

Regeneration Dynamics in Fragmented Landscapes at the Leading Edge of Distribution: *Quercus Suber* L. as A Study Case

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Abstract

Aims

We studied the regeneration dynamics of woodlands and abandoned old fields in a landscape dominated by *Quercus suber* in its lower limits for rainfall and temperature. Two hypotheses were established: (1) recruitment of *Q. suber* is restricted more by abiotic variations than other species adapted to more extreme Mediterranean conditions; and (2) decreases in precipitation reduce growth, but temperature positively affects growth in the leading cold edge of this species distribution area.

Methods

We selected nine sites containing forest stands and old fields with and without tree remnants, and analyzed stand structure, soil parameters and tree growth.

Results

Succession was arrested in plots without tree remnants after cultivation abandonment. By contrast, remnant trees were accelerators of forest recovery. Tree cover played a fundamental role in *Quercus* recruitment throughout seed dispersal and facilitation effects that ameliorate summer drought. However, soil variables also significantly explained much of the variance observed and are important for understanding differences in regeneration. Winter and spring precipitation exerted a positive effect on tree growth, as well as temperatures during winter/spring and September.

Conclusions

Regeneration dynamics are modeled by the density of tree cover in the cold and dry edge of the distribution area of *Q. suber* where *Q. ilex* is increasing in abundance. Although temperature has a positive effect on the tree growth of *Q. suber*, when demographic processes are considered, decreases in water availability likely play a critical role in *Q. ilex* recruitment. This in turn changes dominance hierarchies, especially in abandoned areas with little or no tree cover.

Introduction

Human activity has influenced the structure and dynamics of forests in the Mediterranean region over centuries (Barbero et al. 1990; Scarascia-Mugnozza et al. 2000; Chauchard et al. 2007; Camisón et al. 2015). This is the case of forests and woodlands dominated by the cork oak, *Q. suber* L., managed systems that are protected by the European Union (Habitat directive 92/43/EEC). *Q. suber* is an evergreen tree species present in the western Mediterranean region that extends through the Iberian Peninsula to the western rim of the Italian peninsula, as well as some Mediterranean islands (Corsica, Sardinia, Sicilia and Balearic Islands) and coastal plains and hilly areas in North Africa, ranging from Morocco to Tunisia (Blanco et al. 1997; Magri et al. 2007; Pausas et al. 2009a). Covering about 2.2 million hectares, the current distribution area of this species is rather patchy, suggesting that in the past there was a more

continuous distribution and that much of what we see today is relictual (Blanco et al. 1997, Carrión et al. 2000, Sánchez-Palomares et al. 2007, Pausas et al. 2009a, Jovellar et al. 2010). Additionally, intensification of ongoing climate change is expected to increase temperatures and the length of dry spells in the Mediterranean basin (Kovats et al. 2014). Consequently, areas suitable for cork oak are predicted to become generally reduced over the twenty-first century, owing also to intermediate and high CO₂ emission scenarios (Pereira et al. 2009; Vesella et al. 2017).

Cork oak can form closed woodlands, but in most parts of its geographic distribution in the Iberian Peninsula and North Africa it usually appears in open woodlands with other dominating or co-dominating *Quercus* (*Q. ilex*, *Q. faginea*) or *Pinus* (*P. pinea*, *P. pinaster*) species (Blanco et al. 1997; Bugalho et al. 2009). Moreover, these open woodlands have been traditionally managed as agro-silvopastoral systems providing goods such as pasture for livestock, acorns for the high-quality pork industry, cereal crops, firewood, cork and game species. However, these woodlands also provide services in the form of recreational tourism, as cork oak systems represent an important cultural heritage in the Mediterranean region (Bugalho et al. 2009; Ovando et al. 2009; Vallejo et al. 2009). In addition, in many parts of Europe, since the second half of the twentieth century, the Iberian Peninsula has been subjected to substantial land abandonment due to socioeconomic factors. These factors have led to the decline and/or disappearance of traditional management systems, crop abandonment, shrub encroachment and an increased risk of fire (Bugalho et al. 2009; Vallejo et al. 2009). The recruitment of oaks after land abandonment is usually a slow process, especially in shrublands, where succession is strongly delayed or almost halted in a condition known as arrested succession (Pons and Pausas 2006; Acácio et al. 2007; Pausas et al. 2009ab, Acácio and Holmgren 2014). Regeneration of *Q. suber* from seed shares problems and limitations that are similar to those experienced by other *Quercus* species in Mediterranean woodlands. These adverse conditions can include: (1) High rates of acorn predation by invertebrates and vertebrates (Branco et al. 2002; Pons and Pausas 2006; Pausas et al. 2009b). (2) Limited dispersion by scatter-hoarding rodents (Pons and Pausas 2007a), since the main long-distance vector the Eurasian jay, *Garrulus glandarius* (Pons and Pausas 2007b), is a forest species that is very scarce or absent in open woodlands (Pons and Pausas 2006). (3) High seedling mortality due to water stress during the summer in open areas that dramatically restricts its establishment under the plant canopy (Castro et al. 2006, Caldeira et al. 2014; Ibañez et al. 2015), with stronger nurse plant facilitation effects as environmental stress increases (Pugnaire and Luque 2001; Maestre et al. 2009, Costa et al. 2017). And (4), reduced seedling growth and high mortality by herbivore defoliation and trampling (Gómez 2003; Silla and Escudero 2006; Rossetti and Bagella 2014, Costa et al. 2017).

In this work, we took advantage of the heterogeneity of the landscape generated by anthropogenic activities in the lower ecological limits of rainfall and temperature supported by *Q. suber* as a representative case to study the regeneration dynamics and growth at the leading distribution edge. This area is expected to be highly vulnerable to changes in traditional management and can be useful for detecting early warnings signs. As such, we have established two main hypotheses. This first hypothesis states that recruitment of *Q. suber* is more restricted by the abiotic variations induced by the dominant

elements of vegetation (nursing effects) than other species more adapted to extreme Mediterranean conditions (e.g.: *Q. ilex*). Therefore, we expected that *Q. suber* recruitment would respond more drastically to changes in vegetation structure through the landscape, performing worse than *Q. ilex* in harsh environments like open old fields. To test this hypothesis, stand structure, soil parameters and spatial patterns of trees and saplings were analyzed in three types of landscape use. The second hypothesis states that tree growth is driven by local climatic conditions (mainly precipitation and temperature) affected by climate change, and that a decrease in precipitation reduces tree growth in water limited regions. We therefore predicted that temperature would positively affect tree growth in the leading cold edge of its distribution.

Material And Methods

Study site

The study area is located in the north subdivision of the Central plateau of the Iberian Peninsula (Fig. 1a; 41° 07' N, 5° 47'W; 800-850 m a.s.l.). The mean annual precipitation is around 380 mm, with a typical Mediterranean period of low precipitation during July and August. The mean annual temperature is around 12°C, with mean temperatures between 3-4°C and 20-21°C during the coldest and the warmest months, respectively. This area contains the biggest *Q. suber* woodland in the northwest part of the Iberian Peninsula, with an extension of almost 2,000 ha (Guerra Velasco 2015). The study area is located in the distributional edge of this species, as *Q. suber* typically occurs between 0 and 800 m a.s.l., it requires an annual precipitation between 600 and 1000 mm and an average temperature around 15°C (Blanco et al. 1997, Sánchez-Palomares et al. 2007, Houston Durrant et al. 2016).

The study area is characterized by a variety of land covers, with open and closed woodlands (dominated by *Q. suber* and the presence of *Quercus ilex* subsp. *ballota*, *Q. faginea* and *Q. pyrenaica*), *Pinus pinaster* plantations, shrublands dominated by *Cistus*, *Halimium* and *Cytisus* species, old fields and abandoned vineyards.

Plot establishment and data collection

Nine sites were selected encompassing three forest stands dominated by *Q. suber*, three old fields with tree remnants and three old fields without tree cover. Site selection was based on changes in landscape use determined by comparing aerial photographs of the Spanish Inter-ministry flight from 1973-1986 (<https://fototeca.cnig.es>) and aerial photographs taken from Google Earth from 2015 (Fig. 1bc). The sites located in old fields, with and without tree remnants, had been cultivated in the 70s and 80s and abandoned in the late 1990s. The forest stands were non-cultivated areas managed for traditional cork extraction, which is still harvested today. In each site, we established a 40 x 40 m plot (1600 m²) where the origin point of each one was randomized and their sides were oriented in the directions of the cardinal points. All data were collected between October 2015 and June 2017.

All trees, both live and dead, and the saplings and seedlings in each plot were recorded. Trees were defined as individuals with a diameter at breast height (dbh) ≥ 5 cm and saplings as individuals with a dbh < 5 cm and height > 200 cm. Small seedlings were defined as individuals < 50 cm in height and large seedlings were defined as individuals ≥ 50 and ≤ 200 cm in height. Resprouting individuals were not considered, and in some doubtful cases the soil surrounding each plant was excavated to check for independence from the nearest plants. The positions of the trees and saplings were located to the nearest centimeter using measuring tapes that were aligned with the sides of the plots, providing X and Y coordinates in a Cartesian plane. Shrub cover (%) was measured sampling five line transects (5 m in length) per plot.

Dendrochronological analysis

Increment cores from all trees were extracted using Pressler increment borers (Häglof, Sweden) at 0.3-0.4 m above ground level to obtain the most accurate age for each tree (Veblen 1992), and at 0.6 m when the tree centers were rotten. Increment cores were mounted and sanded following the procedure established by Stokes and Smiley (1968), and the annual rings were counted using a stereomicroscope (SMZ800, Nikon, Japan). When the cores did not reach the pith, the number of rings to the center was estimated using the geometric procedure described by Duncan (1989). If the center was rotten, the rings counted in the non-rotten section of the core were considered as the minimum age for that tree. Cores were scanned at 2000 dpi resolution (Perfection V550, Epson, Japan), and tree-ring widths were measured with a 0.01 mm resolution on the scanned JPG images using the software CooRecorder 7.6 (Cybis, Sweden). The visual and statistical cross-dating of the tree-ring width series was done and checked using the software CDendro 7.6 (Cybis, Sweden) and Cofecha (Holmes 1983), respectively. The tree-ring width series were detrended to obtain mean residual chronologies of the tree-ring width indices. First, a spline function was fitted to each tree-ring width series to obtain standardized indices which preserve the high-frequency variability potentially related to climate. Second, an autoregressive model was applied to remove the first-order temporal autocorrelation in the detrended series and generate residual indices. Third, a biweight robust mean was computed to produce residual mean chronologies for each species. Detrending residual chronologies were obtained with the package *dplR* (Bunn 2008) under R environment (R Development Core Team 2018).

Climate data

Climate data spanning the 1948-2019 time frame were provided by the Meteorological State Agency (AEMET) from the nearest meteorological station located 30 km southeast and at the same altitude as the study site (Matacán; 40° 94' N, 5° 50'W; 790 m a.s.l.).

Soil analysis

Five soil subsamples per plot were taken using a soil core sampler at depths between 0 and 30 cm. One subsample was taken in the center of the plot and four subsamples were taken 20 m apart from the center towards each corner of the plot. The subsamples were pooled in a single sample for structural and

chemical soil analysis. The samples were air-dried and passed through a 2-mm sieve before laboratory analysis.

The pH was determined in distilled water (in a ratio 1:2.5) using a CRISON micropH 2001 pH-meter. Soil texture was quantified by the Robinson's pipette method after previous dispersion with sodium hexametaphosphate (Loveland and Whalley 1991). Organic carbon was determined by wet oxidation with a dichromate-sulphuric acid mixture (Walkley 1947). Residual dichromate was back titrated using ferrous sulphate. The difference in added FeSO_4 compared with a blank titration determined the amount of easy oxidizable organic carbon (Walkley 1947). We used a conversion factor of 1.72 to convert organic carbon to organic matter (Nelson and Sommers 1996). Available Ca, Mg and K were extracted using ammonium acetate 1M and determined by plasma ICP-MS. P was determined by the Bray I method, modified from Bray and Kurtz (1945).

Statistical analysis

To analyze how seedling regeneration (small seedlings, large seedlings and saplings) could be explained by soil and structural stand variables, a Redundant Analysis (RDA) was performed (Borcard et al. 2018). Previously, seedling data was transformed using a *hellinger* transformation, as suggested by Legendre and Gallagher (2001) for abundance data. To avoid collinearity within sets of soil and structural explanatory variables, we examined the correlation between variables and groups of variables using a hierarchical cluster analysis approach (complete linkage method). When the correlation between variables or groups of variables was greater than 0.6, only one variable was selected, which resulted in variance inflation factors (VIF) always less than five units within the set of variables (Quinn and Keough 2002). For the soil variables we selected organic matter, P (%), pH and sand content (%), and for the structural stand variables we selected basal area and tree density (Table S2, Figs S1 and S2). Then, we performed a variation partitioning analysis (9999 random permutations) to analyze how much of the variance in seedling regeneration was explained by these variables (Borcard et al. 2018). Variation partitioning and RDA were analyzed with the *vegan* library under the R environment (R Development Core Team 2018), whereas *HH* library was used to evaluate variance inflation.

For each plot, the univariate spatial patterns of all seedlings were analysed using the O-ring statistic derived from the pair correlation function (Wiegand and Moloney 2014). The pair correlation function is the expected number of points between the largest and smallest radii of a ring of fixed width at increasing distances from an arbitrary point, divided by the intensity k of the pattern (Diggle 2003, Wiegand and Moloney 2014). We used the complete spatial randomness model (CSR) implemented as a homogeneous Poisson process to determine whether the distribution of trees or saplings was random, aggregated or regular. For this univariate analysis, $O(r) > 1$ indicates that the individuals are aggregated at distances r , while $O(r) < 1$ means they are regularly dispersed. Additionally, the spatial relationships between seedlings and trees (of the same species and all *Quercus* trees pooled together) were analysed using the bivariate O-ring statistic. We used the antecedent condition as the null model, keeping the tree positions fixed whereas the seedlings were randomized using a CSR model (Wiegand and Moloney

2014). Values of $O_{12}(r) < 0$ indicate repulsion between the two classes of individuals up to distance r . By contrast, values of $O_{12}(r) > 0$ indicate attraction between the two classes up to distance r . To evaluate the significance of the spatial statistics under the null model considered, 95% of simulated envelopes were generated using 999 Monte Carlo simulations. The twenty-fifth highest and lowest values of the 999 iterated functions were chosen to obtain the upper and lower values of the envelopes. All spatial analyses were performed using the 2014 version of the software Programita (Wiegand and Moloney 2014).

Bootstrapped Pearson's correlation functions were calculated between the mean residual chronologies of each tree species and monthly climatic variables (mean, minimum and maximum temperature, precipitation). Confidence envelopes were obtained by calculating 1000 bootstrap samples and tested for significance using the 95% percentile range method (Dixon 2001). The climatic window for these analyses spanned from the previous September (year $t-1$), i.e. prior to the year of tree-ring formation, up to the following September (year t). This analysis was performed with the function *dcc* implemented in the package *bootRes* (Zang and Biondi 2013) under R environment (R Development Core Team 2018). The trends in monthly or seasonal climatic data were analyzed using the Kendall τ statistic with one-tail tests, as we were interested in decreasing rainfall and increasing temperature patterns.

Results

Woodlands showed tree densities of 95.8 ± 29.2 trees ha^{-1} and basal areas of 11.84 ± 1.34 m^2 ha^{-1} (mean \pm se), where *Q. suber* was the dominant tree species. Mean tree dbh was 29.9 cm (confidence interval of 24.7-36.2 cm at 95% level). *Quercus* species showed a broad and multiage structure, but with most of the trees recruiting between 1930 and 1980 (Fig. 2a). Old fields with tree remnants showed lower tree density and basal area than woodlands (62.5 ± 12.5 trees ha^{-1} , 5.20 ± 1.95 m^2 basal area ha^{-1} , mean \pm se), and with most of the trees (66.7% of total trees) recruiting after 1970 (Fig. 2b). Mean tree dbh was 19.6 cm (confidence interval of 16.1-23.2 cm at 95% level). *Q. suber* was also the main species in the old fields with tree remnants, although *Q. ilex* dominated tree recruitment after 1990 (Fig. 2b).

In relation to seedling recruitment, oak recruitment was quite abundant in the woodlands (2410.4 ± 719.8 seedlings ha^{-1} , mean \pm se) and dominated by *Q. suber*, especially in the shortest height classes (Fig. 3a). Significant differences in the mean recruitment of seedlings were observed between species and height classes, with clear differences in the shortest categories but insignificant ones in the highest categories (Table 2). In the old fields with tree remnants, oak recruitment was moderately abundant (1266.7 ± 445.6 seedlings ha^{-1} , mean \pm se, Fig. 3b) and seedling recruitment varied between species and height class (Table 2). By contrast, *Q. ilex* was the only species that recruited in the old fields without tree cover, although with low densities (33.3 ± 10.4 seedlings ha^{-1} , mean \pm se, Fig. 3c). Sapling density of *Quercus* species was low in woodlands (66.7 ± 29.6 saplings ha^{-1}) and old fields with tree remnants (43.7 ± 64.9 saplings ha^{-1}) and zero in old fields without tree remnants. Differences in sapling density were found between forest types (df= 2, F= 7.06, $p= 0.005$) but no differences were found between species (df= 2, F= 0.62, $p= 0.56$).

In woodlands and old fields with tree remnants, small seedlings of *Q. suber* showed a clear clumped spatial pattern at distances up to 8-12 m and were associated to *Quercus* trees, with more seedlings than expected under a random process up to distances of 5-7 m from each tree (Table 1). Small seedlings of *Q. ilex* presented a clumped pattern, but at greater distances in old fields with tree remnants than in woodlands. *Q. ilex* small seedlings were also associated with trees but at lower distances than *Q. suber*, especially in woodlands (Table 1). By contrast, large seedlings of both *Q. suber* and *Q. ilex* mostly showed a random pattern distribution and were independently distributed from trees in woodlands whereas in old fields with tree remnants only large seedlings of *Q. suber* showed association with trees at distance of 4-7 m (Table 1).

In the RDA, the first axis represented a contrast between old fields without tree cover at the right, where only small seedlings of *Q. ilex* were present, and woodlands at the left, where small and large seedlings of *Q. suber* dominated tree regeneration (Fig. 4). Basal area, pH and soil organic matter showed a strong correlation with the first axis, with higher values of these three variables associated with woodlands. The second axis was related to the abundance of *Q. faginea* with regard to tree regeneration, with the sites with the higher scores in the upper panel showing a higher abundance for this species (Fig. 4).

Variance partition analyses showed that soil and structural variables explained most (81.0%) of the variance in the abundance of regeneration across sites (Fig. 5). Shared explained variation for both sets of variables was 31.6%, with soil variables contributing slightly more to the unshared explained variation in seedling abundances than structural variables (soil variables: 27.1%, $F = 3.85$, $p = 0.014$, permutations = 999; structural stand variables: 22.3%, $F = 5.67$, $p = 0.004$, permutations = 999).

Ring growth was positively correlated with rainfall in January and June (Fig. 6a), and September temperatures (mean, maximum and minimum) and minimum winter-early spring temperatures also positively influenced ring growth (Fig. 6bcd). Monthly rainfall in January and June did not show a decreasing trend over time (Table S1), although annual rainfall significantly decreased (Table S1, Fig. S3). All mean, maximum and minimum temperatures in January, February, March and September did not show temperature increases over time, except for the minimum temperature in September (Table S1).

Discussion

a) Regeneration dynamics of *Quercus*

Succession was considerably arrested in the plots without tree remnants two to three decades after cultivation abandonment. Similar trends have been observed in other studies, where 45 years after abandonment *Cistus*-dominated shrublands prevailed with scarce or absent oak recruitment (Acácio and Holmgren 2014). *Cistus*-dominated shrublands, as well as acorn availability and drought stress, exert competition and inhibition effects over *Quercus* recruitment and appear as highly resilient systems (Pérez-Devesa et al. 2008, Rolo and Moreno 2011, Acácio and Holmgren 2014). However, shrub cover (mainly characterized by *Cistus*, *Halimium* and *Lavandula* spp.) was quite low in plots in old fields (< 10%, Table S2) and vegetation was dominated by annual herbaceous species, likely due to low rainfall

combined with high sand and low organic soil contents that exacerbated hydric stress (Fernández-Ales et al. 1993, Rawls et al. 2003, Nunes et al. 2017). *Q. ilex* was the only tree species successfully recruiting in the open old fields, although slowly and with low densities, despite *Q. suber* being the dominant tree species in the landscape with respect to density and basal area. Two processes are likely involved in the success of *Q. ilex* over other *Quercus* species in the open plots. First, although *Quercus* species share the same animal dispersers, most studies have shown that acorns from *Q. ilex* are preferred over other *Quercus* species by the main oak disperser the Eurasian jay (*Garrulus glandarius*) and by small rodents (Pons and Pausas 2007ac, del Arco et al. 2018). Since the Eurasian jay is a forest species that has never been detected in the study area, mice species are the main candidates for acorn dispersal. Pilferage rates are reduced by caching the seeds preferentially outside the canopies of scattered trees, increasing the survival of cached acorns in open areas (Muñoz and Bonal 2011). However, limited facilitative shrub cover reduces successful recruitment causing a low density of *Q. ilex* seedlings (Pulido and Díaz 2005, Smit et al. 2009, Rolo and Moreno 2011). Second, although both evergreen species, *Q. suber* and *Q. ilex*, are well adapted to the summer drought of the Mediterranean climate (Mediavilla and Escudero 2003, González-Rodríguez et al. 2011, San-Eufrasio et al. 2020), previous studies have shown a higher survival rate for *Q. ilex* seedlings than for *Q. suber* (Plieninger et al. 2010, González-Rodríguez et al. 2011). In addition, *Q. ilex* has lower conductance and maximum transpiration rates than *Q. suber*, which delays leaf desiccation thanks to a more conservative use of water (Mediavilla and Escudero 2003, San-Eufrasio et al. 2020), which decreases growth suppression under high water stress (Caldeira et al. 2014).

Our results indicate that remnant oak trees are great accelerators of forest recovery after cultivation abandonment. Also, old fields with tree remnants boost oak regeneration almost 40-fold in comparison to old fields without trees, and woodlands show a 2-fold increase in oak regeneration in comparison to old fields with tree remnants. Tree cover is considered to play a fundamental role in *Quercus* recruitment through several processes. First, acorns are heavy fruits with limited dispersal, where most of the acorns produced only reach areas situated close to the parent trees (Pulido and Díaz 2005, Acácio et al. 2007, Pausas et al. 2009b). Second, facilitation effects caused by tree cover are key in the regeneration of *Quercus* species, especially in Mediterranean environments (Caldeira et al. 2014, Costa et al. 2017). Facilitation includes several direct and indirect mechanisms with positive effects on both survival and *Quercus* seedling growth. Drought stress and summer seedling survival during the first years of establishment is usually considered one of the main bottlenecks in *Quercus* regeneration (Díaz and Pulido 2005, Silla and Escudero 2004, Smit et al. 2009, Pérez-Ramos et al. 2012). Therefore, tree cover improves microclimate conditions reducing high summer temperatures and alleviating heat stress (Díaz and Pulido 2005, Pausas et al. 2009, Smit et al. 2009) and reduces competition with herbaceous vegetation (Caldeira et al. 2014). Our results are also consistent with these patterns, as tree basal area and density were strongly correlated with the first RDA ordination axis that ordered sites with a strong gradient of seedling density from right to left (Fig. 4). Additionally, the spatial analysis of small seedlings of *Q. suber* and *Q. ilex*, which accounted for most of the seedling abundance (71.6%) showed a clustered pattern associated with *Quercus* trees in both woodlands and old fields with tree remnants, although at shorter distances in *Q. ilex* than in *Q. suber*. By contrast, *Quercus* large seedlings showed a random

spatial pattern independent of the trees, highlighting that tree cover is only a limiting factor during seedling establishment. These changes in the spatial pattern with ontogenetic development support previous findings that suggest that the positive effects of shaded microhabitats are reversed for seedling development (Pérez-Ramos et al 2011, Pausas et al. 2009b). In *Q. suber* and *Q. ilex*, seedling establishment and survival are improved under shade (Espelta et al. 1995, Pausas et al. 2009b, Smit et al. 2009, Pérez-Ramos et al. 2012). However, it has been also shown that low light suppresses growth, that these species establish 'seedling banks' under dense tree cover and that more open canopy conditions are required for saplings and trees to develop (Espelta et al. 1995, Pausas et al. 2009b, Pérez-Ramos et al. 2010, 2012). In addition, we also found significant differences in the regeneration densities of *Q. suber*, *Q. ilex*, *Q. faginea*, with the abundance of *Q. suber* strongly correlated with the basal area and tree density of stands. In woodlands, *Q. suber* dominated seedling regeneration in the categories with the shortest height, but the differences in seedling abundance between *Q. ilex* and *Q. suber* disappeared in the tallest categories and showed similar densities. In the old fields with tree remnants, *Q. ilex* and *Q. suber* showed similar seedling densities, and although the interaction was not significant, the abundance of *Q. ilex* large seedlings was 5-6 fold higher than that of *Q. suber*. These results indicate that *Q. ilex* has a higher seedling survival rate and more likely to reach the sapling stage, probably due to their higher tolerance to shade (Sevilla 2008), and in particular their greater tolerance to hydric stress during the summer (Plieninger et al. 2010, González-Rodríguez et al. 2011, San-Eufrasio et al. 2020). Thus, although *Q. suber* trees dominate the landscape of our study area, the results indicate that *Q. ilex* can produce a greater number of young trees as compared to *Q. suber* (Fig. 2b), owing to the better performance of *Q. ilex* seedlings. Only in woodlands can *Q. suber* partially compensate for their lower performance and survival, producing a higher number of seedlings. Conversely, the deciduous *Q. faginea* present the lowest abundance, especially in the old fields (with or without tree remnants), which is in line with its lower tolerance to drought compared to evergreen species (Silla and Escudero 2004, González-Rodríguez et al. 2011, San-Eufrasio et al. 2020).

The presence of tree remnants showed a strong effect over soil parameters with higher concentrations of organic matter, N, exchangeable cations (K^+ , Ca^{2+} , Mg^{2+}) and slightly less acidic soils than in old fields without tree remnants (Table S2). The deep root system of *Q. suber* trees uptakes and pumps basic cations, especially Ca^{2+} , from the deep to upper soil layers throughout litterfall production and decomposition, which significantly improves soil nutrient availability (Serrasolses et al. 2009, Rossetti et al. 2015). The results of the variance partitioning showed that soil and stand structural parameters explain a significant amount of the shared variation in the amount of regeneration of *Quercus* species (Fig. 5), which makes sense since the influence of trees on soil characteristics are spatially correlated. However, soil variables also explain a significant amount of the variance (27.1%) not explained by structural stand variables and are important for understanding the differences found in *Quercus* regeneration between plots. Among soil variables, organic matter content summarizes most information on nutrient availability with which is strongly correlated (Table S2, Fig. S1), but it is also directly involved in improving soil water retention in abandoned cultivars, especially in soils with high sand content (Rawls et al. 2003, Costa et al. 2017).

b) Climate-growth relationships of *suber* trees

Analysis of the impact of climatic conditions on the tree-ring growth of *Q. suber* is challenging due to the difficulty in identifying ring boundaries in trees being managed for cork extraction (Costa et al. 2003). Only two previous studies have reported successful tree ring chronologies involving *Q. suber* in the Mediterranean region (Costa et al. 2003, Zribi et al. 2016) that have been complemented with short chronologies from cork growth-rings (Caritat et al. 1996, Costa et al 2016, Leite et al. 2019). As shown in these studies, winter and spring precipitation exerted a large positive effect on tree-ring growth due to the replenishment of soil water reserves before the onset of the favorable growing season (Costa et al. 2001, Jovellar et al. 2010, Costa et al 2016, Zribi et al. 2016, Leite et al. 2019). In our study site, *Q. suber* showed the latest leaf emergence of all coexisting *Quercus* species, with budbreak and emergence of the new leaf cohort occurring between the end of May and the beginning of June (del Río et al. 2015). However, in relation to other studies where temperature has been found to have an insignificant or negative effect over cork or ring growth (Caritat et al. 1996, Costa et al. 2001, Zribi et al. 2016), in our study site, the mean and minimum temperatures during winter and/or spring and the mean and maximum temperatures during September exerted a positive influence on tree growth. This is consistent with the cold temperature conditions occurring in winter and early spring in our study area, located at the leading temperature edge of its distribution limits in the Iberian Peninsula. The large vessels in oak trees are very sensitive to winter embolisms caused by freezing temperatures, and in spring the reactivation of growth is greatly dependent on hydraulic conductivity recovery (Hacke and Sauter 1996, Lebourgeois et al. 2004). This highlights the importance of mild winter-early spring temperatures on *Q. suber* tree growth in the “cold leading edge” of its distribution. However, although winter temperatures are a limiting factor and warming has room for net positive effects on tree growth (Sánchez-Salguero et al., 2015), climatic data did not reveal a significant temperature increase during winter and early spring in this study site. On the other hand, minimum temperatures during September have significantly increased during the last decades and have had a positive impact on tree growth by most likely extending the favorable weather (Marqués et al. 2018). However, the positive effect of an extended growing season in early autumn can be counteracted by a decrease in rainfall, especially in the study area where annual rainfall is also in the lower edge of *Q. suber* distribution limits, as has also been recently observed in Mediterranean forest at the limits of species distribution (Madrigal-González et al. 2018, Marqués et al. 2018). Although no significant decreasing trends were found in the more critical months (Fig. 6a), there is a decreasing trend in mean annual rainfall (Fig. S3), so more detailed studies will be needed to understand the combined effects of temperature and rainfall on tree growth within future climate change scenarios.

Conclusions

Regeneration dynamics were strongly modeled by the presence and density of tree cover in the fragmented landscape in the cold and dry edge of *Q. suber* distribution. We have seen that tree cover affects seedling abundance differently through both direct (acorn availability and shading) and indirect (improving nutrient availability and water retention in soils) mechanisms. Consequently, the stress tolerant species *Q. ilex* was the only species found to recruit in open old fields decades after being

abandoned. Also, the presence of isolated tree remnants in old fields allowed the recruitment of *Q. suber*, but *Q. ilex* had a higher abundance of large seedlings and young trees; only in woodlands did both species show similar recruitment success. These results indicate that *Q. ilex* is more abundant in this landscape compared to *Q. suber*, even though the latter species is the dominant tree in our study area.

In addition, the climate-tree growth relationship of *Q. suber* presents contrasting results. Higher temperatures were found to have a positive effect on tree ring-growth, especially during winter/early spring and early autumn. On the other hand, winter and spring rainfall recharges water soil reserves promoting tree growth during the favorable season. These opposing effects increase pose uncertainty in predicting *Q. suber* growth and productivity under climate change scenarios involving higher temperatures and less rainfall. However, when the demographic processes are considered, less water availability is likely to play a critical role in favoring *Q. ilex* recruitment in contrast to *Q. suber*, which could lead to changes in dominance hierarchies, especially in abandoned areas with scarce or absent tree cover.

Declarations

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Authors' contributions

Conceptualization: Fernando Silla; Methodology: Carmen Ureña, Diego Navarro, Valentín Herrera, Pilar Alonso-Rojo, Fernando Silla; Formal analysis and investigation: Carmen Ureña, Diego Navarro, Valentín Herrera, Jorge Montero-Muñoz and Fernando Silla; Writing - original draft preparation: Jorge Montero-Muñoz and Fernando Silla; Writing - review and editing: Carmen Ureña, Diego Navarro, Valentín Herrera, Pilar Alonso-Rojo, Héctor Hernández-Alonso, M^a Fernanda Cepeda, Luis Carlos Jovellar, Belén Fernández-Santos

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Tables

Table 1. Results of the O-ring statistic for the univariate and bivariate analyses. Statistical significance at 95% simulated envelopes for the univariate analyses (grey: random pattern; black: clumped pattern; white: regular pattern) and for the bivariate analyses (grey: independence pattern; black: attraction pattern; white: repulsion pattern).

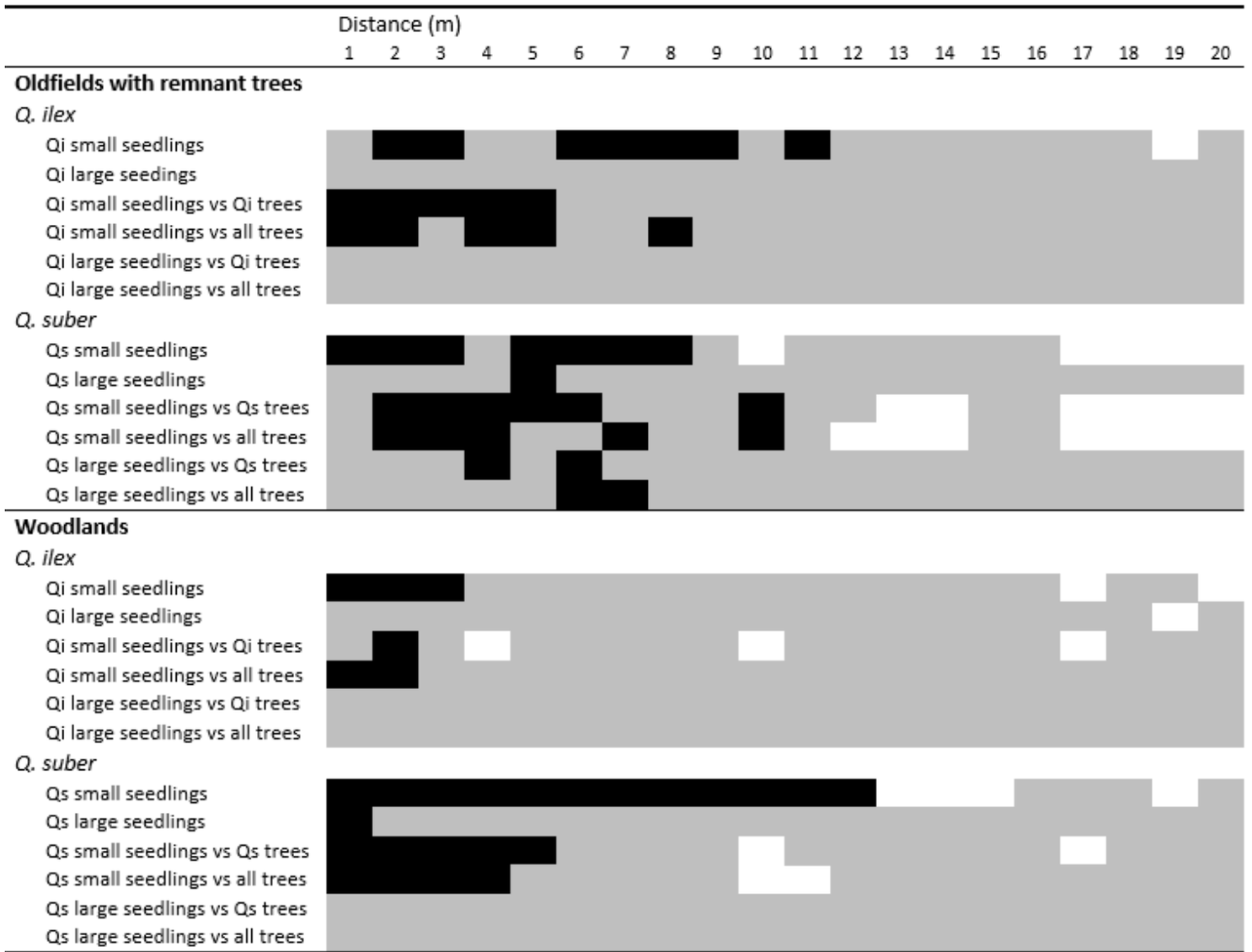


Table 2
Results of the two-way variance analysis testing for differences in seedling recruitment between species and height classes.

	Woodlands			Oldfields with tree remnants		
	df	F	p	df	F	p
Species	2	5.10	0.003	2	7.58	< 0.001
Height	4	6.09	0.006	4	5.91	0.007
Interaction	8	2.33	0.044	8	0.93	0.500