Early diversification and permeable species boundaries in the Mediterranean firs

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- boundaries is critical in order to understand patterns of diversification and their historical drivers. Despite *Abies* (Pinaceae) being the second most diverse group of conifers, the evolutionary history of Circum-Mediterranean firs (CMF) remains under debate.
- Methods We used restriction site associated DNA sequencing (RAD-seq) on all proposed CMF taxa to investigate their phylogenetic relationships and taxonomic status.
- **Key Results** Based on thousands of genome-wide SNPs, we present here the first formal test of species delimitation, and the first fully resolved, complete species tree for CMF. We discovered that all previously recognized taxa in the Mediterranean should be treated as independent species, with the exception of *A. tazaotana* and *A. marocana*. An unexpectedly early pulse of speciation in the Oligocene-Miocene boundary is here documented for the group, pre-dating previous hypotheses by millions of years, revealing a complex evolutionary history encompassing both ancient and recent gene flow between distant lineages.
- **Conclusions** Our phylogenomic results contribute to shed light on conifers' diversification. Our efforts to resolve the CMF phylogenetic relationships help refine their taxonomy and our knowledge of their evolution.

Key words: species delimitation, speciation-with-gene-flow, phylogenomics, RAD-seq, conifers, transposable evolution, Abies.

INTRODUCTION

Extant gymnosperms have relatively limited species diversity presumably as a result of low speciation rates and high extinction (Crisp and Cook 2011). However, recent pulses of radiation have been found in some gymnosperm lineages, challenging the perception that gymnosperm taxa are ancient and barely variable (Davis and Schaefer 2011). Global climate shifts have been suggested as major drivers of diversification in gymnosperms radiations (Nagalingum *et al.* 2011). However, other powerful contributors to conifer evolution and diversity are the frequent hybridization events (Isoda *et al.* 2000; Ru *et al.* 2016) and bursts of transposable elements (Nystedt *et al.* 2013), which could facilitate regulatory, karyotypic or other genetic changes (Oliver *et al.*, 2013). Different transposable elements dynamics can contribute to reproductive isolation and promote species radiation (Serrato-Capuchina and Matute 2018). Delimiting species boundaries and inferring their evolutionary relationships is critical in order to understand the pace and mode of lineage formation in diversifying taxa, as well as their evolutionary drivers (Barley *et al.* 2013).

Firs (*Abies*, Pinaceae) are an important component of temperate-cool forests that characterize the boreal regions of the Northern Hemisphere (Farjon and Rushforth 1989). They represent the second most diverse group of the Pinaceae family and of gymnosperms as a whole (Xiang *et al.* 2009), consisting of more than 50 species. Although fir trees have three major areas of distribution: N America, E Asia and the Mediterranean Basin, evolutionary studies have so far primarily focused on the fir trees outside the Mediterranean Basin (Linares 2011). Based on paleogeographical data (Linares 2011), colonization and diversification of firs in the Mediterranean could have been associated to climatic and geological changes during the late Miocene and the Pliocene. Numerous

investigations have been already conducted on the genus *Abies* but included only part of the proposed taxa and only single DNA regions or low-resolving molecular markers (see below), hence species delimitation and evolutionary history of all Circum-Mediterranean firs (CMF) remain debatable.

In the last 200 years, taxonomic classification of firs has undergone numerous revisions with at least 14 formal classification attempts (Franco 1950; Liu 1971; Farjon and Rushforth 1989). The number of recognized taxa has ranged from six to 12 species and subspecies in different classifications. The most recent taxonomical review (Farjon and Rushforth 1989) classified CMF into two groups (section Abies and sect. Piceaster) and eight species. Section Abies comprises Abies alba Mill., the most widely distributed fir in Europe; A. nebrodensis (Lojac.) Mattei, a critically endangered species (conservation status: CR by IUCN) (Nebrodi mountain range, Sicily, Italy); A. cephalonica Loudon (Greece) and A. borisiiregis Mattf. (N Balkans and Greece), a hybridogenous taxa of A. alba and A. cephalonica (Bella et al. 2015; Krajmerová et al. 2016). In addition, from the Eastern Mediterranean, Farjon and Rushforth (1985) recognized A. nordmanniana (Steven) Spach (NE Caucasus and Turkey) as a single taxon. However, Abies nordmanianna is a complex taxon which other authors (Mattfeld et al. 1925; Liepelt et al. 2010; Linares 2011) have subdivided in A. equitrojani (Asch. & Sint. ex Boiss.) Mattf. (N Turkey) and A. bornmuelleriana Mattf. (N Turkey). This subdivision in only partially supported by molecular evidence (Hrivnák et al. 2017). Finally, A. cilicica (Antoine & Kotschy) Carrière, inhabiting Lebanon, Syria and S Turkey, has been included in the next section by some authors (Franco 1950; Liu 1971).

Section *Piceaster* would comprise *A. numidica* de Lannoy ex Carrière (N Algeria) and *A. pinsapo* Boiss. (S Spain and N Morocco). However, this last taxon has been subdivided into three different taxa, *A. pinsapo* Boiss *s.s.* (S Spain), *A. marocana* Trab. and *A. tazaotana*

Côzar ex Villar (both N Morocco) by several authors (Franco 1950, Arista and Talavera 1994).

Nevertheless, recent studies did not provide molecular support for the Moroccan species distinction (Sánchez-Robles *et al.* 2014, Terrab *et al.* 2007).

The inference of macroevolutionary relationships between *Abies* sections has been shown to be mostly robust to coalescent stochasticity of individual phylogenetic markers (Suyama et al. 2000; Xiang et al. 2009, 2015; Semerikova and Semerikov 2014). However, the attempts to reconstruct the phylogenetic history of CMF have yielded strongly conflicting signals between nuclear and plastidial markers (Sánchez-Robles et al. 2014; Semerikova and Semerikov 2014, 2016; Xiang et al. 2015), resulting in a largely unresolved polytomy; inter-specific relationships and divergence times hence remain to date debated. Although many phylogeographical and phenotypic studies have significantly contributed to clarify species delimitation (e.g. Fady et al. 1992; Scaltsoyiannes et al. 1999; Terrab et al. 2007; Bosela et al. 2010; Liepelt et al. 2010; Hrivnák et al. 2017), it is noteworthy that these previous studies did not include all, or even most CMF taxa. In addition, no genome-wide study has been performed to reconstruct a species tree and determine the causes of the observed molecular discordances. Disagreements among classifications of Abies are largely attributed to insufficient understanding of phylogenetic relationships within the genus, as well as its morphological complexity (Robson et al. 1993; Xiang et al. 2009). This complexity may be further exacerbated by historical gene flow between several species, especially those that are closely related and in geographical proximity (Scaltsoyiannes et al. 1999; Isoda et al. 2000; Semerikova et al. 2011; Sánchez-Robles et al. 2014; Xiang et al. 2015; Krajmerová et al. 2016). Reproductive barriers in the CMF can be permeable as suggested by successful artificial crossings, which is in contrast to the strong barriers existing between N

American and Mediterranean firs (Mergen *et al.* 1962; Alizoti *et al.* 2011; Kormutak *et al.* 2013). Secondary contacts among several CMF species during climatic cycles in the Quaternary have been previously suggested (Linares 2011; Sánchez-Robles *et al.* 2014). However, the existence of ancient gene flow during the evolution of CMF has never been tested, likely because of the weak phylogenetic signal of the previously used molecular markers.

An accurate CMF species delimitation is fundamental to manage biodiversity and forest genetic resources (Scarascia-Mugnozza et al. 2000; Fady et al. 2016; Kavaliauskas et al. 2018; Potter 2018). Due to a high degree of endemism, geographically scattered distribution and fragmentation by human activities, four CMF taxa are currently included in the IUCN red list as critically endangered or endangered (A. pinsapo "EN", A. pinsapo var. marocana "EN", A. nebrodensis "CR", A. numidica "CR"; Farjon et al. 1993; Farjon and Page 1999; Esteban et al. 2010). Hence, the clarification and accurate assessment of species status has important conservation implications (Adams et al. 2014; Melville et al. 2014). New species delimitation analyses on high-throughput data have recently allowed to identify species boundaries in several cryptic species complexes as well as in rapid adaptative radiations, e.g. in some Reptilia (Barley et al. 2013, Grummer et al. 2014), primates (Hotaling et al. 2016), and plants (Jackson et al. 2017; Brandrud et al. 2019). We applied Illumina-based restriction site associated DNA sequencing (RAD-seq) on all CMF taxa in order to investigated the evolutionary history of this group.

The main goals in the present study were: 1) to explore the origin and phylogenomic relationships of the modern CMF, 2) to test taxonomic hypotheses about the current

diversity of CMF, and 3) to investigate the existence of interspecific hybridization and the role of the transposable elements on the evolutionary history of the group.

MATERIALS AND METHODS

Taxon Sampling and Genomic Data

Leaf material from 40 individual trees (2-4 individuals per taxon) of all recognized CMF taxa (Fig. 1), together with one representative of each of the eight remaining sections of the genus *Abies* were collected at 28 natural sites using the geographic information from EUFORGEN project (http://www.euforgen.org/) during several field trips in 2015 and 2016 (Supplementary data Table S1). The samples were preserved in silica gel. Vouchers were deposited in the Herbarium of the University of Seville (SEV). Genomic DNA was extracted using a DNA extraction kit (Qiagen DNeasy Plant Mini Kit) following manufacturer's instructions. The RAD library was prepared as in Paun *et al.* (2016). Genomic DNA was digested by using high-fidelity *Sbfi* (New England Biolabs) and the resulting fragments were double barcoded. The library was sequenced in a separate lane of an Illumina flowcell HiSeq 2500 at the VBCF NGS Unit (www.vbcf.ac.at/ngs) as 100 bp single-end reads.

RADseg catalog building and SNP detection

Quality filtering and demultiplexing of the library were performed with deML (Renaud *et al.* 2015). Additionally, the *process_radtags.pl* program from STACKS ver. 1.29 (Catchen *et al.* 2011, 2013) was used to remove reads without a restriction cut site. Different empirical tests were performed with the *denovo_map.pl* program to optimize the parameters used for catalog building. We varied the minimum number of identical reads required for a stack to be formed (i.e., the setting 'm') from three to 11, the number of differences between alleles in a locus when processing one individual ('M') between one

and five, and the differences between orthologuous loci between individuals ('n') from zero to four. The final combination of settings (m=3, M=2, and n=1) was chosen to maximize the number of polymorphic RAD loci with maximum nine SNPs (to avoid pooling paralogues in the same locus) which were present in at least 30 individuals (out of 40, i.e., 75% of total). For most of the analyses we used a reduced data set, 'abies-reduced set', based on the first SNP of each RAD-locus (6,090 SNPs in total) to avoid linkage disequilibrium and ensure the independence of the loci. For most analyses, we removed the samples of the hybrid *A. borisii-regis* (*A. alba* × *A. cephalonica*, Krajmerová et al. 2016) from the data set to avoid artefacts in the phylogenomic estimates (hereafter referred to as 'abies-reduced-nohyb set'). Finally, some analyses focused on the in-group CMF only and were performed by excluding the outgroup and *A. borisii-regis* (hereafter the 'med-abies-reduced set').

Phylogenomic Analyses and Tempo of Diversification

Phylogenetic reconstruction of the genus *Abies*, based on the *abies-reduced set* and the *abies-reduced-nohyb set*, were conducted by using maximum likelihood (ML) and Bayesian inference (BI). For the ML approach we used RAXML ver. 8.0 (Stamatakis 2014) on the CIPRES Web Portal (Miller *et al.* 2015). The ASC_GTR-GAMMA model as appropriate for a dataset of concatenated SNPs and 1,000 rapid bootstrap inferences were used to estimate clade confidence. BEAST ver. 2.3.1 (Bouckaert *et al.* 2014) was used for BI and molecular dating using a GTR mutation model with gamma-distributed mutation rates across sites which provides a good approximation of the marginal likelihood at inter- and intraspecific evolutionary levels using highly variable sequences (Jia *et al.* 2014). The ucld prior was set to a uniform distribution (min. 0, max. 1), the tree prior was speciation Yule process, and the chain was 20M MCMC steps, logged every 2,000 steps. We ran four runs which were

concatenated and used to estimate the posterior distributions of topology and divergence time. In a first analysis, we dated the nodes in the genome-wide SNP-based phylogeny by using a relaxed-clock model in BEAST2 and four calibration points. The age of the oldest Abies fossil (Schorn and Wesley, 1986) being treated as a minimum age of calibration of the root, by applying a log-normal distribution of 47 Ma to the age of the oldest node. In addition, we used three Abies fossils for the calibration of nodes in the ingroup and outgroup. We used the same fossil calibrations as in Xiang et al. (2015). Fossil ages were assigned as a uniform distribution for their stratigraphic range (Supplementary data Table S2). The minimum age of the stems of Abies and CMF were assigned using lognormal prior distributions (47 Ma and 23 Ma, respectively), meanwhile uniform prior distributions (a minimum of 16 Ma and a maximum of 47 Ma) were assigned to the crown node of the clade including sects. Grandes and Oiamel and the crown node of sect. Momi. In a second independent dating analysis, we used two biogeographical calibration points (six calibration points in total; Supplementary data Table S2). We used the date of the separation of N Africa from Europe (i.e., the end of the Messinian Salinity Crisis, the opening of the Strait of Gibraltar and Tyrrhenian Sea; ~6 Ma, Normal prior distribution) as a calibration age of the divergence between A. pinsapo and A. marocana - A. tazaotana but also for divergence between A. nebrodensis and A. alba.

Species delimitation

We compared eight candidate species delimitation models (Fig. 2) by using the Bayes factor delimitation of species (SNAPP-BFD*, Leaché *et al.* 2014) as implemented in SNAPP (Bryant *et al.* 2012). BEAUti v2.3.1 (Bouckaert *et al.* 2014) was used to set up the analysis file. The null model followed the most recent taxonomic treatment, but considering

subspecies as species (11 species in total; Delimitation Species Hypothesis A). Based on previous studies and our phylogenomic results, seven additional hypotheses were tested (Fig. 2). Specifically, we tested that A. marocana and A. tazaotana, which are locally restricted to Mount Tazaot, form together a single species and that their current distributions are just the result of recent fragmentation (hypothesis B; Sánchez-Robles et al. 2014). In addition, we tested the inclusion of the restricted A. nebrodensis in the widespread species A. alba (hypothesis C). We also tested a potential clustering together of the Turkish taxa (A. nordmanniana, A. bornmuelleriana and A. equi-trojani) as a single species (hypothesis D). These three taxa have been treated with different taxonomic status in several taxonomic revisions (Coode and Cullen 1965; Farjon and Rushforth 1989). Finally, we also tested other combinations of these species delimitations (hypothesis E-I). For these analyses, we used the med-abies-reduced set without any missing data. We analysed the alternative tree topologies by using 24 steps of path sampling with 100,000 Markov Chain Monte Carlo (MCMC) steps and 10,000 pre-burn-in steps to estimate marginal likelihoods. Convergence of the parameters (ESS > 200) was checked using Tracer v1.7.1 (Rambaut et al. 2018) after 10% burn-in removal. Estimated Marginal Likelihoods for competing models were compared by using Bayes Factors (2 \times loglikelihood difference) to identify the most likely species delimitation. The species tree was subsequently inferred by using the Bayesian multispecies coalescent approach implemented in SNAPP with the same previous parameters (Bryant et al. 2012) and visualized with Densitree 2.01 (Bouckaert, 2010). In addition, we used SVDQuartets (Chifman and Kubatko, 2014) with 100 nonparametric bootstrap replicates in PAUP* version 4.0a147 (Swofford, 2002) to generate a species tree for the best species delimitation.

We also produced a lineage-through-time (LTT) plot, which displays the number of speciation events through time, taking into account possible extinction events. We calibrated the species trees from SNAPP by enforcing a strict molecular clock using the 'chronos' function in the R library *ape* (Paradis *et al.* 2004). As calibration point, we used the 95% Highest Posterior Density node age of the CMF inferred from the previous genomewide SNP-based BEAST2 analysis. Additionally, we used the gamma statistic (y) from chronograms (Pybus and Harvey 2000) to determine whether a signal of decreased diversification was noticeable.

Genetic structure and recent admixture

To assess the genetic structure of CMF, we performed a Bayesian Markov Chain Monte Carlo (MCMC) approach with STRUCTURE ver. 2.3.4 (Pritchard *et al.* 2000). Ten independent runs were carried out for the data set for values of *K* ranging from one to 13. The number of iterations was 500,000 with a burn-in of 100,000 steps. STRUCTURE analyses were performed for all the *Abies* species in the Mediterranean area, as well as separately for each section (sect. *Abies* and sect. *Piceaster*, with K ranging from one to ten and one to eight, respectively). STRUCTURE HARVESTER software (Earl and VonHoldt 2011) was used to determine the optimal number of clusters (*K*) implementing the Evanno method. The graphical representation of STRUCTURE results was generated by using CLUMPAK (Kopelman *et al.* 2015). Additionally, fineRADstructure and RADpainter ver. 0.1 (Malinsky *et al.* 2018) were used to infer the shared ancestry among individuals. The estimated haplotype coancestry matrix was used to infer recent admixture and to build a tree by using default parameters. For this analysis, we removed, as recommended, a sample with more missing data (*A. equi-trojani_94_1*) than the rest.

As popular approaches such as STRUCTURE (Pritchard et al. 2000) and ADMIXTURE (Alexander et al. 2009) are only appropriate for detecting gene flow over relatively short evolutionary time scales and may have limited power when only few individuals per population are available (Soraggi et al., 2018), we applied an ABBA/BABA approach (Green et al. 2010) to the SNP matrix. This method distinguishes the contribution of incompletelineage-sorting from introgression by calculating imbalances in alternative tree topology frequencies under a four-taxon scenario (three ingroup taxa and one outgroup). In short, for a particular SNP with an ancestral ("A") and derived ("B") alleles, the alternative patterns "ABBA" and "BABA" should be equally frequent under a scenario of incomplete lineage sorting without gene flow. D-statistics and Z-scores calculations were performed with the R package evobiR (Blackmon and Adams 2015), which measures signals of alternative phylogenetic asymmetry and the proportion of the genome that is shared between two taxa due to admixture, respectively (Durand et al. 2011). ABBA/BABA tests were performed on the abies-reduced set including heterozygous sites by generating pseudohaploid sequences (Green et al. 2010). One thousand bootstrap iterations were used to measure the standard deviation of the *D*-statistic (Eaton and Ree 2013). Based on previous studies (Sánchez-Robles et al. 2014) and our STRUCTURE results we investigated gene flow among A. alba (sect. Abies) and A. numidica and A. pinsapo (sect. Piceaster). Using the approach proposed by Martin et al. (2013) and the previously estimated species relationships, we performed several ABBA-BABA tests with different candidate species to distinguish between admixture in different time periods. Briefly, if gene flow was only ancient (e.g. previous to A. nebrodensis divergence from A. alba), then A. alba and A. cilicica should both be equally admixed with *A. numidica* and *A. pinsapo*. However, if gene flow was more recent (i.e., after *A. nebrodensis* divergence), then *A. alba* should be more admixed with *A. numidica* and *A. pinsapo* than with *A. cilicica*. We performed analyses for all combinations of pairs of conspecific individuals from candidate admixers (*A. alba, A. nebrodensis, A. cilicica, A. numidica* and *A. pinsapo*) using *A. lasiocarpa* as outgroup.

Genomic divergence in CMF

Additionally, to infer transposable elements (TEs) contribution to genomic divergence of CMF species, the presence of TEs was checked in raw reads using the approach proposed by Trucchi *et al.* (2017). We randomly extracted 100,000 single-reads from each individual, which were inspected for transposable elements by using RepeatMasker ver. 4.0.6 against the RepBase TE database (Kohany *et al.* 2006) of Viridiplantae ver. 2016-08-29. Each individual sample was analysed independently and the relative abundances (i.e., TEs per 100,000 reads) of different TEs (Gipsy, Copia, LINEs and DNA transposons) were compared between sections by using Poisson-based GLMs in the software R.

RESULTS

The 100-bp Illumina RAD sequencing for 40 DNA samples (1.7M reads on average per sample) representing virtually all recognized CMF taxa (Fig. 1, Supplementary data Text S1, Supplementary data Table S3) produced in the final STACKS catalog a total of 1,585,024 RAD-loci with an average coverage per sample (\pm SD) of 81.3 \pm 25.9 reads per locus. The demultiplexed data have been deposited in the NCBI Short Reads Archive (BioProject ID PRJNA563575). We found a high variability in the percentage of missing RAD-loci among samples, ranging from 4.9% to 61.2% (mean \pm SD 12.84% \pm 10.24; Supplementary data Table S3) but this pattern was not biased towards particular taxa ($F_{19,20} = 1.593$; P = 0.155). After

retaining only polymorphic RAD-loci that were present in at least 30 individuals and had maximum nine SNPs per locus, we obtained a final data set with 6,090 RAD-loci and a total of 23,931 SNPs. However, for further analyses we retained only one SNP per RAD-locus (6,090 unlinked SNPs in total).

Genome-based phylogeny and species delimitation

The ML and BI resulted in robust and generally consistent topologies with two highly supported (BS = 1.0 and PP = 99) lineages within the CMF (Fig. 2a). These main lineages corresponded to the two previously-recognized sections in the Mediterranean, sect. *Abies* and sect. *Piceaster*. In the *Piceaster* lineage, *A. numidica* (occurring in Algeria) is an early-diverged clade (BS = 1.0 and PP = 99), whereas *A. pinsapo* (Spain) and *A. tazaotana* + *A. marocana* (NW Morocco) are sister clades. Individuals of *A. tazaotana* and *A. marocana* appeared intermixed. In the *Abies* section, two highly supported clades were shown. The first one included *A. alba* and *A. nebrodensis*. The second clade included species from Greece, *A. cephalonica* and an inner clade of Turkish taxa, *A. cilicica*, *A. nordmanniana*, *A. bornmuelleriana* and *A. equi-trojani*. As expected in a bifurcating tree, the individuals of the hybrid taxon *A. borisii-regis* clustered with either parental species, *A. alba* and *A. cephalonica* (Supplementary data Fig. S1).

For the Bayesian Species Delimitation analysis in SNAPP, the species hypothesis with the highest likelihood suggested that all previously recognized taxa (either at the species or at the subspecies level) in the Mediterranean should be treated as independent species, with the exception of *A. tazaotana* and *A. marocana* (MLE = -6500.7 for distinct, hypothesis A; MLE = -6440.5 for the two taxa grouped as one species, hypothesis B; Bayes Factor value = 120.4 representing decisive evidence, Supplementary data Table S4).

Origin and evolution of Mediterranean firs

The topology of the species tree constructed with SNAPP, based on the best hypothesis according to Bayesian Factor Delimitation of Species (BFD*), was mostly congruent with the concatenated-SNPs trees (Fig. 2), but the main lineages evolved comparatively early. The *Piceaster* lineage (including *A. numidica*, *A. pinsapo* and *A. marocana-A. tazaotana*) showed the same internal relationships, whereas the Turkish lineages (including *A. nordmanniana*, *A. bornmuelleriana* and *A. equi-trojani*) diverged rapidly from one another. Additional support to the CMF relationships came from the species tree inferred using SVDquartets, whose topology was highly supported (Supplementary data Fig. S2). This species tree, again, supported the *Abies* and the *Piceaster* sections as monophyletic, with 100% BS. Whereas the SVDquartets tree was largely congruent with inferences from concatenated data sets, the species tree differed in some relationships, especially for deep divergence events.

Dating the diversification of CMF is key to understand the putative contribution of climatic fluctuations and geological events around the Mediterranean Basin to the evolution of CMF. The estimated divergence times using BEAST and several fossil records are shown in Fig. 2. The chronogram was congruent with an early divergence of the lineages of sect. *Abies* and sect. *Piceaster* around the late Oligocene (95% Highest Posterior Density estimates; HPD: 23.1–34.1 Ma). Miocene divergence times for all current CMF lineages are clearly inferred, with speciation times pre-dating by tens of millions of years the Messinian Salinity Crisis (5.59-5.33 Ma; Krijgsman *et al.* 1999) and the Mediterranean climate establishment (~3.2 Ma, Suc 1984). For the *Piceaster* lineage, the split of *A. numidica* and *A. pinsapo-A. marocana-A. tazaotana* was estimated at 13.3–27.1 Ma and at 9.4–21.3 Ma for that of *A. pinsapo* and *A. marocana-A. tazaotana*. For the lineage of sect. *Abies*, an Oligocene-

Miocene split of *A. alba* and the eastern lineage was estimated (18.4–31.1 Ma). The eastern lineage diverged in the early Miocene, with 15.1–27.6 Ma for the split of *A. cilicica* and 12.2–24.0 Ma for that of *A. cephalonica*. The Turkish species were the last to diverge, around 8.4–18.5 Ma. Additionally, the speciation between *A. nebrodensis* and *A. alba* occurred 9.1–21.8 Ma. These estimates were robust to additional calibration points derived from geological events, such as the opening of the Strait of Gibraltar and Tyrrhenian Sea after the Messinian Salinity Crisis (Supplementary data Table S5). Nevertheless, this secondary dating analysis (including geological events) invariably showed shorter ranges for the Highest Posterior Density estimates. Additional dating analyses using other tree prior speciation processes (i.e. Birth-Death and Bayesian Skyline; not shown) provided similar estimates.

We produced a lineage-through-time (LTT) plot, which displays the number of speciation events through time, taking into account extinction events (Fig. 2). The CMF species have accumulated early and quickly in the history of the group. The gamma statistic was further employed to explore the acceleration of speciation rate over the history of CMF. The distribution of the gamma values on the 2,276 SNAPP species trees rejects a constant speciation rate; γ (mean \pm SD) = -1.61 \pm 0.38; p-values < 0.05 (Fig. 2). The negative gamma values revealed an early and rapid accumulation of branching times near the root, indicative of a decelerated diversification. The shift in the number of speciation events occurs quickly in the group, with the highest number of speciation events estimated at the boundary between the Oligocene and the Miocene. The rapid divergence and expansion in the Tertiary and a relative speciation stasis in the Quaternary were congruent with the lack of differences in *Abies* fossil records along the latitude (t-test, t = -0.384, d.f. = 85.268, P = 0.702) or longitude (t = 1.299, d.f. = 90.376, P = 0.197) between these two epochs (Supplementary data Fig. S3).

Genomic divergence between fir species and evidence for ancient hybridization during the speciation process

The rapid CMF radiation was congruent with low genomic divergence among species. More than half of the SNPs (51.8%) represented shared polymorphism between at least two CMF species. Heterozygosity was similar between species, ranging from 0.04 to 0.08 with an average of 0.055 \pm 0.002 ($\chi^2_{11,20}$ = 16.13, P = 0.134). However, the *Piceaster* section showed a significantly lower heterozygosity (0.049 \pm 0.005) than *Abies* (0.058 \pm 0.003; $\chi^2_{1,20}$ = 6.35, P = 0.01; Supplementary data Fig. S4). The pairwise F_{st} values between species ranged from 0.04 to 0.53 (Supplementary data Table S6).

The Bayesian assignment of genetic structure (Fig. 3) showed, again, two main clusters (i.e. best partition K=2 following Evanno method; LK= 50984.8, ΔK = 789.2, Supplementary data Fig. S5a) corresponding to the two taxonomic sections. Although most samples showed a high assignment probability to just one section, admixture signals appeared in several individuals (Fig. 3) of A. numidica and A. pinsapo, which showed intermediate probability to belonging to sect. Abies. Additional suboptimal groupings were K=5 and K=7 (Supplementary data Fig. S5a) which did not show clear taxonomical patterns. Within-sections, the genetic structure generally mirrored the species tree (Fig. 3). For sect Piceaster, K=3 was the best partition (Supplementary data Fig. S5b) corresponding to A. numidica, A. pinsapo and A. marocanna+A.tazaotana. For sect. Abies, K=5 partition was selected (Supplementary data Fig. S5c). The first group corresponded to A. cephalonica individuals, the second one included A. cilicica individuals. Abies alba and A. nebrodensis supported together a third cluster, whereas the last group encompassed A. nordmanniana, A. equi-trojani and A. bornmuelleriana. High haplotype coancestry levels among the accessions of Abies section Piceaster were estimated with RADpainter (Fig. 3). In addition,

and consistently with recent hybridization between the species of both sections, we observed high levels of haplotype coancestry among distantly-related species (e.g. *A. numidica* and *A. alba*) as estimated with RADpainter (Fig. 3).

Additionally, we tested if ancestral gene-flow was also compatible with these admixture signals. ABBA-BABA tests (Patterson's *D*-statistic tests; Fig. 4, Supplementary data Table S7) supported an ancient divergence-with-gene-flow among species from the two CMF sections. Significant negative D-statistics supports a notable ancient gene flow between *A. alba* and two species of sect *Piceaster*, *A. numidica* and *A. pinsapo* but also between *A. cilicica* and the sect *Piceaster* species. In addition, higher absolute admixed values (Z-statistics) were found for comparisons including *A. nebrodensis* (as opposed to *A. cilicica* comparisons) indicating additional, recent admixture events. Remarkably, *A. alba* showed higher admixture with *A. pinsapo* than with *A. numidica*.

Furthermore, a significantly divergence between genomic landscapes of the two *Abies* sections was also mediated by differences in transposable elements activity. We found significant differences between both *Abies* sections (Fig. 5) in the relative amount of DNA transposons (χ^2 = 27.5, P < 0.001), Copia (χ^2 = 324.5, P < 0.001), Gipsy (χ^2 = 447.5, P < 0.001) and LINE (χ^2 = 3.3, P < 0.1) within our RADseq data. Interestingly, the relative abundances of some TE classes showed also a strong correlation with the latitude of the samples (Supplementary data Fig. S6).

DISCUSSION

To our knowledge, this research represents the first genome-wide evolutionary study in firs and the most intensive sampling including all CMF taxa. Although some limitation of RADseq for outlier scans has been suggested and debated (Tiffin and Ross-Ibarra 2014; Lowry et al.

2016, but see Catchen *et al.* 2017), this study highlights the advantages of applying RAD-seq to obtain SNPs for phylogenetic studies in taxa with giant genomes, like in the genus *Abies*. Furthermore, the combined use of phylogenetic methods and population genomic methods (such as STRUCTURE) have contributed to reveal the complex evolutionary history of Mediterranean firs clarifying not just their phylogenetic relationships but also the recent and old secondary contacts.

Phylogeny of the Mediterranean Abies

Our dated phylogenomic tree corroborated the monophyly of all Mediterranean firs, as previously suggested (Suyama et al. 2000; Semerikova and Semerikov 2014, 2016; Xiang et al. 2015). However, our molecular analyses, for the first time, clearly separated CMF into the two previously classified sections, Abies and Piceaster (Farjon and Rushforth 1989). Previous studies using classical markers have been unable to clarify the phylogenetic relationships of these taxa. The monophyly of the Piceaster section, including the SW species A. numidica, A. marocana, A. tazaotana and A. pinsapo, was additionally supported by other molecular markers (Ziegenhagen et al. 2005; Sánchez-Robles et al. 2014). However, the taxonomic status of the Moroccan species has been controversial; sometimes they were considered as mere varieties of A. pinsapo (Farjon and Rushforth 1989), sometimes as subspecies (Govaerts 1995) or distinct species (Maire 1952). The species delimitation analysis suggested that the Moroccan Abies species must be classified as different species from A. pinsapo, but that they should not be split into independent species. This result is congruent with the genetic distinctiveness of A. pinsapo from A. marocana - A. tazaotana and the lack of genetic structure among populations of the latter by using AFLP and plastid markers (Terrab et al. 2007; Sánchez-Robles et al. 2014). Both Moroccan taxa were

previously separated (Sánchez Cózar 1946) on the basis of morphological characters (mainly leaves, cones and bracts), but because of their close genetic similarity, and based on finer morphological studies (Sekiewicz *et al.* 2013) as well, we propose to redefine their taxonomical status as a single species. Based on these results we update the current taxonomy of CMF (Text S1).

In the Abies section, the results from the phylogenetic reconstructions and the species delimitation hypothesis test support the identity of A. nebrodensis, endemic to the Madonie Range of N Sicily (Italy) as a species derived from an ancestor of A. alba, the most widely distributed species inhabiting central Europe. This taxonomical status is supported by differences in traditional nuclear and plastid markers (Vicario et al. 1995; Parducci and Szmidt 1999; Parducci, AE Szmidt, et al. 2001a) and refute its classification as a subspecies or a variety of A. alba or its alleged hybrid origin from A. alba and A. numidica (Landry 1984; Parducci, A Szmidt, et al. 2001b). A close relationship between these two latter species from different sections was previously suggested by Sánchez-Robles et al. (2014) based on AFLP markers but our results refute this hypothesis. Furthermore, the relationships between A. alba, A. cilicica and A. cephalonica have been resolved here for the first time. The two latter species (plus the A. nordmanniana complex clade) formed a deeply diverged sister lineage to A. alba. In this clade, an early divergence of A. cilicica firstly occurred, whereas the Greek A. cephalonica and the E Turkish clade diverged later. Although Abies cilicica was previously wrongly assigned to the Piceaster section (Franco 1950; Liu 1971), our data rather support a phylogenetic relationship with sect. Abies. In addition, the monophyly of the Greek A. cephalonica and the E Turkish Abies species (i.e. the A. nordmanniana complex) was previously highlighted by their shared mitochondrial haplotype (Ziegenhagen et al. 2005). Their genetic distinctiveness was further visible in our STRUCTURE results. The A.

nordmanniana complex formed a single cluster, but the BFD* analysis suggested the existence of three different species: A. nordmanniana s.s, A. equi-trojani and A. bormulleriana. A recent study (Hrivnák et al. 2017) on the A. nordmanniana complex resulted in partially congruent phylogenetic relationships with our species delimitation hypothesis. However, the differentiation between the sister species A. equi-trojani and A. bormulleriana was unclear (Hrivnák et al. 2017). Finally, the clarification of CMF taxonomy has predictable positive effects on conservation. New CMF classification could help authorities to focus conservation efforts (especially for the Moroccan and Turkish firs).

Origin and evolution of the Mediterranean firs

The information on the phylogenetic relationships uncovered in this study, combined with our knowledge of the geographical distribution of the species and the fossil record, can provide insights on the origin and evolution of Mediterranean firs. In short, the dated monophyletic origin of the Mediterranean firs and their close relationships with eastern Asian sections Momi and Pseudopicea (Xiang et al. 2015) would suggest a single colonization event by a CMF ancestor from Asia in the late Eocene-Oligocene. This scenario is consistent with the estimated date for the Mediterranean crown group in previous studies on the genus *Abies* (Aguirre-Planter et al. 2012; Xiang et al. 2015). It is also noteworthy that the global expansion and diversification of firs have been dated during the Eocene as well, in agreement with dispersed fossils all over the N Hemisphere (Xiang et al. 2007, 2015). The global climate cooling down through the Eocene-Oligocene (Zachos et al. 2008) period could have favoured the geographical expansion of firs. In relatively short time, as supported by the fossil records (Cavagnetto and Anadón 1996), the CMF ancestor could have reached the westernmost region of the Mediterranean Basin (i.e., the Iberian Peninsula).

With regard to the speciation sequence in the Mediterranean firs, a previous hypothesis suggests a slow diversification along the Miocene-Pliocene, culminating with the rise of the modern CMF species after the Messinian Climate Crisis and the Pliocene marine transgression (Linares 2011). Contrary to expectations, our conservative molecular dating and the speciation rate analyses suggest a much earlier diversification of the Mediterranean firs in the late Oligocene-Early Miocene which is congruent with the Mesoamerican firs radiation (Aguirre-Planter et al. 2012). The finding of rapid diversification in CMF is particularly striking given that it concerns one of the most important components of temperate-cool forests in Eurasia and it has been widely studied. Our molecular dating method was carefully designed to incorporate the uncertainty associated with paleontological data and it was robust to different settings (including some biogeographical events). Understanding why CMF have diversified during Miocene requires a reconsideration of paradigms concerning patterns of diversification in the Mediterranean basin as previously suggested by Vargas et al. (2018). Quaternary climatic cycles have been evoked as main factor of diversification but the new evidence suggests a secondary role providing secondary contacts and posterior isolation.

Our reconstruction of past evolutionary events suggests that during late Oligocene-Early Miocene, the archaic Mediterranean *Abies* would have split into two groups, corresponding to the two sections *Piceaster* and *Abies*, perhaps in each extreme of the Mediterranean Basin. In the western region, the ancestor of *A. numidica*, *A. pinsapo* and *A. marocana* (i.e., sect. *Piceaster*) would remain genetically isolated and in the central-east region would have differentiated the archaic *Abies* sect. *Abies* species. Additionally, the latter species would have quickly split into two lineages, one for proto-*Abies alba* and another one for the ancestor of all eastern species.

Additionally, our LTT plot and gamma statistics analysis showed signals of a rapid speciation burst shortly after the CMF expansion in the early Miocene. These results predated by tens of million years the previous hypothesis about CMF diversification. Interestingly, the divergence between Ephedra and disjunct Cedrus species in the Mediterranean has been estimated to the Miocene (Qiao et al. 2007; Ickert-Bond et al. 2009) as well. Provided CMF, Cedrus and Ephedra species share a similar biogeographic history in the Mediterranean region, our study supports the idea that living Mediterranean gymnosperms are not relicts and they have originated by an early Neogene pulse of diversification (Davis and Schaefer 2011) together with many Mediterranean angiosperms (Vargas et al. 2018). The likely reason behind this pulse of speciation in the Mediterranean flora could be attributed to the global climate change during the Miocene. After the temperature and rainfall decreased during the Eocene-Oligocene, temperature rose slightly through the Neogene until reaching in Miocene a climatic optimum (15-17 Ma; Zachos et al. 2001). The elevated temperature connected to a global CO₂ increase (Zachos et al. 2008) could have aggravated the effects of increased aridity and caused extinctions and geographic isolation in the CMF (Rundel et al. 2016).

Furthermore, we found differential dynamics in TE between the two sections which contrast with the slow and steady rate of accumulation of TE found otherwise in gymnosperms (Nystedt *et al.* 2013). However, a recent study (Voronova et al, 2017) suggests some retrotransposon elements have been established, diverged and expanded after separation of Pinaceae family. In spite of the genome similarity between firs species, differences in TEs could have profound effects on gene expression (Bennetzen and Wang 2014) and possibly might contribute to adaptative radiation and morphological diversity in CMF. The role of climatic stress in the activation of TE has been widely assessed (Capy *et al.*

2000) and may additionally support the climatic role in the speciation of CMF (Kaplan and Guy 2005; Krasensky and Jonak 2012; George *et al.* 2015; Sánchez-Salguero *et al.* 2017).

Evidence of early divergence-with-gene-flow and secondary contacts in the Mediterranean firs

Generally, our study revealed signs of admixture between CMF species of both sections but also within-sections, causing potential conflicts between gene trees, but also when constructing a bifurcating species tree. In spite of the morphological differentiation between CMF species, a lack of reproductive isolation was evidenced by artificial interspecific crossing experiments (Mergen et al. 1962; Kormutak et al. 2013). The multiple locations of A. borisii-regis in the multiple phylogenomic analysis and the low pairwise F_{ST} values support its recent hybridogenous origin due to multiple introgression events between A. alba and A. cephalonica. This pattern of genomic variation is consistent with a previously found genomic cline (Krajmerová et al. 2016) and suggests that A. borisii-regis is an unstabilized hybridogenous taxon. Furthermore, the existence of recent gene flow between other CMF species was also apparent from multiple lines of evidence. The signals of admixture in the STRUCTURE analysis and the high haplotype coancestry are congruent with other previously-suggested secondary contacts, most likely during the Pliocene-Pleistocene, between A. alba, A. pinsapo and A. numidica (Parducci, AE Szmidt, et al. 2001a; Sánchez-Robles et al. 2014). Based on cpSSR analysis, Sánchez-Robles et al. (2014) suggested ancient secondary contacts among these species as no recent admixture signals were found using AFLP markers. In our study, we uncovered the importance of this ancient gene flow among species from the two main lineages in the evolution of the CMF. The process of fast speciation in CMF involves alternate periods of partial isolation and

secondary contacts among divergent species as a likely consequence of climatic oscillations.

CONCLUSIONS

In summary, our modern phylogenomic approach sheds light on the diversification of the Circum-Mediterranean firs. Our efforts to resolve the phylogenetic relationships among the different lineages have refined their taxonomy and our knowledge of their evolution. An unexpectedly early pulse of speciation in the Oligocene-Miocene boundary is here documented, pre-dating by millions of years previous hypotheses, revealing a complex evolutionary history encompassing both ancient and more recent gene flow between distant lineages. Different transposable elements dynamics could have contributed to the genome divergence of Circum-Mediterranean firs.

SUPPLEMENTARY DATA

Supplementary data are available online at https://academic.oup.com/aob and consist of the following. Text S1: Taxonomic Revision. Table S1: Population details of the samples included in the study. Table S2: Firs fossil calibrations and geological events used in this study. Table S3: SNP screening by using the STACKS software for all the studied individuals. Table S4: Marginal Likelihoods and Bayes Factor values for the alternative species delimitation hypothesis. Table S5: Mean ages and 95% HPD heights for the major nodes of the CMF chronogram using two additional geographic events for calibration. Table S6: Circum-Mediterraneam *Abies* pairwise FST estimates from 6,090 SNPs. Table S7: Details of individual ABBA-BABA tests for ancient gene flow among *Abies* species from sections Abies and Piceaster. Fig. S1: Maximum Likelihood phylogenetic tree of CMF including the hybrid *A. borisii-regis*, based on 6,090 SNPs. Fig. S2: SVD-Quartet species tree. Fig. S3: Geographic pattern of *Abies* fossil records along latitude and longitude in the Tertiary and the

Quaternary periods. Fig. S4: Violin plots of heterozygosity per individual of the two Mediterranean *Abies* sections. Fig. S5: Summarized results from STRUCTURE. Fig. S6: Relationship between DNA transposons and Latitude.

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FIGURE CAPTIONS

- FIG. 1. Sampling map for the Mediterranean firs used in this study. Each population is labelled and species ranges are highlighted with different colours (more information in Supplementary data Table S1). Species distributions are based on the EUFORGEN project (http://www.euforgen.org/) and personal observations.
- FIG. 2. Phylogenetic reconstructions for the CMF based on 6,090 SNPs. (A) Chronogram built in BEAST2 using 6 fossil calibration points (asterisks). Posterior probabilities and bootstrap supports from RAxML are colour marked according to legend. (B) Species-delimitation hypotheses tested in the BFD* analysis. Each grey cell represents an independent species. Other colours indicate taxa that are merged into a single species.
- **FIG 3**. SNAPP species cladogram for the best species-delimitation hypothesis (A). Lineage through time plot and distribution of γ -statistics for the SNAPP trees (B).
- FIG. 4. Different types of evidence of recent gene flow in the CMF. Hierarchical structure among and between CMF samples estimated by fineRADstructure. Results of the Bayesian assignment analysis (STRUCTURE) for CMF (and independently for each section). Haplotype coancestry matrix based on 6090 SNPs with individual *A. equitrojani_94_1* removed due to excessive missing data. The colour in the heat map corresponds to the number of haplotypes imported from a donor genome to a recipient genome. High levels of haplotype coancestry among distantly-related species (e.g. *A. numidica* and *A. alba*) suggest secondary contacts.
- **FIG. 5.** Hypothetical schemes of introgression between species of *Abies* sect *Abies* and *A.* sect *Piceaster* at different time periods. Violin plots show the distribution of the *D* statistics (ABBA-BABA tests) resulting from testing different individuals from each

species. Detailed information about the significance of the *D*-statistics for each test is provided in Supplementary data Table S7. Values different from zero denote introgression between species. ALB *Abies alba*, CIL *A. cilicica*, NEB *A. nebrodensis*, NUM *A. numidica* and PIN *A. pinsapo*. *Abies lasiocarpa* was used as outgroup.

FIG. 6. Violin plots of the relative abundance of different transposable elements families per individual of the two Mediterranean *Abies* sections. Asterisks denote statistically significant differences at P < 0.001 (Poisson Generalized Linear Model).

Figure 1

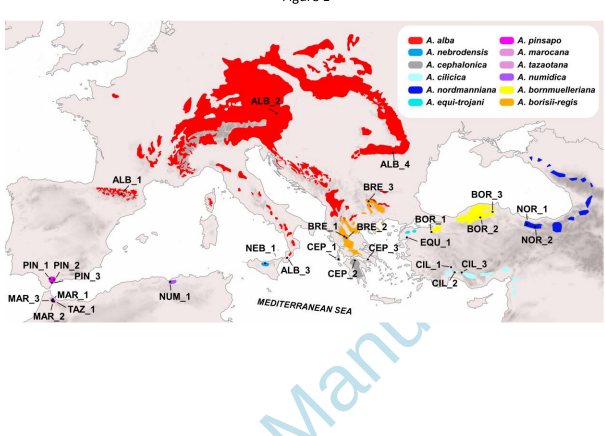


Figure 2

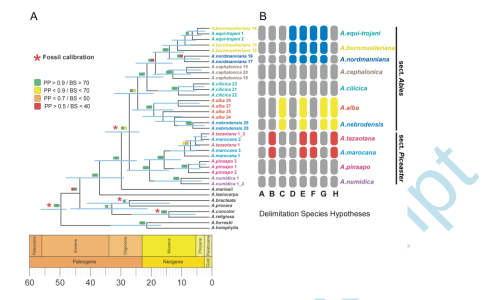


Figure 3



Figure 4

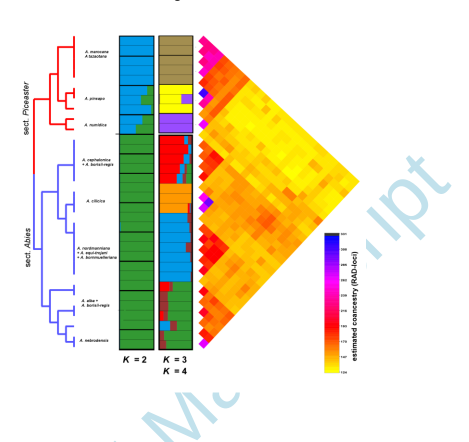


Figure 5

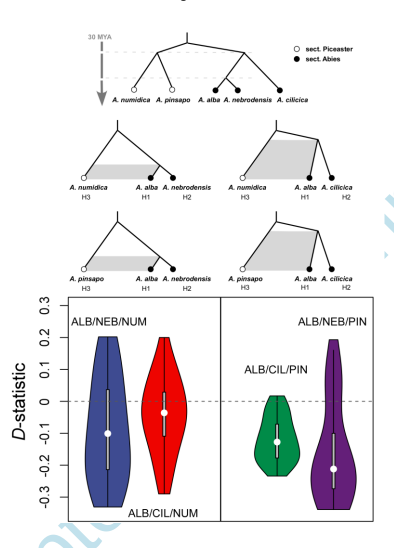


Figure 6

