al.*, 2017; sect. *Protobalanus* and *Quercus* in western North America, cf. ITS data of Manos *et al.*, 2001) always were members of the same subgenus. Inter-generic reticulation possibly can be found even within major core Fagaceae lineages such as *Notholithocarpus* and American members of subgenus *Quercus* (P. Manos, pers. comm., 2012): subgenus *Quercus* and the western North American relict genera *Notholithocarpus* and *Chrysolepis* share the same plastome lineage (Zhou *et al.*, 2022), possibly including the "Euro-Med" haplotype of *Q. ilex* in section *Ilex* (*Q. ilex* included in Yang et al., 2019; cf. Grimm 2022). In contrast, there is up to date no evidence at all for inter-subgeneric gene flow (past or present) in oaks, despite the fact that e.g. species of section *Quercus* frequently can be found in sympatry with (widespread) species of sections *Cerris* and *Ilex* across Eurasia.

Background: Quercus crenata hybrid origins

Hybridisation and introgression are more likely in the case of *Q. crenata*, postulated to be a *Q. cerris* × *Q. suber* hybrid or *Q. cerris* populations introgressed by *Q. suber*, because of its intermediate morphology, specifically the presence of well-developed cork layers in *Q. crenata* (Schwarz, 1936–1939; Pignatti, 1982). Also in this case, broadly-sampled nuclear spacer data did not produce any evidence so far for *Q. cerris*-unique (diagnostic) sequence variants in *Q. crenata*, while strongly supporting an (inclusive) common origin (holophyly, i.e. monophyly in a strict sense; cf. Ashlock, 1971) of *Q. crenata* and *Q. suber* (Denk and Grimm, 2010; Simeone *et al.*, 2018; Piredda *et al.*, 2020; the current study using phylogenomic data from carefully selected placeholder tip-set).

D-statistics

To test for introgression and hybridisation using the *D*-statistic tests, one needs SNP data from at least four tips: an outgroup defining the root of the three tested tips (p4); a tip (p3) representing one potentially introgressing/introgressed population or species; and two other tips (p2 and p1) of which one is hypothesized to introgress with p3, the other of which is not. SNPs are subsetted to the included tips and include only binary SNPs that form one of two patterns: ABBA, where A represents one nucleotide and B the other, and the pattern references p1 to p4 in order; or BABA. The relative frequencies of ABBA and BABA are compared, and *Z*-scores on bootstrap-resamples are used to calculate a *P*-value, which we report in the paper using

Holm-correction (see *Methods*) for each *D*-statistic test performed to correct for multiple testing. A significant over-abundance of the ABBA pattern suggests that phylogenetic discordance implied by the SNPs is likely due to introgression between p3 and p2; a significant over-abundance of the BABA pattern suggests introgression between p3 and p1. A non-significant *D*-statistic result suggests that phylogenetic discordance implied by the SNPs can be accounted for as an outcome of incomplete lineage sorting.

Our tests focused on two primary hypotheses: the hybrid origins of *Q. afares* (as proposed in, e.g., Mir et al., 2006, Welter et al., 2012, and Mhamdi et al., 2013) and of *Q. crenata*. We specifically tested for admixture in *Q. afares* and *Q. crenata* because both taxa have been traditionally viewed as (F1) hybrids: *Q. afares* as *Q. suber* × *Q. canariensis* (the latter belonging to sect. *Quercus*; e.g. Mir et al., 2006); *Q. crenata* as *Q. cerris* × *Q. suber*. In addition, we performed follow-up tests for admixture between *Q. ilex* and *Q. suber* (locally sharing *Q. ilex*-unique "Euro-Med" plastid haplotypes in the western Mediterranean; Simeone et al., 2018), *Q. cerris* and *Q. afares*, and among subsections *Aegilops*, *Suber*, and *Libani*.

Methods

RAD-seq data were clustered de novo in IPYRAD v 0.9.84 for 34 individuals comprising all section Cerris individuals, all Quercus ilex individuals, four individuals from section Cyclobalanopsis, Quercus canariensis, and Notholithocarpus densiflorus as an outgroup, using the same clustering thresholds used above but retaining loci with a minimum of four samples, to increase the number of potential loci usable for D-statistic tests. Tests were conducted by generating all possible combinations of the outgroups, the first potential introgressing individuals (introgressant) in our dataset (first species listed in each comparison) as taxon p3, the individuals of the second potential introgressant in our dataset as taxon p2, and all other taxa compatible with the pectinate topology used in the D-statistic test as taxon p1 (see Durand et al., 2011, fig. 1), with ABBA as the expected dominant pattern under the introgression hypothesis being tested in each case. One test ("canariensis-afares") was limited to 16 tests with a range of possible sisters because the unconstrained test caused errors. Z-scores were calculated for each test, and two-tailed P-values for Z-scores were corrected for multiple test biases using Holm-Bonferroni correction in R, with correction applied separately for each of the ten hypotheses investigated. For all hypotheses performed, effect of the p3 or p1 taxon was investigated by summarizing D-stats, Z-scores, and corrected p-values for the tests in which each taxon was included. Scripts for performing and summarizing analyses are in the 'D stats'

subfolder of https://github.com/andrew-hipp/cerris-fbd (https://doi.org/10.5281/zenodo. (6595965).

A total of 13 hypotheses are summarized in *main-text Table 3*, which summarizes the results over all tests performed for each hypothesis; and **Tabulated Data S1-1**, which summarizes test results for specific focal individuals in either the p3 or p1 position for the tests performed. Summaries are provided by indicating the number of tests performed, and then D, Z, and Holm-Bonferroni-corrected P, all with mean and 95% quantiles, as well as the percent of tests significant at $P \le 0.01$. Test numbers in **Tabulated Data S1-1** correspond to test numbers reported in *Table 3* of the main text.

Two additional tests were performed but not included in the results or discussion of the main paper for clarity: a test of relative importance of introgression with the two *Q. crenata* individuals included, using all species potentially introgressing with *Q. crenata* as tested above with each of the *Q. crenata* individuals as sister species. This test result is reported in **Tabulated Data S1-1**. The second is a five-taxon ("partitioned") *D*-statistic test evaluating the direction of introgression with *Q. crenata* and potential introgressants. This is reported in **Tabulated Data S1-2**.

Taxa included in D-statistic tests performed

The following summarizes what taxa were included in each test performed, and how many topologies (n) were evaluated for each test. Details of which individuals were included are in the associated Python scripts (as indexed and detailed in https://github.com/andrew-hipp/cerris-fbd/blob/main/D stats/finalForPaper/AAA.testsSummarized.md).

- A1 -- cerris-crenata (n = 500): Q. cerris with Q. crenata
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. cerris, potential introgressant 1
 - o p2 Q. crenata, potential introgressant 2
 - o p1 Q. suber as sister species to Q. crenata
- A2 -- crenata-cerris (n = 2560): Q. crenata with Q. cerris
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. crenata, potential introgressant 1
 - o p2 Q. cerris, potential introgressant 2
 - o p1 undefined --- anyone who fits the topology

- B1 -- canariensis-afares (n = 16): Q. canariensis with Q. afares
 - o p4 Notholithocarpus as outgroup
 - o p3 Q. canariensis, potential introgressant 1
 - o p2 Q. afares, potential introgressant 2
 - o p1 Q. libani [sister to Q. afares]
- B2 -- suber afares (n = 700): Q. suber with Q. afares
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. suber, potential introgressant 1
 - o p2 Q. afares, potential introgressant 2
 - o p1 undefined --- anyone who fits the topology
- C1 -- Aegilops-crenata (n = 450 tests): introgression of subsect. *Aegilops* with *Q. crenata*
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Aegilops oaks *Q. macrolepis*, *Q. brantii*, *Q. ithaburensis* as potential introgressant 1
 - o p2 Q. crenata, potential introgressant 2
 - o p1 Q. suber as sister species to Q. crenata
- C2 -- crenata afares (n = 280 tests): Q. crenata and Q. afares
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. crenata potential introgressant 1
 - o p2 Q. afares potential introgressant 2
 - o p1 undefined --- anyone who fits the topology
- C3 -- suber-cerris (n = 6400 tests): Q. suber and Q. cerris
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. suber, potential introgressant 1
 - o p2 Q. cerris, potential introgressant 2
 - o p1 undefined --- anyone who fits the topology
- D -- Libani-crenata (n = 350): subsect. *Libani* with *Q. crenata*
 - o p4 Q. chenii, Q. variabilis, and Q. acutissima as outgroups
 - o p3 Q. libani, Q. trojana, or Q. afares as potential introgressant 1
 - o p2 Q. crenata, potential introgressant 2
 - o p1 Q. suber as sister species to Q. crenata
- E1 -- cerris-afares (n = 270): *Q. cerris* with *Q. afares*
 - o p4 Aegilops oaks as outgroups: *O. macrolepis*, *O. brantii*, *O. ithaburensis*
 - o p3 Q. cerris, potential introgressant 1
 - o p2 Q. afares, potential introgressant 2
 - o p1 Q. libani 628 Q. trojana 585
- E2 -- cerris-afares | libani (n = 90): Q. cerris with Q. afares, Q. libani as p1
 - o p4 Aegilops oaks as outgroups: Q. macrolepis, Q. brantii, Q. ithaburensis
 - o p3 Q. cerris, potential introgressant 1
 - o p2 Q. afares, potential introgressant 2
 - o p1 *O. libani* 628

```
• E3 -- cerris-afares | trojana (n = 90): Q. cerris with Q. afares, Q. trojana as p1
       o p4 - Aegilops oaks as outgroups: O. macrolepis, O. brantii, O. ithaburensis
       o p3 - Q. cerris, potential introgressant 1
       o p2 - Q. afares, potential introgressant 2
       o p1 - Q. trojana 585
 E4 -- cerris-trojana (n = 90): Q. cerris with Q. trojana, Q. libani as p1
       o p4 - Aegilops oaks as outgroups - Q. macrolepis, Q. brantii, Q. ithaburensis
       o p3 - Q. cerris, potential introgressant 1
       o p2 - Q. trojana 585
       o p1 - Q. libani 628
 F -- suber-ilex (n = 6,240): Q. suber with Q. ilex
       o p4 - sect. Cyclobalanopsis as outgroup
       o p3 - Q. ilex, potential introgressant 1
       o p2 - Q. suber, potential introgressant 2
       o p1 - any species sister to p2 w/ respect in context of p4 and p3 used
  Supplemental Test 1 – crenata vs. crenata (n = 165)
       o p5 - Q. chenii, Q. variabilis, and Q. acutissima as outgroups
       o p3 1 = p4 - varies (see below)
       o p3 2 = p3 - varies (see below)
       o p2 - Q. crenata, potential introgressant 2
       o p1 - Q. suber as sister species to Q. crenata
   Supplemental Test 2 - 5-taxon D-statistic tests (n = 3)
       o p5 - O. chenii, O. variabilis, and O. acutissima as outgroups
       o p3 1 = p4 - varies (see scripts)
       o p3 2 = p3 - varies (see scripts)
       o p2 - Q. crenata potential introgressant 2
       o p1 - Q. suber as sister species to Q. crenata
```

Results: summary of D-statistic tests

D-statistic tests recover strong and consistent support for admixture between Q. cerris and Q. crenata (100% of tests significant at $p \le 0.01$; main-text $Table\ 3$). Five-taxon D-statistic tests (Supplemental Test 2) suggest that the direction of past or recent introgression has been from Q. cerris to Q. cernata, as the alleles that are shared by Q. cerris and either section Aegilops or section Libani are also significantly shared with Q. cernata relative to Q. suber (Tabulated Data S1-1). There is also strong and consistent support for admixture between Q. cerris and Q. afares when Q. libani is used as the sister to Q. afares (Test E2: 98.9% of tests significant; main-text Table 3). When Q. trojana (Italy, Balkans, Turkey) is used as the sister to Q. afares (endemic to Tunisia, N. Algeria), only the two westernmost Q. cerris individuals show significant levels of admixture: OAK-MOR-591 (Latium, Italy) and OAK-MOR-736

(Samsun, Turkey; 88.9% of tests significant; Test E3, Tabulated Data S1-1). There is minimal evidence of admixture/ introgression between *Q. ilex* and *Q. suber* (Test F: 19.0% of tests significant; Table 3). Partitioning the section *Aegilops* (p3)-*Q. crenata* (p4) test (Test C1) by individuals in the p3 position indicates that two of the *Q. ithaburensis* individuals (OAK-MOR-599, OAK-MOR-735; Levante) may have been weakly involved in introgression with *Q. crenata* (endemic to Tyrrhenian side of Italy; 61.5–88.0% of tests significant) in the past, while the other section *Aegilops* individuals (SE. Italy, Turkey, Israel) were more strongly involved (99.3–100% of tests significant; **Tabulated Data S1-1**). Partitioning the "crenata vs. crenata" test (Supplemental Test 1) by p3 individuals demonstrates that *Q. crenata* TUS13-003 (OAK-MOR-593), which falls sister to *Q. crenata* 1977-541 (OAK-MOR-986) + *Q. suber*, has been disproportionately affected by admixture from the *Cerris* core clade (**Tabulated Data S1-1**).

Tests involving *Q. afares* showed no evidence of introgression with *Q. canariensis* in any topologies (Test B1, *main-text Table 3*), but high variability in test results for *Q. suber*, with *P*-values ranging from <0.001 to 0.859 (Test B2, main-text Table 3). Subsequent tests between subsections *Aegilops* and *Suber* (C1, C2, C3) as well as subsection *Libani* and *Q. crenata* (D) partitioned tests ("cerris – afares | libani"; "cerris – afares | trojana") show that some ABBA-patterns probably represent a deeper history of introgression or other source of imbalanced shared ancestral alleles rather than recent introgression. Against the background of the FBD-dated tree (main-text Fig. 4), the fossil and niche history (following sections), the *D*-statistics point to ancestral gene flow from the precursors of subsection *Cerris* into those of subsection *Libani*, especially in the central Mediterranean region. We interpret this result as a likely outcome of introgression during the crown diversification of the Western Eurasian *Cerris*, resulting in phylogenetic discordance regarding resolution of the subsections rather than recent introgression,

However, there is evidence for more recent geographically structured introgression that bears additional study with more samples. We observe generally higher and more discriminative D-values for admixture when Q. libani is used than when Q. trojana is used as p4 in the test. The most involved Q. cerris individuals in the latter test (E3) are OAK-MOR-591 from Italy ($D \ge 0.22$; 89% of tests significant; geographically closest Q. cerris) and, to nearly the same level, the northern Turkish OAK-MOR-736 ($D \ge 0.18$; 89% of tests significant; Tabulated Data S1-1). These two individuals furthermore constitute the subsection Cerris subclade that includes also the Iranian Q. castaneifolia, a species with a distinctly primitive

(potentially ancestral within section *Cerris*) leaf morphology not unlike *Q. afares* (following section) and the widespread fossil-species *Q. kubinyii*.

The *D* estimates for *Q. crenata* differ considerably from those of *Q. afares*. We find low to high levels of admixture for all tested scenarios (*main-text Table 3*), continuously increasing along the phylogenetic tree and peaking – as in the case of *Q. afares* – in the Italian and northern Turkish individuals of *Q. cerris*. At least two reticulation scenarios might explain such a result:

Scenario 1—Quercus crenata and its precursors, representing the first diverging and westernmost section Cerris lineage (main-text Figs 2–3), were repeatedly affected by introgression when newly diverging eastern Mediterranean lineages radiated. The D-statistic results would thus reflect in this scenario a history of gene flow with other sympatric western Eurasian subsections.

Scenario 2—Alternatively, recent introgression of *Q. cerris* with sympatric *Q. crenata* in its western range may account for the observed *D*-statistics. The inferred admixture from other lineages under this scenario would reflect patterns shared by *Q. cerris* and its sister lineage, subsection *Libani* (*D*-values approaching the level as seen in southeasternmost *Q. cerris* individuals from the Levante) and the phylogenetically more distant subsection *Aegilops*.

We consider Scenario 2—modern-day introgression from *Q. cerris* into *Q. crenata*—the less likely: the partitioned *D*-statistics identified OAK-MOR-593 as the *Q. crenata* individual most affected by admixture from outside subsection *Suber*. Yet this individual exhibits relatively low root-tip distance within its clade (*main-text Fig. 1*). We would expect relatively recent introgression to introduce tree-incompatible signal into the RAD-seq data, inflating the tip branch length beyond that observed in the less-impacted second *Q. crenata* individual, OAK-MOR-986. We consequently consider either (1) ancient introgression among the precursors of modern subsections of west Eurasian *Cerris* or (2) demographic processes that result in imbalanced maintenance of ancestral polymorphisms to be a more plausible explanation for the significant *D*-statistic results.

Taxonomy and Previous Schemes

Some of the clades recovered from the RAD-seq data in the present study have previously been recognized as infra-sectional groups. Camus (1936–1938) united the East Asian members of section *Cerris* within her subsection *Campylolepides*. *Quercus suber* has previously been accommodated within its own subsection *Suber* (Menickij, 1984). Nevertheless, the majority of clades recognized here do not correspond to previously recognized infrageneric taxa (*main-text Table 2*) but agree with the 5S-IGS-defined species groups in Simeone *et al.* (2018; **Fig. S1-1**) and are here treated as subsections. The discrepancy between morphological groupings recognized previously and the clades recognized in this study on molecular grounds may represent a combination of convergence (parallelism; **Fig. S1-6**, **Table S1-1**) among clades and ecological divergence within clades.

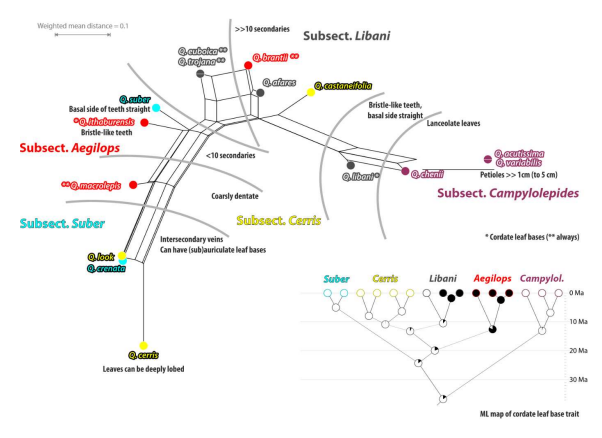


Figure S1-6. Neighbour-net splits graph inferred from pairwise (weighted) distances calculated from a leaf-morphological matrix (**Data File S1-1**; cf. **Table S1-1**). Note the poor phylogenetic sorting (subsections represent molecular clades inferred based on nuclear data) of leaf-morphological groups (species being part of the same neighbourhood) in spite of the relative tree-like graph. Bottom left: Maximum likelihood mapping of the cordate leaf base trait on the dated phylogeny. For terminology, see **Figure S1-7**.

Subsection Suber

Although the cork oak Q. suber is generally considered an evergreen tree, and therefore has been associated with section *Ilex* by Menickij (1984), it behaves differently from fully evergreen oak species. Its leaves are thinner and there is a tendency to shed most of the leaves in spring or early summer as response to early moisture stress, giving the tree a pronounced deciduous aspect (Schwarz, 1936-39; Escudero and del Arco, 1987; le Hardÿ de Beaulieu and Lamant, 2010). Menickij (1984) speculated (translation of the Russian original 2005, p. 444) "The only species of the subsection Suber – the Mediterranean Q. suber – gives the impression of an intersectional hybrid between some initial species of the preceding subsection [the subsection Heterobalanus; = sect. Ilex, the Ilex oaks] and species of the Q. aegilops group (section Cerris)." Indeed, Q. suber has a number of highly specialized ecological adaptations. Most conspicuous is the thick cork layer of the stem, which protects the trees from wildfires. Further, the sudden abscission of leaves in dry periods is an effective adaptation to seasonal dry climates, and the plastic cycle of fruit maturation may be a response to changeable availability of nutrients and water. A thinner cork layer is also produced by Q. crenata, which our analysis resolves as the only other member of the clade comprising O. suber. This cork layer has been one reason for assuming that Q. crenata was a hybrid between Q. suber and Q. cerris (Schwarz, 1936-1939; Pignatti, 1982). In sum, the now available data point to an ancestor-descendant relationship, with O. suber representing a highly adapted species that may have evolved from a Q. crenata-like ancestor in course of niching (cf. main-text Fig. 1; Supplementary Data 3; Simeone et al., 2018, figs 5–6). Plate S1-1 illustrates the ancestral-primitive (plesiomorphic) character of *Q. crenata* leaves in comparison to one of the earliest and widespread leaf fossils that can be assigned to section Cerris: O. kraskinensis Pavlyutkin. The disparate partitioned Dstatistics results (**Tabulated Data S1-2**) could result from an ancestor-descendant relationship. Being a recent and strongly bottlenecked offshoot of the subsection Suber lineage, we find no evidence for past or recent admixture in the Q. suber individuals from outside the lineage. The statistically significant D-values observed for the Q. crenata individuals may reflect retained ancestral polymorphism inherited from the subsection's precursors as well as a higher susceptibility for introgression of this less evolved and niched sibling (main-text Fig. 4, Table *3*; Tabulated Data S1-1, S1-2).

Subsection Aegilops

The 'Vallonea-oaks', subsection *Aegilops*, share several morphological similarities (**Fig. S1-6**; Avishai, 2016). The definition of the Vallonea-oaks has changed through time (*main-text Table 2*). The concept of subsect. *Aegilops* proposed here, including only *Q. brantii*, *Q. ithaburensis*, and *Q. macrolepis* (see also Schwarz, 1936), is supported both by our RAD-seq dataset and by 5S-IGS differentiation patterns (Denk and Grimm, 2010; Simeone *et al.*, 2018). The species share unique 5S-IGS variants not found in any other species of section *Cerris*. Their nuclear SNP patterns show a high coherence (well-developed, unambiguously supported root branch despite substantial tip divergence, *main-text Fig. 1*).

Cerris core group: Subsection Libani

Ørsted (1871) and later Schwarz (1936) and Menickij (1984) united the East Asian and western Eurasian fully deciduous species of section *Cerris* with a prominent mucronate leaf margin (Fig. S3-2) in the same section or subsection. Likewise, most monographers considered *Q. afares* and *Q. castaneifolia* as belonging to the same section/ subsection (*main-text Table 2*; cf. Fig. S1-6), associations that can be rejected based on nuclear data (Simeone *et al.*, 2018; this study). In our analysis, *Q. afares* is resolved in a clade together with *Q. libani* and *Q. trojana*; these three species share a lineage of 5S-IGS variants ('Oriental lineage', Cluster 1 in Simeone *et al.*, 2018; cf. Fig. S1-1) not found in subsection *Cerris* as defined below. As in the case of nuclear-encoded ribosomal spacer data (Denk and Grimm, 2010; Simeone *et al.*, 2018), we found no evidence for a white oak parentage in *Q. afares* (*main-text Table 3*; Tabulated Data S1-1).

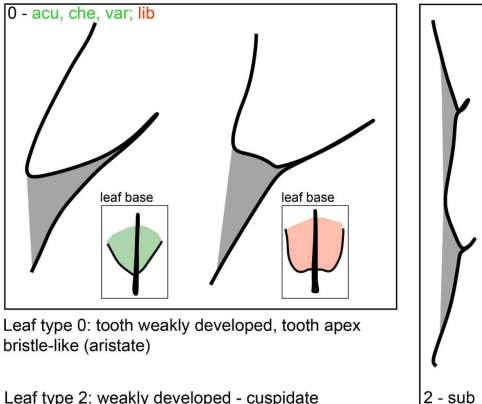
Cerris core group: Subsect. Cerris

Schwarz (1936) suggested a close relationship between the geographically isolated and ecologically narrow *Q. castaneifolia* and *Q. cerris*. Our data demonstrate that the narrow endemic *Q. look*, traditionally associated with the Vallonea-oaks (subsect. *Aegilops*), also belongs to subsection *Cerris*. As seen in the full-tip ML tree (*main-text Fig. 1*), the genetic variation found in *Q. cerris* encompasses its two endemic siblings, in striking analogy to earlier 5S-IGS data ('Occidental lineage', Cluster 2 in Simeone *et al.*, 2018). Thus, subsect. *Cerris* represents a case example for the retention of ancestral morphologies in locally isolated populations leading either to the formation of relict species (*Q. castaneifolia*) or to highly adapted (*Q. look*) Mediterranean species. Further in-depth nucleome analyses are likely to

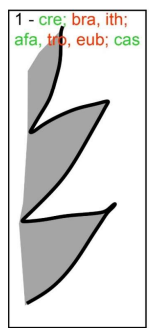
reveal cryptic or previously unrecognized species (pseudo-cryptic species) in *Q. cerris* and/or recurrent ancient hybridisation events in subsection *Cerris*, probably involving members of subsection *Libani* (5S-IGS dimorphism in *Q. trojana*; Simeone *et al.*, 2018; cf. Test E4: very low *D*-values, 66–89% of the tests statistically significant for introgression between *Q. cerris* and *Q. trojana*).

Species not yet studied

The only section *Cerris* species not included in the present study, *Q. euboica*, has recently been investigated using one chloroplast and one nuclear marker (Simeone *et al.*, 2018). Traditionally considered a subspecies of *Q. trojana*, the only species of section *Cerris* with individuals showing a prominent 5S-IGS dimorphism, *Q. euboica* appeared to be morphologically intermediate between *Q. cerris* and *Q. trojana* (**Fig. S1-6**). Thus, *Q. euboica* clearly is a member of the *Cerris* core group established here. Based on the available molecular data, its taxonomic placement remains ambiguous, belonging either to subsection *Libani* or *Cerris*, in analogy to the geographic isolates *Q. castaneifolia* and *Q. look*. Further in-depth studies are needed to assess the amount of secondary homogenisation because of sub-recent hybridisation with or without introgression by *Q. cerris*.



Leaf type 2: weakly developed - cuspidate



Leaf type 1: tooth strongly developed, tooth apex mucronate to cuspidate

Leaf types 3, 4: further increase of tooth area, subsidiary teeth, lobation (mac; loo, cer)

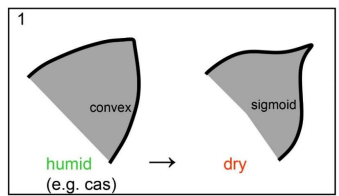


Figure S1-7. Classification of general leaf types found in section Cerris introducing the main discriminative features scored for the morphological matrix included in Data File S1-1 (cf. Table S1-1). Specific character suites (see also Fig. S1-6) were used to assign leaf fossils to clades to date the phylogeny (see main text). Species names are abbreviated by the first three letters.

Table S1-1. Leaf traits scored for the morphological matrix included in Data File S1-1.

Character	[1]	1	2	[3]	3	4	[5]	5
Species	[Leaf size	Leaf size	Leaf organisation	Secondary	Secondary veins	Intersec.	[Petiole]	Petiole
	(length, cm)]			veins		Veins		
Q. acutissima	8-19	Large	Simple	13-18	>>10	Absent	>>1cm (to 5 cm)	Long
Q. chenii	7-12	Medium	Simple	12-16	>>10	Absent	1cm (to 1.5 cm)	Short
Q. variabilis	8-15(-20)	Large	Simple	13-18	>>10	Absent	>>1cm (to 5 cm)	Long
Q. crenata	4-9	Medium	Simple	5-8	<10	Present	1-1.5 cm	Short
Q. suber	3-7	Small	Simple	5-6	<10	Absent	0.5-1 cm	Short
Q. brantii	6-10(-13)	Medium	Simple	8-14	>>10	Absent	0.5-2 cm	Short
Q. ithaburensis	4-9	Medium	Simple	5-8	<10	Absent	1-2 cm	Short
Q. macrolepis	5-9	Medium	Simple	5-8	<10	Absent	1-3.5 cm	Medium
Q. afares	7-14	Medium	Simple	8-17	>>10	Absent	1-2 cm	Short
Q. libani	7-12	Medium	Simple	11-16	>>10	Absent	0.8-2 cm	Short
Q. trojana	3-8(-10)	Medium	Simple	9-10	c.10	Absent	0.5-1 cm	Short
Q. euboica	5-9(-11)	Medium	Simple	8-10	c.10	Absent	0.5-1 cm	Short
Q. castaneifolia	10-20	Large	Simple	8-14	>>10	Absent	1-2.5 cm	Short
Q. look	5-7.5	Small	Simple	4-8	<10	Present	0.5-1.5 cm	Short
Q. cerris	5.5-14(-20)	Large	Simple to deeply lobed	5-9	<10	Present	0.3-2cm	Short

Character	6	7	8	9	10		
Species	Leaf base	Leaf shape	Leaf margin	Tooth type	Basal tooth side	Tooth type categories	
Q. acutissima	Obtuse, acute	Narrow ovate, lanceolate	Dentate	Bristle-like	Weakly developed, Straight	plesiomorph 1	0
Q. chenii	Obtuse, acute	Narrow ovate, lanceolate	Dentate	Bristle-like	Weakly developed, Straight	plesiomorph 1	0
Q. variabilis	Obtuse, acute	Narrow ovate, lanceolate	Dentate	Bristle-like	Weakly developed, Straight	plesiomorph 1	0
Q. crenata	Obtuse, acute, subauriculate	Narrow ovate, elliptic	Coarsely dentate	Coarse triangular	Convex to sigmoid	plesiomorph 2	1
Q. suber	Obtuse, acute	Ovate, elliptic	Dentate	Cuspidate	Weakly developed	derived 1 sclerophyllous	2
Q. brantii	Cordate	Ovate, elliptic	Dentate	Cuspidate	Convex to sigmoid	plesiomorph 2	1
Q. ithaburensis	Obtuse to cordate	Ovate, elliptic	Dentate	Cuspidate to bristle-like	Convex to sigmoid	plesiomorph 2	1
Q. macrolepis		•		_	-	•	
[incl. Q.							
vallonea]	Cordate	Ovate to broad ovate	Coarsely dentate	Coarse triangular	Convex	derived 2	3
Q. afares	Obtuse, acute	Narrow ovate, elliptic	Dentate	Coarse triangular to cuspidate	Convex to sigmoid	plesiomorph 2	1
Q. libani	Cordate, obtuse	Narrow ovate, elliptic, oblong	Dentate	Bristle-like	Weakly developed, Straight	plesiomorph 1	0
Q. trojana	Cordate	Narrow ovate, elliptic, oblong	Dentate	Cuspidate	Convex to sigmoid	plesiomorph 2	1
Q. euboica	Cordate	Narrow ovate, elliptic	Dentate	Cuspidate	Convex to sigmoid	plesiomorph 2	1
Q. castaneifolia	Obtuse, acute	Narrow ovate, elliptic	Coarsely dentate to dentate	Coarse triangular to cuspidate	Convex to sigmoid	plesiomorph 2	1
Q. look	Obtuse, acute, (sub)auriculate	Narrow ovate, elliptic	Coarsely dentate	Coarse triangular	Convex to sigmoid	derived 2	3
Q. cerris	Obtuse, acute, (sub)auriculate	Narrow to broad elliptic, obovate	Coarsely dentate	Coarse triangular	Convex	derived 3	4

Ad char. 10: Weakly developed = tooth area close to 0, tooth reduced to cusp or bristle-like extension (see Fig. S1-7).

Abbreviations used in legend of Fig. S1-7 (preceding page): $acu - Quercus \ acutissima$, $afa - Q. \ afares$, $bra - Q. \ brantii$, $cas - Q. \ castaneifolia$, $cer - Q. \ cerris$, $che - Q. \ chenii$, $cre - Q. \ crenata$, $cub - Q. \ euboica$, $ith - Q. \ ithaburensis$, $lib - Q. \ libani$, $loo - Q. \ look$, $mac - Q. \ macrolepis$, $sub - Q. \ suber$, $tro - Q. \ trojana$, $var - Q. \ variabilis$.

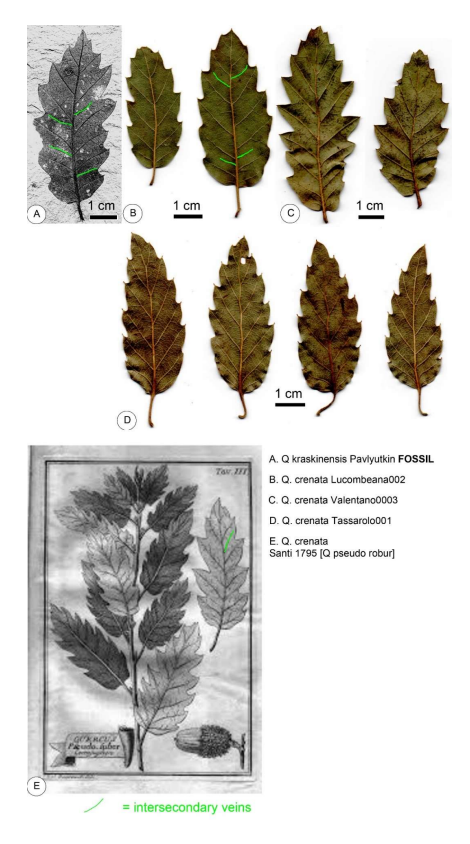


Plate S1-1. Leaf morphologies shared by modern-day *Q. crenata* and fossil-species *Q. kraskinensis* (Oligocene to Miocene, Central Asia to Europe); representing a likely symplesiomorphic leaf type of section *Cerris*.

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