

Adapting through glacial cycles: insights from a long-lived tree (*Taxus baccata*)

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Summary

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- Despite the large body of research devoted to understanding the role of Quaternary glacial cycles in the genetic divergence of European trees, the differential contribution of geographic isolation and/or environmental adaptation in creating population genetic divergence remains unexplored. In this study, we used a long-lived tree (*Taxus baccata*) as a model species to investigate the impact of Quaternary climatic changes on genetic diversity via neutral (isolation-by-distance) and selective (isolation-by-adaptation) processes.
- We applied approximate Bayesian computation to genetic data to infer its demographic history, and combined this information with past and present climatic data to assess the role of environment and geography in the observed patterns of genetic structure.
- We found evidence that yew colonized Europe from the East, and that European samples diverged into two groups (*Western*, *Eastern*) at the beginning of the Quaternary glaciations, c. 2.2 Myr before present. Apart from the expected effects of geographical isolation during glacials, we discovered a significant role of environmental adaptation during interglacials at the origin of genetic divergence between both groups.
- This process may be common in other organisms, providing new research lines to explore the effect of Quaternary climatic factors on present-day patterns of genetic diversity.

Introduction

It is currently accepted that Quaternary climatic oscillations have played a major role in shaping the geographical distribution of European species and their patterns of genetic structure (Hewitt, 2004). In the particular case of temperate taxa, geographical isolation and long-term persistence in southern refugia during the glacial episodes have been considered essential for population divergence and the emergence of new lineages (Hampe & Petit, 2005). However, climatic conditions experienced during glacial and interglacial intervals could also have provided opportunities for genetic divergence through selective pressures and adaptation associated with different local or regional environments. In populations adapted to ecologically dissimilar habitats, gene flow can be limited by selection against maladapted immigrants (Nosil *et al.*, 2005), and this might in turn have indirect effects on the whole genome, because reduction of gene flow promotes neutral divergence through increased genetic drift (Wright, 1931). In this case, genetic differentiation inferred from neutral markers is expected to be correlated with differences in local environments,

a pattern that has been described as 'isolation-by-adaptation' (IBA; Nosil *et al.*, 2008), in analogy to standard patterns of genetic differentiation with geographical distance, that is, 'isolation-by-distance' (IBD).

Despite the large body of research devoted to understanding the effects of climatic changes of the Quaternary, few studies to date have investigated the differential contribution of geographic isolation and/or climatic adaptation in creating population genetic divergence in temperate species. Long-lived organisms, such as trees, are especially well-suited models to address these questions. Many temperate trees are distributed over large areas characterized by a wide heterogeneity of both biotic and abiotic factors, and show local adaptation to environmental gradients at multiple spatial scales (Savolainen *et al.*, 2007), which can generate IBA patterns. Because of the buffering effects of their life history traits (great longevity, overlapping generations, prolonged juvenile phase) on changes in genetic structure (Austerlitz *et al.*, 2000), long-lived trees offer additional advantages over short-lived organisms for investigating the generation of IBA through the Quaternary, allowing us to explore how much genetic

variation is associated with current or past environmental conditions. For instance, the effects of climate during the last glaciation are still evident on contemporary patterns of genetic variation of long-lived *Quercus engelmannii* (Ortego *et al.*, 2012) and *Q. lobata* (Gugger *et al.*, 2013), suggesting that the genetic signal of past climate can persist over extended time periods in organisms with large effective population sizes and long generation times. Nevertheless, unravelling the effect of different climatic periods on spatial genetic divergence is challenging, because current observed patterns may result from the interplay among processes acting at different temporal scales. Assessing the most likely time course for the appearance of environmental barriers to gene flow is the first step towards accurately dissecting their role as actual contributors to IBA. Today, the existence of various palaeoclimatic databases allows us to evaluate the effect of period-specific climatic conditions on neutral genetic diversity by testing each period separately, and recently developed approximate Bayesian computation (ABC) methods can be used to elucidate complex demographic scenarios with relatively low demands in terms of computation effort (Beaumont, 2010), as well as to estimate the time of the inferred demographic processes.

English yew (*Taxus baccata*) is a long-lived, slow growing Tertiary relict (Hao *et al.*, 2008) native of Eurasian temperate and Mediterranean forests. Extending from North Africa to Scandinavia, and from the Iberian Peninsula to the Caspian Sea, yew grows under a wide range of environmental conditions, from oceanic to continental and Mediterranean climate (Thomas & Polwart, 2003). Although palaeoecological information on past yew distribution is scarce, some of the longest European Pleistocene pollen records indicate that *Taxus* expanded its range during several interglacials and made a much more significant contribution to vegetation in Europe than today (Mamakova, 1989; Turner, 2000; de Beaulieu *et al.*, 2001; Müller *et al.*, 2003; Koutsodendris *et al.*, 2010). Palynological records also indicate that yew was able to persist during the last glaciation, not only in southern refugia (Allen *et al.*, 2002; Carrión, 2002; Carrión *et al.*, 2003), but also in Central and Eastern Europe (Stewart & Lister, 2001; Willis & van Andel, 2004), although some debate still exists on the presence of cryptic refugia in northern Europe (Tzedakis *et al.*, 2013). The wide extent of environmental heterogeneity within the species' range, together with its long presence in Europe, make English yew an ideal species to investigate the impact of Quaternary climatic changes on genetic diversity via neutral and selective processes.

In this study, we used an integrated approach combining genetic and palaeoenvironmental data to elucidate the demographic history of *T. baccata* throughout its range, and determined the role of environmental and geographical factors in generating the observed patterns of genetic structure. About 5000 trees from 238 localities covering the yew natural range were genotyped with neutral microsatellite markers to identify distinct genetic clusters. ABC was used to select the most likely scenario shaping genetic diversity in this species and to set an approximate time frame for the inferred history. Finally, we used the available climatic information for three time periods – the last interglacial (LIG, *c.* 120 000–140 000 yr before present, BP), the last glacial

maximum (LGM, *c.* 21 000 yr BP) and present conditions (PRE, *c.* 1950–2000) – to evaluate the relative importance of current and past climatic conditions on the observed patterns of genetic variation. Coupling these approaches helped to determine whether standing patterns of genetic divergence are the result of historical isolation or, alternatively, of local adaptation to ecologically differentiated areas.

Materials and Methods

Sampling, DNA extraction and nuclear microsatellite genotyping

A total of 4992 samples ($n = 1\text{--}60$ per locality, mean 21) were collected at 238 localities covering the entire distribution range of *Taxus baccata* L. (Fig. 1; Supporting Information Table S1). Total DNA was isolated from 50 to 100 mg of dry leaf material using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) or a modified protocol from Dellaporta *et al.* (1983). Seven primer pairs for the amplification of nuclear microsatellites (nuSSRs) developed specifically for *T. baccata* were used for the genetic analysis following conditions described in Dubreuil *et al.* (2008).

Chloroplast DNA sequencing

Six chloroplast regions were tested following the PCR conditions given in Shaw *et al.* (2005): *rbcl*, *rpl36-rps8*, *trnH-psbA*, *trnC-ycf6*, *trnT-trnL* and *trnL-trnF*. Additionally, the *trnS-trnQ* spacer region was amplified as in Schirone *et al.* (2010). Only three of the regions were successfully amplified: *rbcl*, *trnS-trnQ* and *trnL-trnF*. The amplified products were screened for polymorphism using 1–2 individuals from 18 to 26 populations (Tables S2, S3) sampled across the distribution range of the species. PCR products were purified using the QIAquick gel extraction kit (Qiagen), and sequenced from both ends with an ABI 377

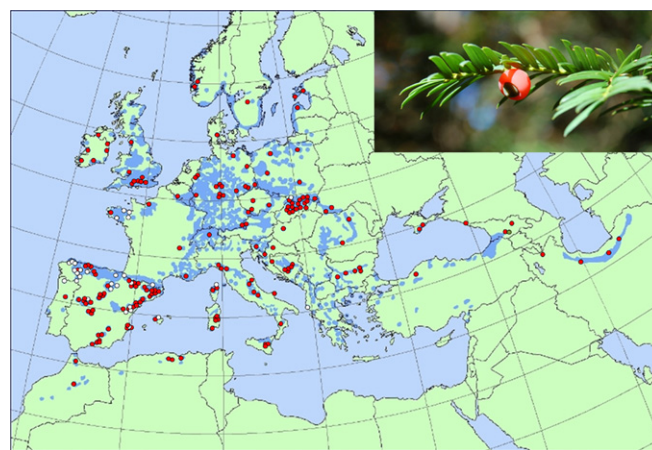


Fig. 1 Location of the 238 populations of *Taxus baccata* included in this study (red circles, populations with $n \geq 8$; white circles, populations with $n < 8$). The geographical distribution of the species is shown in blue (kindly provided by EUFORGEN, the European Forest Genetic Resources programme, www.euforgen.org). A female cone of the species is shown in the inset.

automated sequencer using the ABI BigDye™ Terminator Cycle Sequencing Ready Reaction Kit (Applied Biosystems, Foster City, CA, USA). Sequences available from previous studies (Shah *et al.*, 2008; Schirone *et al.*, 2010) were downloaded from GenBank and aligned with the newly obtained sequences (accession numbers KP115899–KP115935).

Genetic diversity and structure based on nuclear microsatellites

In all populations with at least eight individuals (195 locations, $n = 4829$), observed (H_O) and expected (H_E) heterozygosity were computed using GENETIX v4.04 (Belkhir *et al.*, 2001). The number of private alleles and allelic richness (A_R) were calculated using GENALEX v6.5 (Peakall & Smouse, 2012) and FSTAT v2.9.3.2 (Goudet, 2001), respectively. Linkage disequilibrium among all pairs of loci within each population was assessed by the Markov-chain approximation of the Fisher's exact test implemented in GENEPOP v4.0.7 (Rousset, 2008).

The same 195 populations were used to investigate genetic structure using different approaches. AMOVA (Excoffier *et al.*, 1992) was used to partition total molecular variance within and among populations using ARLEQUIN v3.5.1.2 (Excoffier *et al.*, 2005). Significance was obtained by nonparametric permutation using 10 000 replicates. Multilocus F_{ST} was estimated correcting for the possible presence of null alleles with the program FREE-NA (Chapuis & Estoup, 2007), using 1000 bootstraps to compute 95% confidence intervals. IBD was evaluated by testing the correlation between the matrix of pairwise ($F_{ST}/(1 - F_{ST})$) and the matrix of geographic distances (logarithmic scale) using the Mantel test implemented in GENETIX v4.04, with 10 000 permutations. Finally, Jost's estimator D_{est} (Jost, 2008) was computed to assess population differentiation using GENALEX v6.5, and significance was evaluated using 1000 replicates.

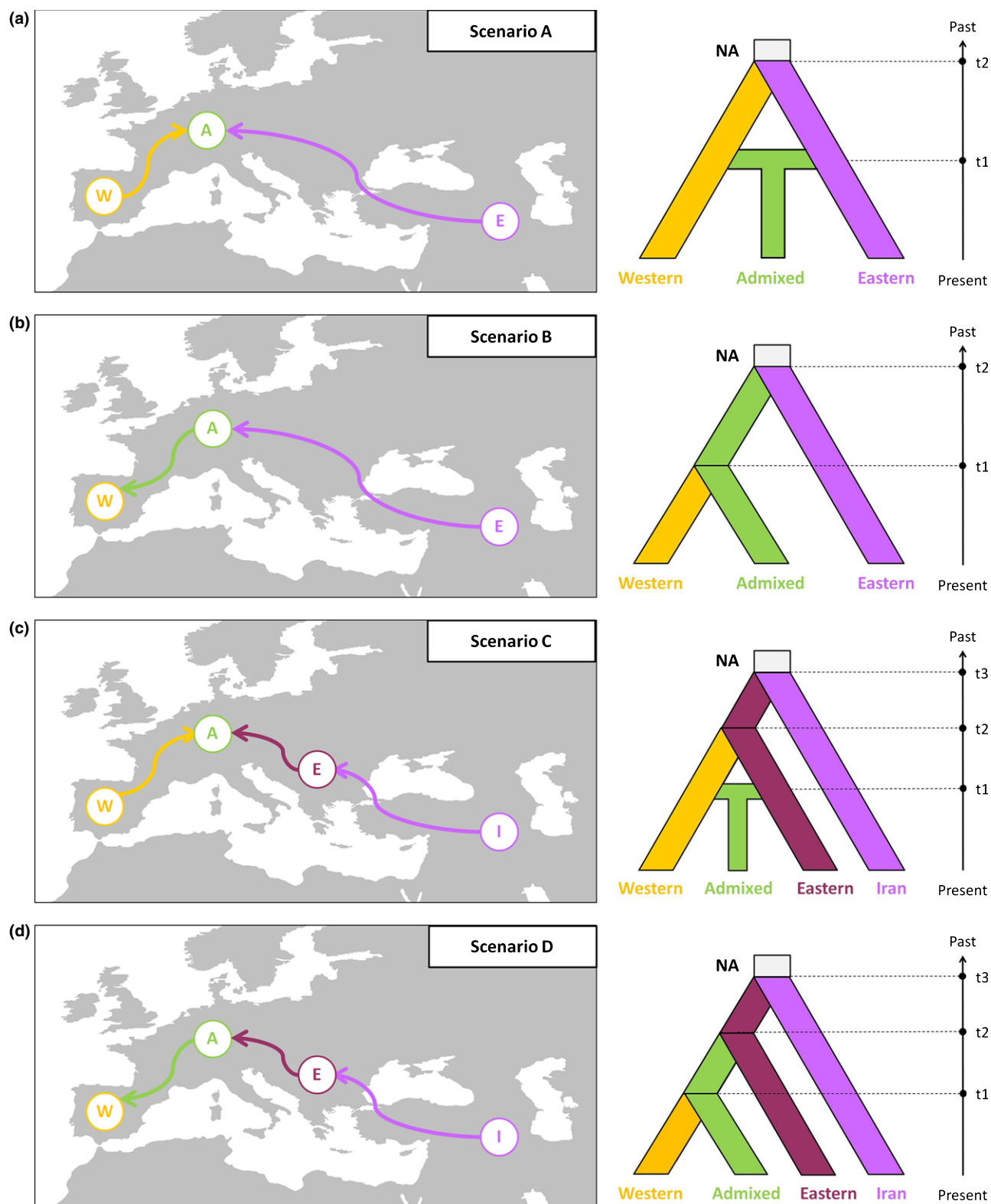
Two Bayesian clustering methods were used to infer distinct gene pools within the full dataset (238 locations, $n = 4992$). The program STRUCTURE v2.2 (Pritchard *et al.*, 2000) was run without prior population information, selecting the correlated allele frequencies model and assuming admixture. Ten independent runs for each K cluster (from $K = 1$ to 7) were performed, setting burn-in and run lengths of 50 000 and 500 000 iterations, respectively. The best number of clusters was determined following the recommendations by Pritchard & Wen (2004) and Evanno *et al.* (2005). Mean F_{ST} values measuring the divergence of each inferred cluster from a single hypothetical 'ancestral population' were also obtained (Falush *et al.*, 2003). Additionally, TESS v2.3 (François *et al.*, 2006) was employed to estimate the number of genetic clusters (K) present in the data incorporating the geographical location of individuals as *a priori* information. Both admixture (BMY model; spatial interaction parameter: 0.6) and nonadmixture models were used to perform five independent runs for K ranging from 1 to 7, with a burn-in of 10 000 and a total number of 50 000 sweeps. The optimal value of K was determined by plotting the deviance information criterion (DIC) against K and choosing the values of K_{max} corresponding to a plateau of the curve (François & Durand, 2010). To graphically

represent the results obtained, the averaged results of the assignments of STRUCTURE and TESS were plotted on maps generated with ARCGIS v9.1. (ESRI, Redlands, CA, USA).

Demographic history (ABC models)

We applied the ABC framework implemented in DIYABC v1.0.4.46 (Cornuet *et al.*, 2010) to nuSSRs to infer the demographic history of *T. baccata*. The information obtained from the earlier analyses was the starting point for designing the different scenarios to test (for further details, see the Results section). STRUCTURE and TESS identified slightly different best number of clusters ($K = 2$ (*Western, Eastern*) and $K = 3$ (*Western, Eastern, Iran*), respectively), but both supported a first partition between *Western* and *Eastern* samples, and the presence of *Admixed* populations at the intersection of both clusters, suggesting a secondary contact of divergent lineages. However, a clear westward cline of decreasing diversity from Iran to the Mediterranean area was also detected, which might be indicative of a colonization pattern. Thus, we designed four scenarios to test alternative hypotheses considering two or three genetic pools (Fig. 2). Scenario A tested a 'secondary contact' of two separated gene pools (*Western, Eastern*). Scenarios B and D tested a 'colonization' event from the east, the former considering two genetic pools (*Western, Eastern*), and the latter with three (*Western, Eastern, Iran*). Scenario C considered a 'colonization' from *Iran*, the separation of European samples into two genetic pools (*Eastern, Western*), and a posterior 'secondary contact'.

For scenarios A and B, three groups of populations were created: *Eastern*, *Western* and *Admixed*. A population was considered as admixed when the proportion of individuals assigned to the eastern or the western cluster was $< 70\%$. The proportion of membership and the assignment of populations to each specific group are reported in Table S1. Given that recent studies suggest that pooling data across populations can be a problem to infer demography (Chikhi *et al.*, 2010), the following procedure was designed to avoid the potential confounding effects of population structure on the inference of demographic parameters. Ten different sets of populations were constructed, each containing *c.* 500 individuals, representing *c.* 10% of the whole dataset (Fig. S1). Each set was composed of *c.* 200 individuals belonging to the *Eastern* pool, *c.* 200 individuals from the *Western* one and *c.* 100 of *Admixed* composition (hereafter called '500-sample datasets'). To minimize the effect of spatial genetic structure, we sought that the populations included in each dataset were geographically close or that genetic divergence among populations was low. For scenarios C and D, two additional datasets with four groups of populations were constructed (Fig. S1), considering the *Iran* pool (Iran, Georgia) to be independent from the *Eastern* one, as inferred using TESS for $K = 3$ and STRUCTURE for $K = 4$. In this case, datasets were composed by *c.* 200 individuals belonging to the *Eastern*, *Western* and *Iran* pools, respectively, and *c.* 100 of *Admixed* composition (hereafter called '700-sample datasets'). The composition of these two datasets for each ABC run was different for all pools except for the *Iran* one, where only 199 individuals were available.



One million simulations were run for each dataset. Prior parameter distributions were chosen as broad as possible to explore a wide range of population sizes and time frames (measured in generations): uniform (10; 100 000) for current effective population sizes, uniform (1; 100 000) for divergence times t_1 , t_2 and t_3 (with $t_3 > t_2$ and $t_2 > t_1$), uniform (10; 1000 000) for ancestral effective population size, and uniform (0.001; 0.999) for admixture rate. A generalized stepwise mutation model was assumed and default values were used for all prior mutation parameters, except for the mean mutation rate, for which minimum and maximum default values (10^{-4} – 10^{-3} mutations per locus per generation) were enlarged to 10^{-5} – 10^{-3} after previous runs giving biased posteriors towards the lower mutation rate. Each simulation was summarized by the following statistics: mean number of alleles and mean genetic diversity (Nei, 1987) for each cluster, and mean number of alleles, mean genetic diversity, F_{ST} , mean index of classification (Rannala & Mountain, 1997) and shared allele distance (Chakraborty & Jin, 1993) between pairs of clusters. After ensuring that this combination of scenarios and priors was able to produce datasets similar to the observed one (Fig. S2), the posterior probabilities of each scenario were calculated with a local logistic regression procedure using the 1% closest simulated points. Retained simulations were used to infer parameter posterior distributions by local linear regression using a logit transformation of the parameters. The reliability of the model and chosen scenario was evaluated for each of the 12 simulations by performing model checking and computing the confidence in scenario choice (see Notes S1 for further details).

Past and present impact of environmental factors on genetic structure

We used the climatic information available at the WorldClim database (Hijmans *et al.*, 2005) to evaluate the effect of past and present climatic conditions on current genetic structure. For the present time (PRE, *c.* 1950–2000), 19 bioclimatic variables were downloaded for the 195 populations with $n \geq 8$. Two bioclimatic variables that were highly correlated with the others ($r > 0.9$) were excluded, and the remaining variables were summarized into the first two axes of a principal component analysis (PCA) using R (R Core Team, 2013). The environmental variables with loadings on the PCA axes > 0.5 (Table S4) and the 238 occurrence points were used to model current climatically suitable areas for English yew with maximum entropy (MAXENTv3.3.3) and BIOCLIM (DIVA-GIS v7.5.) algorithms. Predictions were also generated separately for the *Western* (153 sampling sites) and *Eastern* (64 sampling sites) gene pools to examine whether genetic divergence among them was environmentally induced. The modelled distributions were generated with 75% of the points (training data) and cross-validated with 25% of the remaining localities (test data), averaged over 10 runs. The performance of the models was tested by measuring the area under the receiver operating characteristic curve (AUC). The logistic outputs of MAXENT models were transformed to presence–absence maps using the maximum training sensitivity plus

specificity (MTSS) threshold. For BIOCLIM, maps were obtained leaving only values with high, very high or excellent suitability (i.e. within the 5–95th percentile interval).

In order to determine the contribution of present environment on genetic differentiation, we tested for the relationship between pairwise F_{ST} and climatic distance while controlling for geographic distance. We computed climatic (Euclidian) distance matrices based on population scores for both PCA axes (PC1, PC2), and for each environmental variable. Tests were performed for the whole dataset and for each genetic cluster (*Eastern*, *Western*) using partial Mantel tests ('mantel.partial' function; R Core Team, 2013) and multiple matrix regressions (MMRR script; Wang, 2013). Significance tests were based on 10 000 permutations. To reduce the risk of spurious correlations, in particular for less conservative MMRR tests, we only considered those correlations that were significant with both methods.

The same procedures were applied to investigate the contribution of past climate to current genetic differentiation. We projected the models for the present onto three paleoclimate layers, the Community Climate System Model (CCSM) and the Model for Interdisciplinary Research on Climate (MIROC) for the last glacial maximum (LGM, *c.* 21 000 yr BP), and the model for the last interglacial (LIG, *c.* 120 000–140 000 yr BP). Then, to investigate the correlation of past climate and observed genetic differentiation (F_{ST}), we retained only those populations where suitable environment have existed for yew persistence during these periods. Although distribution of *T. baccata* may have not been exactly the same across the Quaternary, projections suggest a rather stable distribution of the species in relatively large parts of its range (see the Results section), so using only populations that could have been located at or near the present locations can be considered a reasonable approximation of the distribution of the species in the past. Thus, we selected those occurrence points with logistic output values in MAXENT above the respective MTSS thresholds in each model, and with suitability values above the 5–95th percentile interval in BIOCLIM. Because of the present distribution of English yew was more accurately predicted when combining modelled distributions of single gene pools than from the full model (see the Results section), we constructed datasets combining the predicted suitable populations obtained for *Western* and *Eastern* models of past climate (indicated in Table S1), and used them to perform partial Mantel and MMRR tests as for the present time.

Results

Chloroplast DNA sequencing

Chloroplast regions comprised 1359, 656 and 344 aligned positions for *rbcl*, *trnS-trnQ* and *trnL-trnF*, respectively. No polymorphism was found for the *rbcl* gene, and *trnS-trnQ* and *trnL-trnF* showed two closely related haplotypes, respectively (Tables S2, S3). For both markers, populations harbouring different haplotypes were located at Guilan and Golestan provinces (Iran), at the eastern extreme of the distribution of English yew (Fig. S3).

Genetic diversity and structure based on nuclear microsatellites

A_R , expected (H_E) and observed (H_O) heterozygosity ranged from 2.243 to 5.295, 0.354 to 0.855, and 0.171 to 0.768, respectively (Table S1). Among a total of 3957 tests for linkage disequilibrium between pairs of loci, 98 were significant ($P < 0.05$) after sequential Bonferroni corrections, but almost all involved the same eight populations. Because the application of Bayesian methods rely on the assumption of linkage equilibrium between loci, we performed additional runs with the STRUCTURE program excluding these populations.

Very similar pairwise F_{ST} were obtained when correcting or not for the presence of null alleles. Corrected values ranged from 0.001 to 0.599, with an overall $F_{ST} = 0.149$. Only 84 out of 18 915 population pairs were not significantly ($P < 0.05$) differentiated from each other after a sequential Bonferroni correction for multiple tests (Table S5). These results were in agreement with AMOVA, with a 16.41% of the total variance explained by differences among populations (Table S6). Overall Jost's D_{est} was 0.478, indicating that the proportion of allelic differentiation among populations was higher than the proportion of variance in allele frequencies. The correlation between genetic and geographic distances was highly significant ($r = 0.281$, $P < 0.001$), suggesting the existence of an isolation-by-distance pattern.

STRUCTURE runs including or excluding populations with significant linkage between loci produced almost identical results. The method identified an optimal partition in two genetic pools with a clear geographical pattern: populations from central Europe to Iran (*Eastern*) and populations from the western range (*Western*), with a contact zone of admixed populations (*Admixed*) located along Central Europe, Italy and the Mediterranean islands (Fig. 3). An additional partition ($K = 3$) subdivided the western group into two differentiated pools, predominantly located in Central Europe and the Mediterranean area, respectively (Fig. S4). Increasing the number of partitions ($K = 4$) produced the splitting of samples from Iran and Georgia as an independent pool (*Iran*) within the *Eastern* one (Fig. S4). The F_{ST} values obtained by the correlated frequencies model in STRUCTURE increased toward the west, suggesting that eastern populations were closer to the hypothetical 'ancestral population'.

The best model with TESS was the one considering admixture, and generated a very similar population clustering for $K = 2$ (Fig. S4), but the best number of clusters was inferred at $K = 3$ (Fig. 3), and the splitting of the easternmost group (*Iran*) occurred earlier (i.e. for $K = 3$ in TESS, and for $K = 4$ in STRUCTURE). Nevertheless, the same trends were identified with both approaches: the first level of divergence produced the partition of *Western* and *Eastern* samples; populations showing higher levels of admixture were located at the intersection of both clusters; divergence from the hypothetical 'ancestral population' increased towards the west; and increasing number of partitions led to the split of the easternmost samples (Georgia, Iran) as an independent group (*Iran*).

The mean genetic diversity of populations assigned to the *Eastern* pool and those of *Admixed* composition was significantly

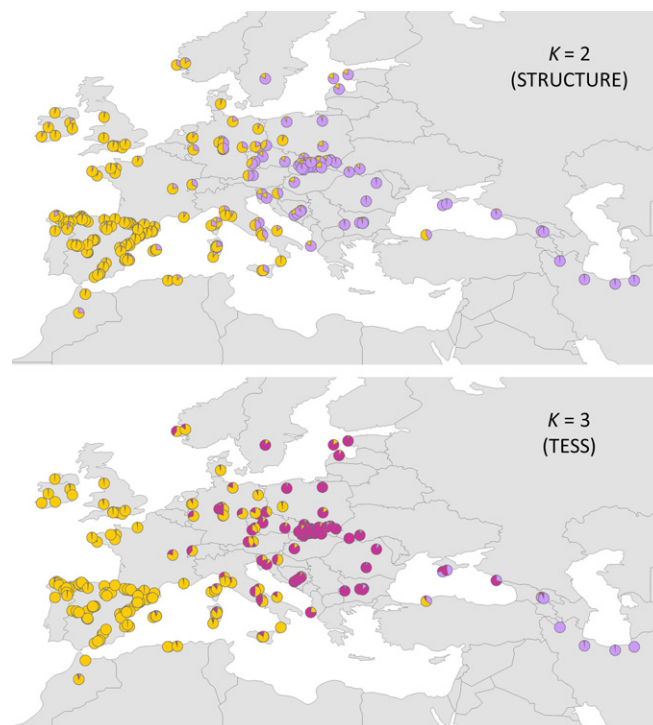


Fig. 3 Best number of genetic clusters (K) obtained for *Taxus baccata* using STRUCTURE ($K = 2$) and TESS ($K = 3$). The eight populations with significant linkage among loci are excluded. Pie charts show averaged values of the different runs for the proportion of membership to each genetic cluster. Different colours indicate different genetic clusters, and are the same as in Fig. 2.

higher than that of the *Western* cluster (mean $A_{R(E)} = 4.46$, $A_{R(A)} = 4.36$, $A_{R(W)} = 3.61$; mean $H_{E(E)} = 0.773$, $H_{E(A)} = 0.759$; $H_{E(W)} = 0.653$; Duncan's test after ANOVA: $P < 0.001$), indicating a pattern of decreasing genetic diversity from east to west (Fig. 4). In addition, the number of populations displaying private alleles was higher in the *Eastern* pool than in the *Western* one (23 vs 15), as well as the number of private alleles (43 vs 17). Of these 43 private alleles, 23 were detected in populations from Iran and Georgia (*Iran* pool). Only seven *Admixed* populations had private alleles, and the proportion was low (seven out of 67). Jost's estimator indicated higher differences in allele composition among populations in the east ($D_{est(E)} = 0.489$, $D_{est(A)} = 0.387$, $D_{est(W)} = 0.351$), whereas greater deviations from panmixia (F_{ST}) were detected in the west ($F_{ST(E)} = 0.127$, $F_{ST(A)} = 0.103$, $F_{ST(W)} = 0.147$).

Demographic history (ABC models)

All the ABC simulations were able to discriminate between the tested scenarios, with high posterior probabilities and 95% confidence intervals never overlapping those of the other scenario (Table 1). The most likely scenario using the '500-sample datasets' was scenario A, with a strong support in almost all cases ($PP \geq 0.8$; Table 1). However, scenario B was always chosen when the datasets used for simulations included the easternmost samples (Georgia, Iran) as representatives of the *Eastern* pool (sim4, sim5 and sim8, see Fig. S1), albeit with lower support

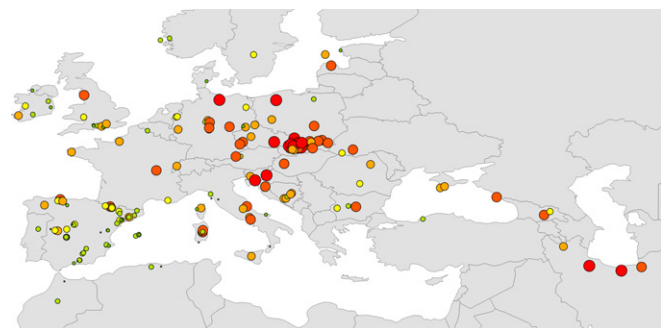


Fig. 4 Distribution of genetic diversity (A_R , allelic richness; H_E , unbiased expected heterozygosity) in *Taxus baccata*. Values for A_R and H_E are indicated by the circle size and (○>○>○) the colour gradient (red > orange > yellow > green), respectively.

(Table 1). This was in accordance with genetic diversity distribution and together suggests an eastern colonization of Europe, with easternmost populations constituting an independent gene pool (*Iran*) that was not the source of admixture in Central Europe. Similarly, simulations performed on the two '700-sample datasets', considering three differentiated pools (*Iran*, *Eastern*, *Western*), unambiguously indicated support for scenario C (> 0.9; Table 1), which tested a first migration wave from the east (*Iran*), a more recent separation of European samples in two distinct pools (*Eastern*, *Western*), and a secondary contact (*Admixed*) between them (Fig. 2). Model testing procedures further supported the reliability of this scenario (see Notes S1 for further information). Parameter posterior distributions are shown in Fig. S5.

Under this model (scenario C), estimated time of divergence among Iranian and European samples would have occurred, on average, c. 6 Myr BP (90% credible intervals: 1.35–14.78 Myr BP, Table 1), assuming a generation time of c. 100 yr for English yew. Although reproduction can begin earlier when growing under open canopy conditions, yew usually grows as isolated understorey tree, and reach maturity later, between 70 and 120 yr of age (Thomas & Polwart, 2003; L. Paule & M. Riba, unpublished data). The posterior separation of *Eastern* and *Western* clusters, and the subsequent admixture event would have taken place c. 2.2 Myr BP (90% credible intervals: 0.5–7.5 Myr BP) and 200 000 yr BP (90% credible intervals: 50 000–800 000 yr BP), respectively (Table 1). These estimates are mostly in agreement with those obtained with the '500-sample datasets', with averaged estimates of both events c. 2 Myr BP and 230 000 yr BP, respectively (Table 1). Although 90% credible intervals are large for all the simulations, model checking confirmed that the model was consistent with the observed data, suggesting that large confidence intervals are due to data information content and not to a model misfit (Notes S1).

Past and present impact of environmental factors on genetic structure

The first two PCA axes explained 52% of the variation for the present climate. PC1 was mainly correlated with temperatures, whereas PC2 was positively correlated with precipitation (Table

S4). Despite some overlap, populations belonging to *Western* and *Eastern* clusters defined clear groups along the first axis of the multivariate space (Fig. S6), indicating that substantial differences exist in climate for each geographic region. On average, populations within the *Western* cluster experienced smaller seasonal temperature fluctuations, warmer temperatures (annual mean, minimum and maximum), higher temperatures during the driest quarter, and less precipitation during the warmest quarter (ANOVA: $P < 0.001$).

Averaged AUC values for the replicate runs were > 0.870 for all distribution models, supporting their predictive power. The predicted full species model for the present generated with MAXENT was fairly congruent with yew current distribution, (Fig. S7), but this was more accurately predicted when combining modelled distributions of single gene pools, especially with regard to Eastern Europe (Fig. 5). Similar results were obtained using the BIOCLIM algorithm (Notes S2).

The CCSM and MIROC models for the LGM yielded large differences in predicted distributions, and were highly dependent on the algorithm used (Figs 5, S7; Notes S2). MAXENT models suggested much wider suitable areas than BIOCLIM, especially with regard to CCSM. However, all models supported the existence of large suitable areas for English yew in several southern refugia (i.e. the Balkans, Iberia and Italy). Projections for LIG produced similar models with MAXENT and BIOCLIM, showing a westward shift with respect to present-day climatic conditions, both for the *Western* and *Eastern* clusters (Figs 5, S7; Notes S2).

Despite a substantial lack of precision for LGM models, a common trend was that models produced using localities from either *Western* or *Eastern* gene pools alone showed little overlap of their predicted distributions for all periods considered, especially during both interglacials (Fig. 5), suggesting that *Eastern* and *Western* clusters have occupied environmentally different regions since the long past.

After controlling for geographic distance, and at the scale of the whole species range, there was a significant positive association between pairwise F_{ST} and the PC1 axis for the present climate ($r_{Env-PRE} = 0.157$, $b_{Env-PRE} = 0.154$, $P < 0.001$), whereas relations with PC2 variables were not significant (Table 2). Analysed separately, annual mean temperature ($r_{Env-PRE} = 0.171$, $b_{Env-PRE} = 0.161$, $P < 0.001$) and minimum temperature of the coldest month ($r_{Env-PRE} = 0.141$, $b_{Env-PRE} = 0.145$, $P < 0.001$) were the most relevant variables explaining population genetic structure. Very similar results were found within the *Western* pool, with a significant correlation among genetic differentiation and PC1 variables ($r_{Env-PRE} = 0.174$, $b_{Env-PRE} = 0.175$, $P < 0.01$), and more specifically with annual mean temperature ($r_{Env-PRE} = 0.219$, $b_{Env-PRE} = 0.218$, $P < 0.01$) and minimum temperature of the coldest month ($r_{Env-PRE} = 0.163$, $b_{Env-PRE} = 0.165$, $P < 0.01$). Within the *Eastern* pool, no significant correlations were found for both tests except for temperature seasonality ($r_{Env-PRE} = 0.136$, $b_{Env-PRE} = 0.150$, $P < 0.05$).

MAXENT projections onto past climatic models suggested that suitable conditions would have existed for the persistence of 102–123 and 94 populations during LGM and LIG, respectively

Table 1 Demographic approximate Bayesian computation (ABC) models for *Taxus baccata*

Simulation	Supported scenario (PP)	I (10 ⁴)	E (10 ⁴)	A (10 ⁴)	W (10 ⁴)	NA (10 ⁵)	ra	t1	t2	t3
Considering two gene pools (<i>Western, Eastern</i>)										
sim1_500	A (> 0.9) [0.9663, 0.9895]	–	7.87 (4.34–9.81)	0.78 (0.18–4.25)	4.43 (2.1–7.13)	8.13 (2.77–9.83)	0.589 (0.204–0.867)	0.071 (0.016–0.281)	2.110 (0.444–7.950)	–
sim2_500	A (> 0.9) [0.9165, 0.9681]	–	6.39 (2.74–9.53)	3.09 (0.87–8.19)	4.61 (2.12–7.68)	6.42 (1.28–9.59)	0.431 (0.108–0.789)	0.164 (0.037–0.644)	2.480 (0.516–8.380)	–
sim3_500	A (c. 0.6) [0.5135, 0.7619]	–	6.43 (2.83–9.51)	4.26 (1.33–8.89)	2.92 (1.01–6.90)	3.19 (0.33–8.47)	0.207 (0.032–0.661)	0.195 (0.041–0.907)	1.870 (0.365–7.960)	–
sim4_500	B (c. 0.6) [0.5213, 0.7602]	–	6.65 (2.89–9.64)	3.99 (1.87–6.80)	4.21 (1.77–7.67)	6.92 (2.00–9.72)	–	0.313 (0.079–1.150)	1.870 (0.408–7.660)	–
sim5_500	B (c. 0.7) [0.6357, 0.8293]	–	3.31 (0.96–8.38)	4.71 (1.93–8.60)	4.54 (1.83–8.33)	4.77 (0.84–9.19)	–	0.573 (0.145–2.070)	3.200 (0.736–8.620)	–
sim6_500	A (> 0.9) [0.8941, 0.9701]	–	6.08 (2.67–9.40)	0.62 (0.14–3.93)	4.15 (1.47–8.36)	0.51 (0.04–3.50)	0.414 (0.070–0.871)	0.062 (0.012–0.316)	0.841 (0.151–5.360)	–
sim7_500	A (> 0.9) [0.9515, 0.9865]	–	7.25 (3.58–9.67)	0.86 (0.20–4.86)	3.58 (1.09–7.28)	4.34 (0.57–9.02)	0.507 (0.112–0.880)	0.069 (0.015–0.346)	1.800 (0.348–7.560)	–
sim8_500	B (> 0.9) [0.8454, 0.9581]	–	7.39 (3.65–9.76)	2.60 (1.22–4.58)	6.63 (3.40–9.33)	5.95 (1.42–9.57)	–	0.480 (0.138–1.410)	1.730 (0.385–7.350)	–
sim9_500	A (c. 0.8) [0.6901, 0.8630]	–	5.53 (2.11–9.40)	4.83 (1.61–9.00)	3.71 (1.57–6.60)	7.22 (1.95–9.76)	0.285 (0.064–0.622)	0.288 (0.065–1.060)	1.940 (0.376–7.900)	–
sim10_500	A (> 0.9) [0.9301, 0.9745]	–	6.72 (3.14–9.63)	3.37 (0.92–8.21)	5.08 (2.38–8.18)	5.78 (0.93–9.48)	0.374 (0.088–0.756)	0.124 (0.026–0.500)	2.270 (0.479–8.160)	–
Considering three gene pools (<i>Western, Eastern, Iran</i>)										
sim1_700	C (> 0.9) [0.8746, 0.9537]	8.06 (5.04–9.78)	4.30 (1.74–8.45)	3.95 (1.36–8.61)	2.96 (1.33–5.85)	3.86 (0.36–9.35)	0.405 (0.065–0.859)	0.193 (0.044–0.826)	2.690 (0.732–8.270)	8.190 (1.640–20.40)
sim2_700	C (> 0.9) [0.8837, 0.9585]	7.59 (4.24–9.74)	4.00 (1.69–8.00)	2.96 (0.90–7.77)	3.09 (1.56–5.27)	7.42 (2.50–9.76)	0.529 (0.127–0.880)	0.202 (0.054–0.630)	1.790 (0.586–5.070)	4.030 (1.050–9.150)

PP, posterior probability (numbers in parenthesis) and 95% confidence intervals (numbers in square brackets); I, current effective population size of the *Iran* gene pool; E, current effective population size of the *Eastern* gene pool; A, current effective population size of the *Admixed* samples; W, current effective population size of the *Western* gene pool; NA, ancestral effective population size; ra, admixture rate; t1–t3, estimated times of the different events depicted in Fig. 2 (in Myr). Values are medians (5 and 95% quartiles) of posterior distributions.

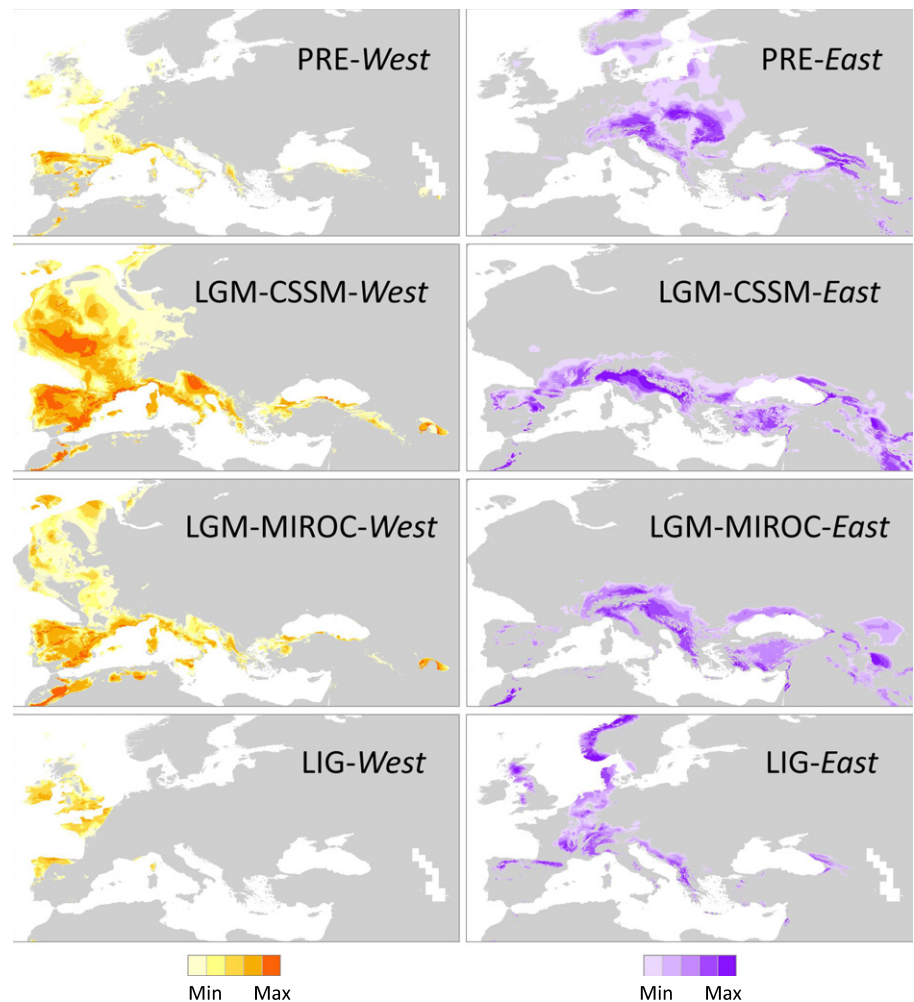


Fig. 5 MAXENT predicted suitability for Western and Eastern gene pools of *Taxus baccata* during three time periods: LIG, last interglacial (c. 120 000–140 000 yr before present, BP); LGM-CCSM and LGM-MIROC, last glacial maximum (c. 21 000 yr BP) employing two different paleoclimate layers, the Community Climate System Model (CCSM) and the Model for Interdisciplinary Research on Climate (MIROC); PRE, present conditions (c. 1950–2000). Darker colours indicate higher probabilities of suitable climatic conditions. Unsuitable areas and those with logistic output values below the maximum training sensitivity plus specificity (MTSS) threshold are indicated in grey.

(Table 3). In these populations, we found that LIG climate contributed similarly as the present to genetic divergence, with a positive association between F_{ST} and annual mean temperature ($r_{Env-LIG} = 0.104$, $b_{Env-LIG} = 0.115$, $P < 0.05$), and minimum temperature of the coldest month ($r_{Env-LIG} = 0.096$, $b_{Env-LIG} = 0.106$, $P < 0.05$). In addition, we found a significant contribution of mean diurnal temperature range ($r_{Env-LIG} = 0.177$, $b_{Env-LIG} = 0.170$, $P < 0.01$), and no significant correlation with the PC2 axis (Table 3). Positive significant associations for both Mantel and MMRR tests were not detected during LGM models. These results were confirmed when using suitable populations predicted with BIOCLIM (Notes S2).

Discussion

Demographic history of English yew

The combination of Bayesian clustering and approximate Bayesian computation (ABC) methods shed light on the demographic history of English yew. According to our ABC results, nuSSRs in *Taxus* seem to retain the imprint of very ancient events, as suggested by the divergence time estimates for the inferred demographic processes. Even so, a near absence of variation and spatial

structure for chloroplast DNA markers was observed, in accordance with the slow chloroplast nucleotide substitution rate reported in conifers (Willyard *et al.*, 2007).

ABC simulations suggest that the most likely demographic scenario for *T. baccata* involves a first migration wave from eastern territories (Iran) to the west, a more recent separation of the European samples into two gene pools (Eastern, Western), and a secondary contact (Admixed) of both clusters along Central Europe, Italy and Mediterranean islands (Fig. 2, scenario C). The ancient migration from the east is also supported by the westward decline of genetic diversity (Fig. 4), and by the fact that F_{ST} values from the hypothetical 'ancestral population' for each inferred cluster always increased towards the west, suggesting that easternmost populations were closer to the ancestral one. In agreement with our results, recent studies set the origin of *Taxus* in North America or South West China during the late Cretaceous to middle Eocene (66.55 ± 11.22 Myr before present (BP)), from which was dispersed to the current distribution areas (Hao *et al.*, 2008). The genus probably reached Europe through the Irano-Turanian region, which has been postulated as a key source for the colonization of the Mediterranean region (Thompson, 2005; Mansion *et al.*, 2008). This event probably occurred before the Lower Miocene, as indicated by the oldest fossil record (16–23 Myr BP;

Kunzmann & Mai, 2005). An eastern origin and westward colonization of the Mediterranean, still reflected in current genetic structure, has also been postulated for other tree genera (e.g. *Abies*, Linares, 2011; *Frangula*, Petit *et al.*, 2005; *Laurus*, Rodríguez-Sánchez *et al.*, 2009).

Our ABC simulations place the separation among the Iranian and European genetic pools *c.* 6 Myr BP, although 90% credible

Table 2 Partial Mantel (PM) correlation (r) and multiple matrix regression (MMRR) coefficients (b) between genetic distance (F_{ST}) and environmental variables for the present time (PRE, *c.* 1950–2000)

	PRE		
	MMRR		
	$b_{Geo-PRE}$	$b_{Env-PRE}$	PM $r_{Env-PRE}$
Whole range			
$F_{ST} - PC1/Geo$	0.305***	0.154***	0.157**
$F_{ST} - PC2/Geo$	0.355***	−0.043 ns	−0.046 ns
$F_{ST} - BIO1/Geo$	0.328***	0.161***	0.171***
$F_{ST} - BIO6/Geo$	0.289***	0.145***	0.141***
Eastern pool			
$F_{ST} - PC1/Geo$	0.406***	0.037 ns	0.040 ns
$F_{ST} - PC2/Geo$	0.457***	−0.143*	−0.148 ns
$F_{ST} - BIO4/Geo$	0.325***	0.150*	0.136*
Western pool			
$F_{ST} - PC1/Geo$	0.098**	0.175***	0.174**
$F_{ST} - PC2/Geo$	0.126**	−0.002 ns	−0.002 ns
$F_{ST} - BIO1/Geo$	0.108**	0.218***	0.219**
$F_{ST} - BIO6/Geo$	0.096*	0.165***	0.163**

Analyses were conducted considering the 195 populations of *Taxus baccata* with $n \geq 8$ (whole range), and only populations within *Western* (153 sampling sites) and *Eastern* (64 sampling sites) gene pools. Variables accounting for PC1 (BIO1, BIO4, BIO9, BIO10, BIO11, BIO18) and PC2 (BIO12, BIO13, BIO16, BIO19) are described in Supporting Information Table S4. BIO1, annual mean temperature. BIO4, temperature seasonality. BIO6, minimum temperature of the coldest month. ***, $P < 0.001$; **, $P < 0.01$; *, $P < 0.05$; ns, not significant. Positive significant tests for both MMRR and PM tests are in bold.

Table 3 Partial Mantel (PM) correlation (r) and multiple matrix regression (MMRR) coefficients (b) between genetic distance (F_{ST}) and environmental variables for the last glacial maximum (LGM, *c.* 21 000 yr before present, BP) and the last interglacial (LIG, *c.* 120 000–140 000 yr BP)

	LGM-MIROC (102)			LGM-CCSM (123)			LIG (94)		
	MMRR			MMRR			MMRR		
	$b_{Geo-MIROC}$	$b_{Env-MIROC}$	$r_{Env-MIROC}$	$b_{Geo-CCSM}$	$b_{Env-CCSM}$	$r_{Env-CCSM}$	$b_{Geo-LIG}$	$b_{Env-LIG}$	$r_{Env-LIG}$
F_{ST} – PC1/Geo	0.166**	–0.088*	–0.086 ns	0.317***	–0.128**	–0.132 ns	0.343***	0.056*	0.058 ns
F_{ST} – PC2/Geo	0.130**	0.077 ns	0.077 ns	0.285***	0.069 ns	0.072 ns	0.358***	0.015 ns	0.016 ns
F_{ST} – BIO1/Geo	0.164**	–0.062 ns	–0.057 ns	0.315***	–0.096*	–0.098 ns	0.357***	0.115*	0.104*
F_{ST} – BIO2/Geo	0.137**	0.031 ns	0.031 ns	0.306***	–0.068 ns	–0.070 ns	0.318***	0.170***	0.177**
F_{ST} – BIO4/Geo	0.219**	–0.199***	–0.185 ns	0.314***	–0.120***	–0.123 ns	0.490***	–0.235***	–0.209 ns
F_{ST} – BIO6/Geo	0.142**	–0.005 ns	–0.004 ns	0.312***	–0.065 ns	–0.065 ns	0.303***	0.106**	0.096*

The number of *Taxus baccata* populations retained for the analyses (i.e. with suitability values of MAXENT predicted distributions above the maximum training sensitivity plus specificity threshold) are indicated in brackets behind each period considered, and are specified in Supporting Information Table S1. Variables accounting for PC1 were: BIO1, BIO5, BIO6, BIO9, BIO10, BIO11 for LGM-MIROC; BIO1, BIO2, BIO5, BIO6, BIO8, BIO9, BIO10, BIO11 for LGM-CCSM; and BIO1, BIO2, BIO4, BIO6, BIO9, BIO11, BIO18 for LIG. Variables accounting for PC2 were the same for all periods considered, and the same as for PRE (BIO12, BIO13, BIO16, BIO19; Table S4). BIO1, annual mean temperature. BIO2, mean diurnal range (mean of monthly (max temp – min temp)). BIO4, temperature seasonality. BIO6, min temperature of the coldest month. ***, $P < 0.001$; **, $P < 0.01$; *, $P < 0.05$; ns, not significant. Positive significant tests for both MMRR and PM tests are in bold.

intervals are wide (on average 1.35–14.78 Myr BP, Table 1). In addition, DIYABC does not model continuous gene flow at each generation, which could have led to underestimating the divergence time. Nevertheless, assuming that lack or reduced gene flow is a reasonable assumption in English yew (see Dubreuil *et al.*, 2010; González-Martínez *et al.*, 2010; Chybicki *et al.*, 2011; Burgarella *et al.*, 2012), as also suggested by high levels of pairwise genetic differentiation in our study (see the Results section). An ancient separation is also evident from the results obtained using chloroplast DNA markers, because the only distinct haplotypes were located at the eastern extreme of the distribution (Guilan and Golestan provinces, Iran), suggesting that both groups became isolated a long time ago. This is additionally confirmed by the significantly higher number of private alleles detected at nuclear microsatellites within the *Eastern* pool, particularly in populations from Iran and Georgia. This ancient vicariance might be associated to the intense changes that occurred during the latest Miocene (6.1–5.7 Myr BP; Popov *et al.*, 2006), which could have favoured both migration and differentiation within the Mediterranean Basin.

Around 2 Myr BP (90% credible intervals *c.* 0.5–7.5 Myr BP), the European populations split into two distinct genetic pools (*Eastern*, *Western*), and a posterior admixture of both lineages seem to have occurred *c.* 200 000 yr ago (90% credible intervals *c.* 50 000–800 000 yr). These results are consistent with the expected pattern assuming that *T. baccata* survived in two allopatric refugia since the beginning of the Quaternary, from which they expanded and converged further north during warm interglacial periods. Such an east–west pattern of differentiation across the Mediterranean region has been reported for other trees (*Laurus nobilis*, Rodríguez-Sánchez *et al.*, 2009; *Olea europaea*, Besnard *et al.*, 2007; *Quercus suber*, Lumaret *et al.*, 2002), herbaceous species (*Arabidopsis thaliana*, François *et al.*, 2008) and coastal plants (*Carex extensa*, Escudero *et al.*, 2010), and has been interpreted as a result of an east–west isolation during glaciations of the Quaternary (e.g. François *et al.*, 2008; Escudero *et al.*,

2010). Our results, however, suggest that interglacials could have played a key role in maintaining genetic divergence between both groups, as we discuss in the next subsection.

Past and present impact of environmental factors on genetic structure

Our analyses support the hypothesis that both geography and climate have played a significant role in shaping genetic structure of English yew. The divergence between *Western* and *Eastern* clusters can be explained by their persistence in spatially isolated refugia during glacial periods. However, species distribution models (Fig. 5) revealed an almost nonoverlapping distribution of both groups linked to distinct climatic regimes, particularly during interglacials, which may have reinforced the divergence of the two lineages through differential adaptation to their respective environments. This was in accordance to the results of partial Mantel tests and MMRs, showing significant positive correlations for the present interglacial between genetic distance and temperature variables, both when considering the whole species range or the *Western* and *Eastern* samples separately (Table 2). Similar correlations were also found for the last interglacial, with temperature variables remaining as significant predictors of genetic distance after accounting for geography at the species level (Table 3). During the last glacial maximum, however, we did not find any significant positive association between climate and present-day patterns of genetic differentiation (Table 3).

Contrary to previous ecological studies highlighting the importance of water availability on *T. baccata* demographic processes, such as regeneration success (Sanz *et al.*, 2009) or population sex ratio (Iszkuło *et al.*, 2009), our results did not reveal a direct effect of rainfall variables (PC2) on genetic divergence, but rather pointed to a major effect of the temperature. The importance of temperature as a selective agent has been well documented in several tree species, usually linked to altitudinal, latitudinal or longitudinal clines (e.g. Jump *et al.*, 2006; Grivet *et al.*, 2011; Prunier *et al.*, 2013). Even though the IBA patterns detected in this study do not imply causality and selection cannot be explicitly tested with our current data, the importance of temperature as a selective agent on English yew is supported by common garden observations, where significant regional differences associated with temperature clines are found in growth and phenology (M. Mayol & M. Riba, unpublished data). Moreover, in a study comparable in scale to the present work (92 populations), we found a significant association between sex-ratio and temperature, but western (Western Mediterranean and British Isles) and eastern (Central and Northern Europe) populations were clearly clustered into two distinct groups (see Fig. S8, after Berganzo, 2009), suggesting the existence of two evolutionary lineages adapted to contrasted temperature ranges. This gives additional support to the role of climate-driven adaptation in the divergence of *Eastern* and *Western* groups after initial isolation in allopatric refugia.

Several studies have reported the joint influence of isolation by distance and environmental adaptation to promote genetic divergence of plant populations (Lee & Mitchell-Olds, 2011; Temunović *et al.*, 2012; Mosca *et al.*, 2014). For example, both

geological and climatic changes during the Pliocene and Pleistocene have been proposed to explain the divergence of lineages of some conifers of the Qinghai-Tibet Plateau, such as *Taxus wallichiana* (Liu *et al.*, 2013) or *Picea likiangensis* (Li *et al.*, 2013). Nevertheless, none of them have reported evidence of a differential contribution of warm and cold periods of the Quaternary in generating genetic divergence of populations or groups through geographic isolation and/or climatic adaptation. Our results suggest that environmental factors during warm interglacials could have been crucial in shaping genetic variation of English yew. The correlation of last interglacial (LIG) climate with present genetic variation also supports that the effects of past climate on genetic variation can persist for many generations, as already reported for other long-lived trees (Ortego *et al.*, 2012; Gugger *et al.*, 2013). However, unravelling the exact contribution of different interglacials (Present (PRE), LIG) on environmentally driven isolation is challenging, mainly because of the absence of an extensive fossil record. Although we cannot discard temporally varying selection, there is some evidence suggesting that adaptive processes would most likely have occurred during the last interglacial. Palaeoecological records indicate that English yew was much more abundant than today during interglacials preceding the last glaciation (e.g. Turner, 2000; de Beaulieu *et al.*, 2001; Koutsodendris *et al.*, 2010). After the Eemian (c. 115 000–130 000 yr BP), *Taxus* is generally scarce in most of the European pollen records, suggesting a strong and continuous decline in its distribution. Molecular data also support strong reductions in effective population size starting between 100 000 and 300 000 yr BP, and continuing up to the present in the Iberian Peninsula (Burgarella *et al.*, 2012). Because large effective population sizes are expected to favour selection processes in relation to drift (Kimura *et al.*, 1963; Charlesworth, 2009), environment-driven adaptation seem to be more likely in the past, when larger effective population sizes of *T. baccata* would have enhanced the efficiency of selection.

In conclusion, our results provide a distinct perspective for the climatic impact of Quaternary glaciations, suggesting that, despite being substantially shorter, selective pressures during interglacials could have had additional impacts on population genetic divergence to those of (extensively reported) geographical isolation during glacial periods. This opens new lines of research to explore the effect of Quaternary climatic factors on the present-day patterns of genetic diversity in other long-lived organisms.

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Supporting Information

Additional supporting information may be found in the online version of this article.

Fig. S1 Geographical location of the 12 different sets of populations used for ABC simulations.

Fig. S2 Pre-evaluation of scenarios and prior distributions.

Fig. S3 Geographical distribution of the chloroplast haplotypes detected in the *trnS-trnQ* and *trnL-trnF* intergenic spacers.

Fig. S4 Summary of the clustering results using TESS for $K=2$, and STRUCTURE for $K=3$ and $K=4$.

Fig. S5 Prior and posterior distributions in the ABC analysis.

Fig. S6 Principal component analysis (PCA) plot of environmental variables for the present time described in Table S4.

Fig. S7 MAXENT predicted suitability for *Taxus baccata* at the whole range for present (PRE) and past (LGM, LIG) climatic conditions.

Fig. S8 Relationship between sex-ratio and temperature.

Table S1 Location, sample size (N), genetic diversity (A_R , H_E , H_O) and genetic cluster assignment of the studied populations. Retained populations for correlations with past climate are indicated (R)

Table S2 Sampled populations and polymorphic sites for the *trnS-trnQ* intergenic spacer

Table S3 Sampled populations and polymorphic sites for the *trnL-trnF* intergenic spacer

Table S4 Bioclimatic variables and standardized loadings for the two first axes of the PCA analysis (present climate)

Table S5 Pairwise F_{ST} values corrected for the possible presence of null alleles using the program FREENA

Table S6 Analysis of molecular variance (AMOVA)

Notes S1 Details and results of model checking and confidence in scenario choice.

Notes S2 Species distribution models and correlations between genetic distance (F_{ST}) and environmental variables obtained using the 'BIOCLIM' algorithm implemented in DIVA-GIS v7.5.

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