






## RESEARCH ARTICLE

# Long-term dynamics of shrub facilitation shape the mixing of evergreen and deciduous oaks in Mediterranean abandoned fields

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## Funding information

Fundación Internacional para la Restauración de Ecosistemas; University of Alcalá, Grant/Award Number: FIRE-UAH 127/2017; 'Tatiana Pérez de Guzmán el Bueno' Foundation; REMEDINAL Postdoctoral Fellowship Programme, Grant/Award Number: TE-CM S2018/EMT-4338; TALENTO Fellow Programme, Grant/Award Number: 2016-T2/AMB-1665

Handling Editor: Cristina García

## Abstract

1. Recovery of Mediterranean forests after field abandonment is a slow process, even without propagule limitations. This is mainly due to stressful conditions for seedling establishment. In this context, shrubs play a critical role in facilitating tree recruitment, but how this process unfolds after field abandonment is not entirely known. We evaluated the long-term dynamics of facilitation by the nurse shrub *Retama sphaerocarpa* in the recruitment of two ecologically contrasting oaks, the evergreen *Quercus ilex* and the deciduous *Quercus faginea*.
2. Thirty years after field abandonment, we dated shrubs and oaks established in an old field to estimate the annual recruitment rates and investigate temporal recruitment patterns. For oaks, we differentiated recruitment at two microsites: open areas or under shrub. To assess how nurse shrubs modulated environmental stressors, we modelled oak recruitment as a function of climatic variables. For the evergreen oak, we assessed these effects within each microsite. Finally, we estimated the annual interaction index between shrubs and evergreen oak juveniles as a function of climatic conditions.
3. Each species showed different recruitment pulses during colonization. Recruitment rate was the highest for the shrub, followed by the evergreen oak. Oak colonization under shrubs was appreciable 20 years after field abandonment, when shrub cover reached 2.2%, and concentrated under medium and large shrubs older than 7 years. Shrubs not only accelerated evergreen oak colonization but also attenuated the fluctuations of recruitment pulses. For the evergreen oak, the interaction index indicated facilitation dominance in years with more arid summers (precipitation < 47 mm and heat waves longer than 8 days) and competition in wetter summers.
4. *Synthesis*. Oak colonization in Mediterranean abandoned fields progressed slowly, and only two decades after abandonment pioneer shrub population grew to a level in size and number that could effectively trigger facilitation and accelerate tree recruitment. The shrub nurse effect was unbalanced between oak species, only benefiting the evergreen oak, and it was more prevalent in arid years. Our

study illustrates the build-up of shrub facilitation during forest recolonization and the varying nature of this process among climatically different years and ecologically distinct species. This information provides insights for assessing and managing Mediterranean forest recovery.

#### KEYWORDS

colonization, facilitation, forest recovery, nurse shrub, *Quercus*, recruitment pulses, *Retama sphaerocarpa*, secondary succession

## 1 | INTRODUCTION

Plants have to overcome several abiotic environmental filters to establish (Harper, 1977; Lortie et al., 2004). The balance of positive and negative biotic interactions, particularly of facilitation and competition, modulates the effect of such filters (Connell & Slatyer, 1977; Pulsford, Lindenmayer, & Driscoll, 2016). In forest succession, plant colonizers can facilitate the establishment of mid- and late-successional plant species into the community through environmental modifications (e.g. Butterfield et al., 2010). Generally, the importance of facilitation increases with environmental stress (the so-called Stress Gradient Hypothesis; Bertness & Callaway, 1994). However, competition can prevail in plant–plant interactions if environmental conditions become very stressful and the abiotic stress is resource driven (e.g. by water shortage or light competition; Maestre, Callaway, Valladares, & Lortie, 2009; Maestre, Valladares, & Reynolds, 2005).

In arid and semi-arid ecosystems such as those found in Mediterranean-climate areas, high irradiance and drought are the main abiotic factors affecting early plant establishment (Joffre, Rambal, & Damesin, 2007; Sánchez-Gómez, Valladares, & Zavala, 2006), constraining spring seedling emergence and first-summer survival (Mendoza, Zamora, & Castro, 2009; Rolo, Pliening, & Moreno, 2013). Semi-arid ecosystems also show high inter-annual climatic variation that can cause irregular recruitment pulses (Giorgi & Lionello, 2008; Kouba, Camarero, & Alados, 2012; Pérez-Ramos, Padilla-Díaz, Koenig, & Marañón, 2015; Pugnaire, Luque, Armas, & Gutiérrez, 2006). Thus, chronic water stress and inter-annual climatic variation slow down forest recovery after disturbances (Rey Benayas, Martínez-Baroja, Pérez-Camacho, Villar-Salvador, & Holl, 2015).

Under stress conditions, shrubs can ameliorate environmental stressors at low competitive costs (Gómez-Aparicio, 2009), facilitating the establishment of mid- and late-successional woody species (Smit, Díaz, & Jansen, 2009; Smit, Ouden, & Díaz, 2008). Nurse shrubs can ameliorate climatic constraints and promote soil fertility (i.e. direct facilitation; Gómez-Aparicio, Zamora, Castro, & Hódar, 2008; Prieto, Padilla, Armas, & Pugnaire, 2011; Pugnaire et al., 1996), but also attract seed dispersers and reduce herbivory or competition with other plants (i.e. indirect facilitation; Cuesta, Villar-Salvador, Puértolas, Rey Benayas, & Michalet, 2010; Perea, López-Sánchez,

& Dirzo, 2017). The facilitative effect of shrubs have been related to specific functional attributes such as plant size (Alday, Zaldívar, Torroba-Balmori, Fernández-Santos, & Martínez-Ruiz, 2016; Navarro-Cano, Goberna, Valiente-Banuet, & Verdú, 2016), the root architecture (Prieto, Kikvidze, & Pugnaire, 2010; Prieto et al., 2011; Rolo et al., 2013) and the ability to fix nitrogen into the soil (Gómez-Aparicio, Zamora, Gómez, Hódar, & Castro, 2004).

The presence of nurse shrubs could be particularly relevant in facilitating the recruitment of less stress tolerant and more palatable species (Gómez-Aparicio et al., 2008; Liancourt, Callaway, & Michalet, 2005), expanding their ecological niche and allowing the coexistence of ecologically distinct species (Bruno, Stachowicz, & Bertness, 2003). In Mediterranean forests, like our study system, evergreen and deciduous oaks frequently co-occur (Madrigal-González et al., 2017). Evergreen oaks have more robust leaves with more leaf mass per area and less N concentration than deciduous oaks (Castro-Díez, Villar-Salvador, Pérez-Rontomé, Maestro-Martínez, & Montserrat-Martí, 1997; Gil-Peigrín, Peguero-Pina, & Sancho-Knapik, 2017). These traits are usually linked to high drought tolerance and low palatability in oak species (Corcuera, Camarero, & Gil-Peigrín, 2002; Quero, Villar, Marañón, & Zamora, 2006). Thus, we could expect evergreen oaks establishing earlier than deciduous oaks during forest succession in open environments, and deciduous oaks being more dependent on nurse shrubs for their recruitment.

Most research on facilitation has been addressed by short-term studies. However, some studies have focused on the shifts of facilitation over ontogenetic stages of the nurse shrub or of the benefactor (Alday et al., 2016; Moreno-Gutiérrez, Battipaglia, Cherubini, Delgado Huertas, & Querejeta, 2015; Navarro-Cano et al., 2016; Rousset & Lepart, 2000; Torroba-Balmori, Zaldívar, Alday, Fernández-Santos, & Martínez-Ruiz, 2015), and on the covariation of seedling and shrub abundance (Ramírez & Díaz, 2008), while others have assessed the effect of inter-annual climatic variations on facilitation (Butterfield et al., 2010). Despite these efforts, as far as we know, a long-term and holistic approach evaluating how facilitative effects change after field abandonment has never been addressed. This knowledge gap hinders our understanding of the dynamics associated with facilitation in plant community recovery (Bertness & Callaway, 1994; Maestre et al., 2005). For example, we know little about how long after field abandonment the nurse shrub community

takes to start facilitating recruitment of other species, how the effects of facilitation vary among competing beneficiary species and how these effects vary with environmental conditions.

To investigate the dynamics of nurse shrub facilitation we studied colonization of a Mediterranean oak forest over 30 years after field abandonment. Specifically, we assessed the effect of the nurse shrub *Retama sphaerocarpa* (L.) Boiss. on the long-term recruitment of the evergreen oak *Quercus ilex* L. and the deciduous oak *Quercus faginea* Lam. Our objectives were to: (a) compare the long-term recruitment dynamics of the nurse shrub and the oaks after field abandonment; (b) quantify the time lag between field abandonment and the nurse shrub facilitative effect on oak colonization; and (c) evaluate the effects of the nurse shrub on the recruitment patterns of the oaks and the climatic constraints that limit their recruitment. We hypothesised that: (a) recruitment patterns follow temporal pulses that differ among pioneer nurse shrubs and mid- or late successional oaks; (b) there is a time lapse between field abandonment and the acceleration of oak recruitment linked to shrub size and the development of shrub population; and (c) oak recruitment under nurse shrubs is decoupled from recruitment in open sites due to shrub buffering of harsh climate conditions. Understanding how climate and the facilitation–competition balance drive the temporal patterns of oak recruitment will allow for more informed assessment of future forest recovery and a better design of restoration efforts.

## 2 | MATERIALS AND METHODS

### 2.1 | Study area

The studied abandoned fields are located on two flat fluvial terraces of the Jarama river in Uceda, Central Spain (700–750 m a.s.l., UTM X: 462 830; UTM Y: 4 524 000; ETRS89 30N; Figure 1). The area has a continental Mediterranean climate, with a mean annual temperature of 13°C that ranges between 23 and 4°C in the hottest and coldest months respectively. The annual rainfall is 500 mm, being c. 60 mm during the summer (Tornero Sánchez, 1998). The soils are mainly alfisols and inceptisols, with sandy loam to sandy clay loam textures (Peñuelas-Rubira et al., 1996). Agricultural use ceased in c. 1984, although occasional sheep and goat grazing has remained in the area. Currently, oak seedling mortality by herbivory represent less than a quarter of mortality by desiccation (unpublished data). Since 1985, the fields have progressively changed to a shrubland dominated

by *R. sphaerocarpa*, a leguminous shrub and a paradigmatic nurse species in the Mediterranean basin (Andivia, Villar-Salvador, Tovar, Rabasa, & Rey-Benayas, 2017; Cuesta et al., 2010; Pugnaire et al., 1996; Rolo et al., 2013). The abandoned fields are adjacent to mixed forests dominated by *Q. ilex* and *Q. faginea* in different proportions (Table S1.1). *Quercus faginea* acorn dispersal is earlier than *Q. ilex*'s (Castro-Díez & Montserrat-Martí, 1998). In 2017, when our field-work took place, the density of *Q. ilex* and *Q. faginea* juveniles were  $45 \pm 49$  and  $15 \pm 31$  individual/ha (mean  $\pm$  SD) respectively (Kruskal–Wallis  $\chi^2 = 9.17$ ;  $p = .002$ ).

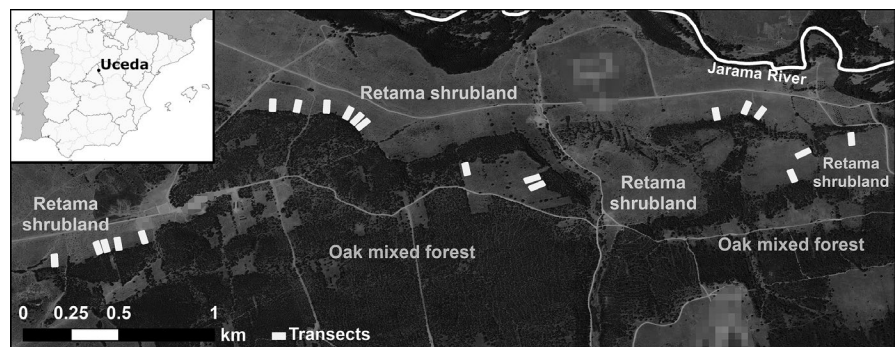
### 2.2 | Field work and oak recruitment dating

In the abandoned fields of the study site, we surveyed 20 rectangular transects (20  $\times$  60 m) perpendicular to the forest border (Figure 1). In the adjacent forest, the diameter at breast height (d.b.h.) of each oak species was measured in one 20  $\times$  35 m plot next to each transect. In each plot, the basal area was calculated using individuals with d.b.h. > 8 cm. The location of transects was selected to sample a wide range of shrub cover and abundance of the two oak species that were representative of the adjacent forest (Table S1.1).

In each transect, we located and harvested all juvenile oaks (namely 105 evergreen oaks and 36 deciduous oaks) during early spring of 2017. To harvest the juveniles, we excavated the ground around the upper part of the tap root and cut a transversal section at the root collar, that is, slightly below the ground level (Figure S2.1a). We then used a dendrochronological approach to date the year of oak recruitment (details in Appendix S2) and reconstruct the *R. sphaerocarpa* shrubland structure along time using allometric models (details in Appendix S3). The microsites where oak juveniles grew were recorded as 'open' or 'under shrub'. To ensure that we assigned the right microsite during recruitment, some records under shrub were corrected to open if (a) the oak was older than the shrub (15 cases in 7 transects –25% of all records–), or (b) the shrub was < 5 years older than the oak and the distance between the oak and shrub stems was less than the predicted crown radius of the shrub in the oak recruitment year (5 cases in 3 transects –8% of all records–).

### 2.3 | Climate data

We downloaded daily weather data for the study period (1985–2014) from nine meteorological stations close to our study area (<http://>



**FIGURE 1** Location of the study area and the 20 transects where *Retama sphaerocarpa* (retama) and oak (*Quercus ilex* and *Q. faginea*) recruitment were sampled

www.aemet.es/es/datos\_abiertos/AEMET\_OpenData; Figure S1.1). We interpolated the maximum temperature and the precipitation to our study area using the METEOLAND package (De Cáceres, Martín, Granda, & Cabon, 2018) in R 3.4.3 (R Core Team, 2017). This method interpolates climatic data to a specific area by correcting the original data according to the proximity and elevation differences between the meteorological station and the target area. For each year, we calculated the spring precipitation as the sum of April and May precipitation, the summer precipitation as the sum of July, August and September precipitation, and the length of heat waves as the maximum number of consecutive days with maximum temperature  $\geq 33^{\circ}\text{C}$ .

## 2.4 | Data analysis

### 2.4.1 | Long-term dynamics of *Retama sphaerocarpa* and oak recruitment

We calculated recruitment rate (no. recruits transect<sup>-1</sup> year<sup>-1</sup>) of *R. sphaerocarpa*, evergreen and deciduous oaks from 1985 to 2014. We used a generalized additive model and the recruitment rate of each species to analyse recruitment patterns over time. For the evergreen oak we differentiated between microsites (open and under shrub). We could not account for microsite differences for the deciduous oak due to the low number of recruits under shrubs. We choose a Poisson error distribution and the smoothing spline method to fit the models with the GAM package (Hastie, 2018) in R 3.4.3 (R Core Team, 2017). We used the Akaike Information Criterion (AIC; Burnham & Anderson, 2002) to compare models adjusted with different number of knots (i.e. data division points), and we selected the number of knots resulting in minimum AIC (e.g. in Thorson, Punt, & Nel, 2012).

### 2.4.2 | Oak colonization in relation to nurse shrub structure

We analysed the relationship between oak recruitment and the structure of the *R. sphaerocarpa* shrubland (i.e. density and size structure of shrubs) with a three-step analysis. First, we adjusted generalized additive models for the accumulated recruitment (total and per microsite) of both oak species (recruits transect<sup>-1</sup>), using the year as explanatory variable and the model details (i.e. error distribution, algorithms and number of knots) described in the previous section. Second, we identified change points over the fitted curves (i.e. the first year of a new regime of variance of accumulated recruitment) using the *cpt.var* function of the CHANGEPOINT package (Killick, Haynes, & Eckley, 2016). The change point was identified with the PELT method and the penalty set manually based on the elbow plot criterion (Figure S1.2). Change points represented acceleration in the recruitment when they separated curve segments of increasing slope. Third, we compared the change years of the accumulated oak recruitment with the change years in each microsite to assess if the acceleration of the oak recruitment was due to recruitment under

shrub (i.e. facilitated recruitment). Then, we described the density and size structure of the shrubland when facilitated recruitment occurred.

To study the relationship between oak recruitment and *R. sphaerocarpa* size, we compared (a) the maximum branch diameter of nurse shrubs (i.e. with oaks recruited under their canopies) and of shrubs without recruits using a Kruskal–Wallis test; and (b) the size structure of the *R. sphaerocarpa* populations with and without recruited oaks. For those comparisons, we only used the data of years and transects with oak recruitment under shrubs.

### 2.4.3 | Shrub modulation of the climatic effects on oak recruitment

To assess the effect of climatic variables on the evergreen oak recruitment rates, and how the nurse shrubs might have influenced this process, we used the annual oak recruitment rates estimated since 2005, which was the recruitment year of the oldest oak found under shrub. There were not enough deciduous oak recruits under shrub to make this analysis, so we assessed only the effect of the climate in total recruitment. We focused on climatic conditions during spring and summer, the most limiting periods for seedling survival (Gómez-Aparicio et al., 2004; Mendoza, Gómez-Aparicio, Zamora, & Matías, 2009). We selected climatic variables that drive spring emergence (spring precipitation in mm, SpPrec<sub>*i*</sub>) and summer survival (summer precipitation in mm, SuPrec<sub>*i*</sub>; length of heat waves in no. days, HtWaves<sub>*i*</sub>). To control for differences on seed availability among transects, we included the basal area of each target oak species in the adjacent forest (BA<sub>*i*</sub>, m<sup>2</sup>/ha). We also considered the year of recruitment as a random variable (YRE) to account for likely temporal processes such as masting, and included an intercept term for each transect to, again, account for differences among transects. For each transect *i* and year *y* the observed number of recruits per microsite (COUNT<sub>*i,y,microsite*</sub>) was analysed as:

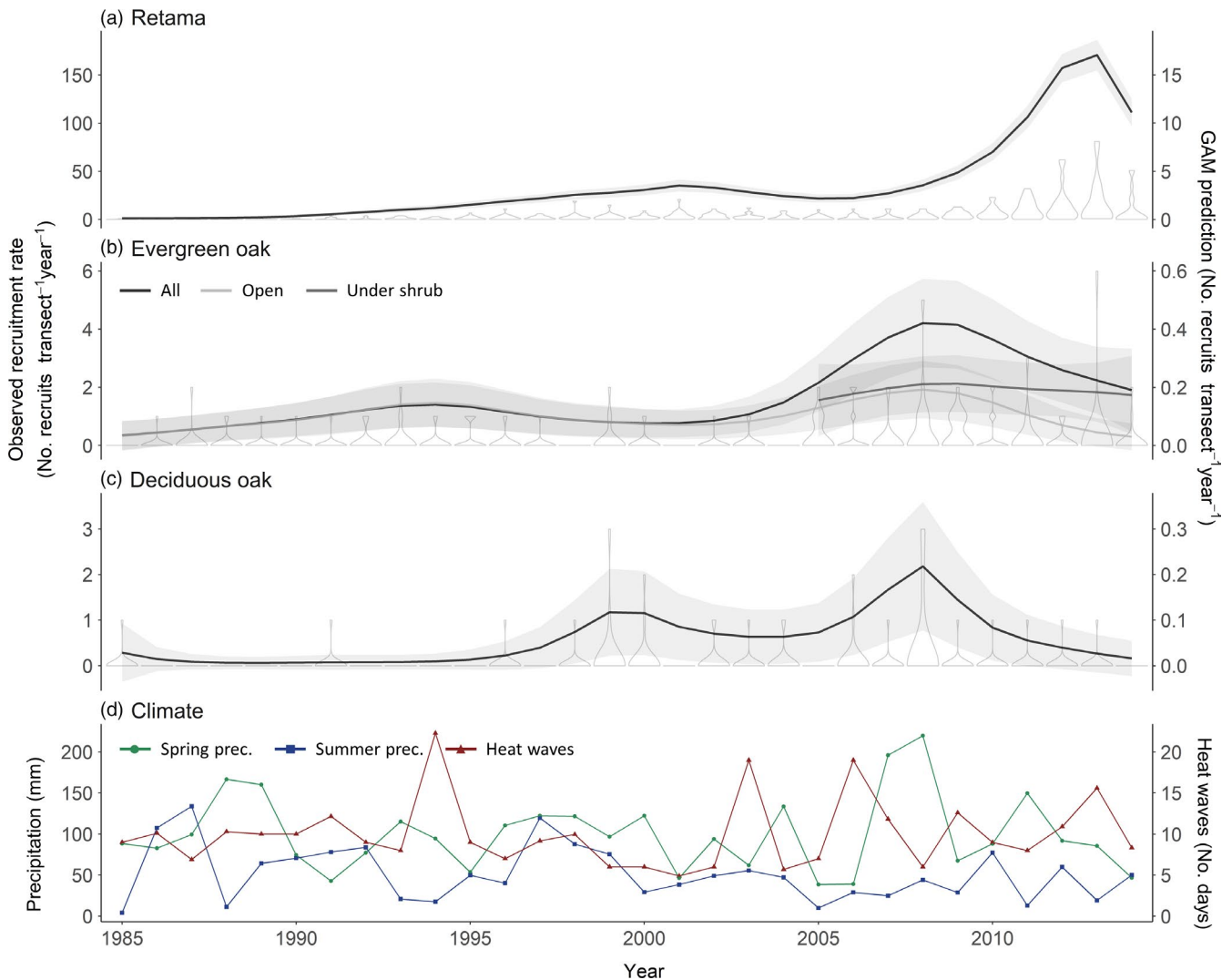
$$\text{Likelihood: Count}_{i,y,\text{microsite}} \sim \text{Poisson}(\lambda_{i,y,\text{microsite}})$$

Process model:

$$\begin{aligned} \ln(\lambda_{i,y,\text{microsite}}) = & \alpha_i + \beta \cdot \text{BA}_i + \gamma 1_{\text{microsite}} \cdot \text{SpPrec}_y \\ & + \gamma 2_{\text{microsite}} \cdot \text{SuPrec}_y \\ & + \gamma 3_{\text{microsite}} \cdot \text{HtWaves}_y + \text{YRE}_y + \varepsilon_{i,y} \end{aligned} \quad (1)$$

We also included an error term  $\varepsilon_{i,y}$  to account for over-dispersion in the recruitment rates (i.e., the variance of the number of recruits was higher than the mean; Kéry, 2010).

We estimated the parameters in Equation (1) with a Bayesian approach and non-informative hyperparameter values (Simpson, Rue, Martins, Riebler, & Sørbye, 2015). We used the RJAGS package (Plummer, 2016) in R 3.4.3 (R Core Team, 2017) and simultaneously ran three Markov chains with a burn-in period of 500,000 iterations and a sampling period after convergence of 100,000 iterations,



**FIGURE 2** Predicted (lines) and observed (violin plots) recruitment rate over time of: (a) the nurse shrub *Retama sphaerocarpa* (retama); (b) *Quercus ilex* (evergreen oak); and (c) *Quercus faginea* (deciduous oak). For the evergreen oak, predicted recruitment rates in open and under shrub microsites are also shown. Note that the scale of recruitment rate differs among species and that all violin plots are represented with the same maximum width. (d) Spring and summer precipitation and length of heat waves along the study period [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

during which we recorded the parameter thinning every 100th iteration (see Table S1.2 for results of convergence diagnosis).

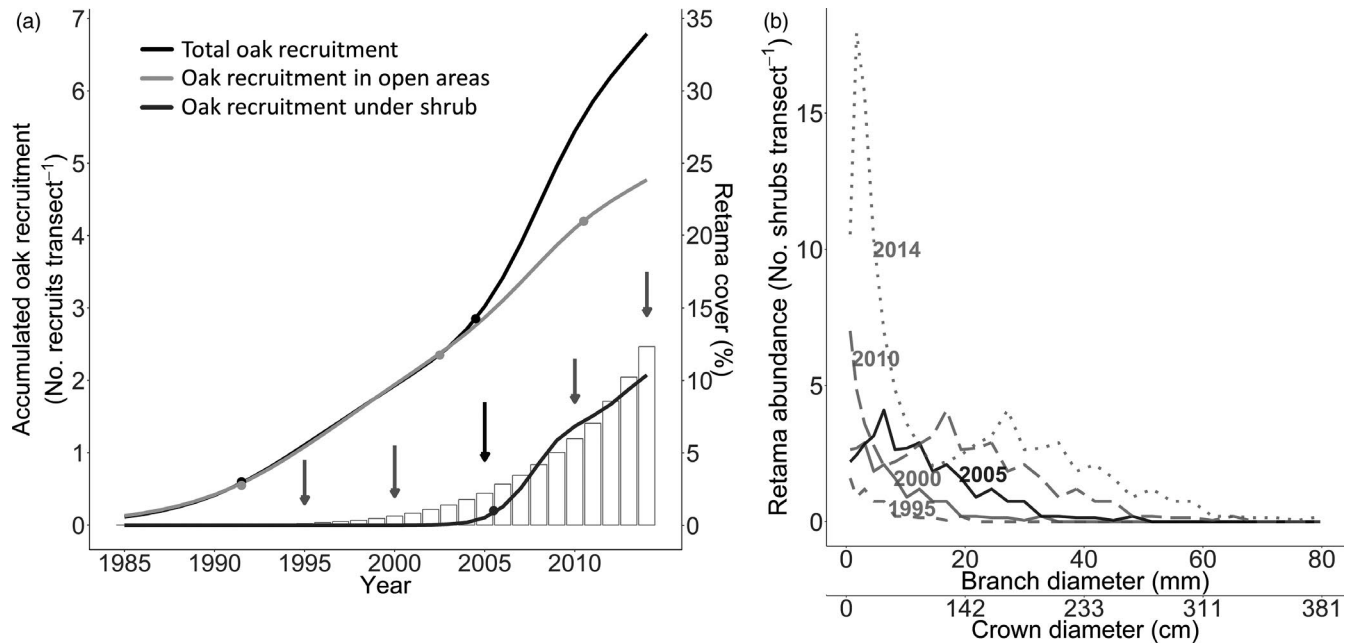
To assess if the effect of the shrub on oak recruitment varied across years as a function of climatic conditions, we estimated the intensity of shrub facilitation or competition on evergreen oak establishment with an interaction index. This index was estimated as the difference between oak recruitment under shrub and recruitment in open sites, in years and transects when at least one oak recruited (Gómez-Aparicio et al., 2004). We could not calculate the interaction index for the deciduous oak due to the lack of recruitment under shrub. We then run linear mixed models to ascertain the extent of the interaction index. We used quadratic and linear variants, as a function of each climatic variable (i.e. spring precipitation, summer precipitation or length of heat waves, all standardized). We used LMER function of LME4 package in R 3.4.3 to fit the models, with a Gaussian error distribution (Bates et al., 2017; R Core Team, 2017).

To account for the higher chances of recruiting under a shrub with increasing shrub cover, we included shrub abundance (i.e. the sum of the basal area of each shrub's widest branch,  $m^2/ha$ , standardized) as a covariate in the models. We also included transect as a random effect to account for fine grain environmental heterogeneity related to the clustered nature of our sampling.

### 3 | RESULTS

#### 3.1 | Dynamics of *Retama sphaerocarpa* and oak recruitment

All species showed two pulses of recruitment that varied in amplitude (i.e. maximum recruitment rate) and length (i.e. duration of the pulses; Figure 2, Table S4.1). Species recruitment pulses occurred at different years: in 2001 and 2013 for *R. sphaerocarpa*, 1994 and 2008



**FIGURE 3** (a) Accumulated recruitment over time for both oak species together (lines) and cover of the nurse shrub *Retama sphaerocarpa* (retama) each year (bars). The dots on lines represent change points in the temporal series of oak recruitment (i.e. acceleration in the recruitment rate when the curve has an increasing slope). The arrows represent the years selected to show the size structure of *R. sphaerocarpa* shrubland in panel b. The black arrow pointing to the black dot represents the change point when oak recruitment under shrub started. (b) Size structure of *R. sphaerocarpa* (retama) shrubland in the abandoned fields for the selected years

for the evergreen oak, and 1999–2000 and 2008 for the deciduous oak. The second pulse was higher than the first pulse for the three species (Figure 2). The amplitude and length of *R. sphaerocarpa* recruitment pulses were remarkably higher than those of oak species (Figure 2a). Pulses of deciduous oak recruitment were less ample and more episodic than pulses of the evergreen oak (Figure S4.1, violin plots in Figure 2b,c). Specifically, the deciduous oak recruitment was null in many years between 1985 and 1996 and it became more regular after 1997 but at a lower rate than the recruitment of the evergreen oak.

The observed recruitment of evergreen oaks under shrub started in 2005 (Figure 2b), 20 years after field abandonment. After that year, evergreen oak recruitment density was 1.5 times greater under shrub than in open areas. This recruitment under shrub represented 38% of total evergreen oak recruitment in the studied period (1985–2014). After the peak recruitment in 2008, recruitment rates remained constant under the shrub but considerably decreased in the open areas (Figure 2b). There were only three recruits of the deciduous oak under shrub (i.e. 9% of total deciduous oak recruitment), all of them in 2008.

### 3.2 | Oak colonization in relation to nurse shrub structure

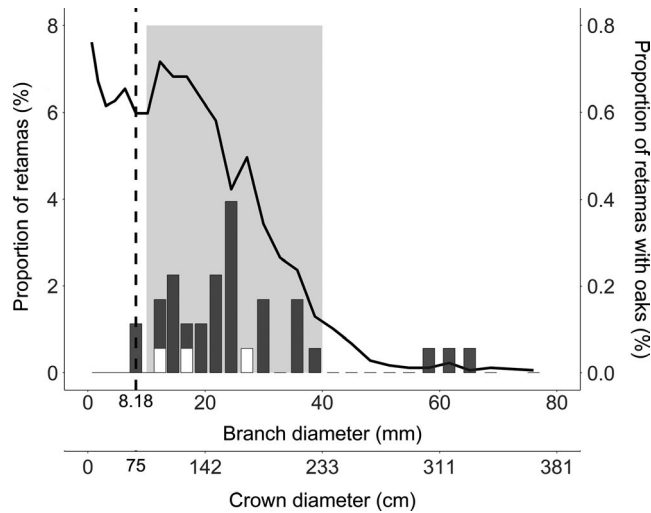
The change point analysis revealed that total oak recruitment accelerated twice during the study period, between 1991 and 1992 and between 2004 and 2005 (black line in Figure 3a, Table S4.2). While the first change coincided with an acceleration of oak recruitment in open areas, the second change coincided with the beginning and acceleration of oak recruitment under shrub (2005–2006; Figure 3a). By 2005, the

estimated *R. sphaerocarpa* shrubland density was  $276 \pm 246$  [0–1108] shrubs/ha (mean  $\pm$  SD [min–max]), and *R. sphaerocarpa* canopy cover was  $2.21 \pm 1.80\%$  [0%–7.16%]. Most of the shrubs were small (branch diameter < 10 mm) or medium (branch diameter 10–40 mm, Figure 3b).

*Retama sphaerocarpa* shrubs with recruited oaks underneath were larger (25.4 [8.2–65.1] mm of branch diameter and 168.6 [75.4–329.2] cm of crown diameter; mean [min–max]) than shrubs without oaks (14.5 [0.6–76.0] mm and 112.9 [12.4–367.4] cm respectively; Kruskal–Wallis  $\chi^2 = 25.4$ ;  $p < .001$ ). Oak recruitment only took place in a small fraction of the *R. sphaerocarpa* population (3.1% [0.5–9.1]), but this fraction was higher when only big (branch diameter > 40 mm) or medium *R. sphaerocarpa* shrubs were considered (15% [0–100] and 5.4% [0–16.7], respectively). Oak recruits were recorded only under *R. sphaerocarpa* shrubs that were at least 7–8 years old with branch and crown diameters > 8 mm and 74 cm respectively (dashed vertical line in Figure 4). Most of the *R. sphaerocarpa* shrubs that contained oaks under their canopy were medium-sized (85.7%; shaded area in Figure 4), which represented 57.9% of all the *R. sphaerocarpa* shrubs. In contrast, small and big shrubs contained only 5.7% and 8.6% of the oak recruits and represented 39.3% and 2.8% of the shrubs respectively.

### 3.3 | Shrub modulation of the climatic effects on oak recruitment

The results of the model of Equation (1) indicated that there were no statistically significant effects of the climatic variables on the recruitment rate of evergreen oaks in any of the two microsites (Figure 5a, model parameters in Table S4.3). Collinearity among

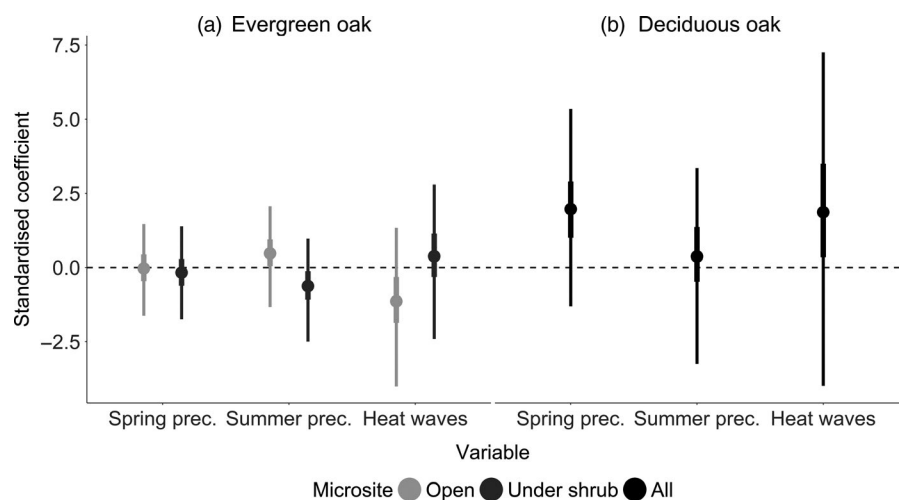


**FIGURE 4** Size class distribution of *Retama sphaerocarpa* (retama) population (line). The filled bar portion represents the fraction of *R. sphaerocarpa* individuals with *Quercus ilex* (evergreen oak) and the unfilled portion represents the fraction of individuals with *Quercus faginea* (deciduous oak) underneath. The dashed line represents the minimum *R. sphaerocarpa* size with an oak underneath. The shaded area represents the medium size individuals of *R. sphaerocarpa*

explanatory variables in Equation (1) was low ( $-0.27 \leq \rho \leq 0.34$ ;  $VIF \leq 1.20$ ; Dormann et al., 2013), and the fit of the model ( $R^2$ ) was .94 (Figure S1.3). There were not statistically significant effects of the climatic variables on recruitment rate of deciduous oaks either (Figure 5b, model parameters in Table S4.3).

The linear models for the interaction index between *R. sphaerocarpa* and evergreen oak recruitment had the best fit (AIC in Table S4.4). Spring precipitation did not have a significant effect on the interaction index (Table S4.5). However, the interaction index was inversely related to summer precipitation and directly related to the length of heat waves (Figure 6, Table S4.5). The effect of the nurse shrub become predominantly negative (i.e. recruitment under shrubs was lower than in open areas; Figure S4.2) when summer precipitation was  $> 47$  mm or the heat waves

**FIGURE 5** Standardized coefficients (parameter  $\times$  variable mean) of the climatic variables in the model of *Quercus ilex* (evergreen oak) recruitment (a; Equation (1) in Data analysis section) and in the model of *Quercus faginea* (deciduous oak) recruitment (b). Whiskers represent 95% of the credible interval of the estimated standardized coefficient, and bold whisker segments represent 75% of the credible interval. The microsite all (open+under shrub) applies to the deciduous oak



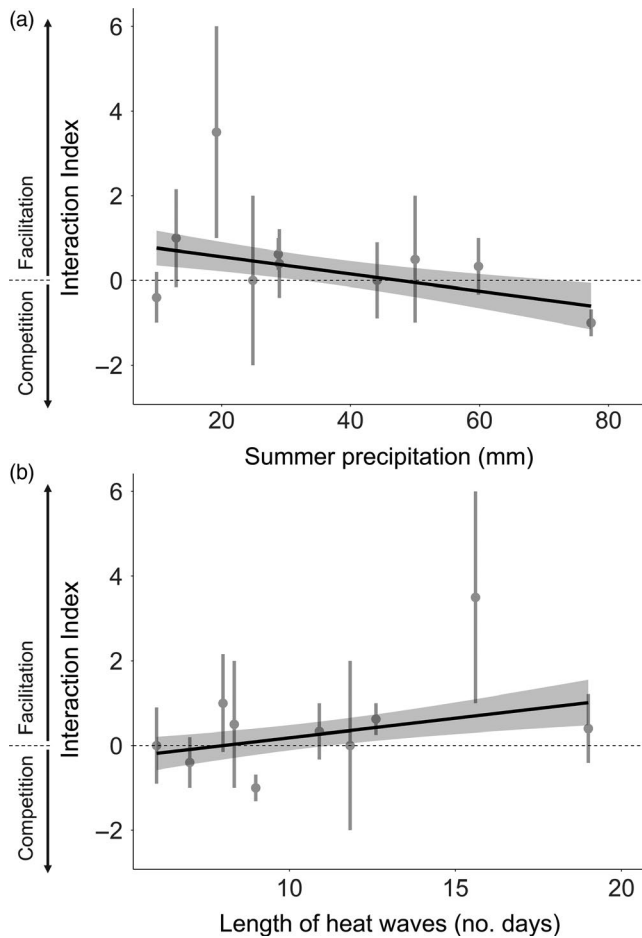
lasted  $< 8$  days (Figure 6). Precipitation  $> 47$  mm or heat waves shorter than 8 days occurred in 30% of the studied years each one (see climate from 2005 to 2014 in Figure 2d).

## 4 | DISCUSSION

Under stressful conditions, the facilitation of nurse shrubs is critical to ensure tree colonization, especially in areas that lack woody vegetation like abandoned fields. Despite their relevance in forest recovery, we know little about the dynamics of these effects. Our analysis of long-term colonization dynamics of oaks showed that their recruitment accelerated when oaks started recruiting under shrubs. It took 20 years of secondary succession to get to this point, likely when shrub cover and size were large enough to provide a facilitative effect. Our results also showed that evergreen oaks recruited more than deciduous oaks and, against our expectation, only the evergreen species benefited from recruiting under the nurse shrub. Facilitation was more pronounced in years with more arid summers, and the prevailing interaction between the nurse shrub and the evergreen oak switched to competition in wet and mild summers. Our study documented facilitation dynamics for oak recruitment at a single site, but these results could be likely general in the Mediterranean forest context. Still further studies at wider spatial scales with long temporal information would be necessary to corroborate how the nuances of this temporal perspective change.

### 4.1 | Long-term dynamics of forest expansion reveal different recruitment pulses of the nurse shrub and oak species

In accordance with our first hypothesis, the temporal trends in pulses of recruitment that we documented indicate distinct regeneration dynamics for the shrub and the two oak species. *Retama sphaerocarpa* —a pioneer shrub species— was the earliest colonizer, presumably due to its high dispersion capacity and drought tolerance (Haase, Pugnaire, Clark, & Incoll, 1999; Haase et al.,



**FIGURE 6** Fitted linear model of the relationship between the interaction index of *Retama sphaerocarpa* and *Quercus ilex* (evergreen oak) recruitment and (a) summer precipitation or (b) the length of heat waves. The dots are the mean interaction index in the 20 transects per precipitation and length of heat waves level and the whiskers represent the standard errors of the means. Positive values of the interaction index mean facilitation and negative values mean competition

1996; Padilla & Pugnaire, 2007). Since most juvenile mortality occurs during the first summer in Mediterranean forests (Mendoza, Zamora, et al., 2009; Rey Benayas, Navarro, Espigares, Nicolau, & Zavala, 2005; Rolo et al., 2013), the large pulse of *R. sphaerocarpa* recruitment towards the end of the studied period is likely due to the beginning of seed production by already established shrubs, as observed for other plant species (Harper, 1977). The oak recruitment pulses were probably caused by a combination of masting cycles (Koenig, Knops, Carmen, & Pearse, 2015; Pérez-Ramos et al., 2015), reduced competition with pre-existing plants (herbs in our study; Rey Benayas et al., 2015), and availability of nurse shrubs (Navarro-Cano et al., 2016; Perea et al., 2017). The increase in oak recruitment over time could also be attributed to the greater activity of acorn dispersers and to soil improvement after field abandonment (i.e. increase of C:N ratio, organic matter or aggregate stability; Hooker & Compton, 2003; Rey-Benayas, Galván, & Carrascal, 2010; Robledano-Aymerich et al., 2014). In

particular, the facilitative effects of the shrubs started 20 years after abandonment.

The two oaks also showed differences in their recruitment dynamics, in agreement with hypothesis 1. In consonance with other studies, the evergreen oak recruited more abundantly, and during longer time windows, than the deciduous oak (Gómez-Aparicio et al., 2004; Perea et al., 2017). Differential drought resistance between these species could be the main cause (Gil-Pelegri et al., 2017; Montserrat-Martí et al., 2009). Specifically, sensitivity of stomatal conductance to water stress is lower for the evergreen oak than for the deciduous oak (Acherar & Rambal, 1992; Mediavilla & Escudero, 2004). Moreover, acorn production and dispersion of the evergreen oak is usually higher than those observed for the deciduous oak (Del Arco, Beltrán, & Martínez-Ruiz, 2018; Pons & Pausas, 2007a, 2007b; Rodríguez-Estévez, García, Perea, Mata, & Gómez-Castro, 2007), and shoots of deciduous oaks are preferred by browsers over shoots of evergreen oaks (Espelta, Habrouk, & Retana, 2006).

Our results show that after 30 years of succession the evergreen oak (more stress-tolerant) had higher recruitment under the nurse shrub, while most deciduous oak juveniles (less stress-tolerant) recruited in open microsites (91% of recruited deciduous oak). This unexpected trend (i.e. the more stress-prone species benefits less from facilitation) has also been documented by Plieninger, Rolo, and Moreno (2010) and Torroba-Balmori et al. (2015). The difference between the recruitment microsites of evergreen and deciduous oak does not support our third hypothesis and contradicts a previous study that found similar occurrence of juveniles under shrub and in open areas for both evergreen and deciduous oaks (Perea et al., 2017), as well as the notion that facilitation benefits more the low stress-tolerant and herbivory-prone species (Costa et al., 2017; Madrigal-González, García-Rodríguez, & Zavala, 2014; Soliveres et al., 2014). The abundance and location patterns of evergreen oak recruitment could be due to the preference of the main dispersers, *Garrulus glandarius* (European jay) and *Apodemus sylvaticus* (wood mice), for the evergreen acorns (Del Arco et al., 2018; Pons & Pausas, 2007a, 2007b), and also to their preference for caching acorns under shrubs (Gomez, 2003; Gómez, Puerta-Piñero, & Schupp, 2008; Morán-López, Alonso, & Díaz, 2015). The lack of recruitment of deciduous oaks under shrubs could be also due to strong competition between the *R. sphaerocarpa* and the deciduous oak or to insufficient facilitation effect (Alday et al., 2016; Plieninger et al., 2010). Future experiments should further assess the underlying mechanisms of the recruitment differences between these two oak species.

For the evergreen oak, the recruitment under shrub was sustained over time in comparison to the pulse patterns observed in open areas. These dynamics may indicate that facilitation can increase the window of opportunity for evergreen oak colonization, which is probably due to both direct and indirect facilitation mediated by *R. sphaerocarpa* (Cuesta et al., 2010; Prieto et al., 2011). However, we could not quantify which proportion of the recruitment under shrub through time was due to the lower extension of open areas. The facilitation



of the evergreen oak together with the low recruitment of deciduous oaks observed suggest a more likely future dominance of evergreen oaks in Mediterranean abandoned fields (Perea et al., 2017).

## 4.2 | Relationship between retama population structure and oak facilitation

It is likely that direct and indirect facilitative mechanisms increase with nurse size and abundance (Allegrezza et al., 2016; Cuesta et al., 2010; Gómez et al., 2008; Navarro-Cano et al., 2016). Our results document the lagged facilitation effect associated with the shrub size and population structure (hypothesis 2). This hypothesis coincides with the results of Ramírez and Díaz (2008), that documented an acceleration of *Q. ilex* recruitment after 15 years of abandonment and its positive relationship with the cover of a leguminous shrub. The *R. sphaerocarpa* canopy cover was in average 2.21% at the beginning and acceleration of oak recruitment under shrub, which highlights the high facilitative capacity of this species.

*Retama sphaerocarpa* started effectively facilitating oak recruitment 20 years after shrubland recovery, and shrubs reached a threshold size of 74 cm in crown diameter. Recruitment rates under shrub were maximum at intermediate shrub sizes. The predominance of medium-sized individuals among the nurse shrubs suggests that facilitation under the shrub peaks and levels off once the shrub attains certain size (Navarro-Cano et al., 2016). At this point, the microclimatic conditions under the shrub canopy may be more moderate than outside, the soil may be specially enriched and structured or the shrub could attract dispersers for hiding and acorn caching (Gomez, 2003; Gómez et al., 2008; Pugnaire et al., 1996). Shrubs older than 21 years frequently had irregular and open canopies linked to branch senescence (V. Cruz-Alonso personal observation). This canopy opening could deteriorate the microclimate under the shrub and the physical protection against browsers (Schöb, Armas, Guler, Prieto, & Pugnaire, 2013), reducing the facilitation capacity of the older and/or larger shrubs. The *R. sphaerocarpa* size threshold for oak facilitation was reached when they were 7–8 years old, but the first oak recorded under shrub occurred 20 years after field abandonment. The lag between reaching a shrub structure able to facilitate oak recruitment and the effective facilitation suggests that recovery of other general ecosystem properties, further than the microenvironment created by the shrubs, such as soil structure and organic matter after agriculture ceased (Robledano-Aymerich et al., 2014), may also be needed for oak recruitment under nurse shrubs.

## 4.3 | Climate modulates the facilitative effect on oak recruitment

We could not find a clear support for the hypothesis that *R. sphaerocarpa* buffers the climate effect on oak recruitment since this effect did not differ between microsites. However, our analysis showed that the facilitative effect of *R. sphaerocarpa* on the evergreen oak was stronger (i.e. the interaction index was more positive) in years with more arid summers, whereas competition prevailed in wet and mild summers. These results support the Stress Gradient Hypothesis (Bertness &

Callaway, 1994; He, Bertness, & Altieri, 2013). The herb community that usually dries out in summer could grow longer under the *R. sphaerocarpa* shade in milder summers, and thus increase competition with oak seedlings (Tielborger & Kadmon, 2000). Surprisingly, we did not find any significant effects of spring precipitation or summer conditions on oak recruitment. The lack of a climatic effect could be due to limitations of our study, which might need additional data and not take into account other processes that might affect recruitment, such as mortality in later years (Debussche & Lepart, 1992). Moreover, modelling results may be affected by the non-inclusion of variables such as acorn production, abundance of dispersers and soil features which vary from year to year (Koenig et al., 2013; Rey-Benayas et al., 2010; Robledano-Aymerich et al., 2014).

## 5 | CONCLUSIONS

Long-term forest recovery in the studied Mediterranean abandoned fields exhibited differential pulses of woody species recruitment. Despite being closely related, oaks with contrasting leaf traits (evergreen vs. deciduous) and drought tolerance showed different patterns of recruitment. The *R. sphaerocarpa* shrubland buffered *Q. ilex* recruitment pulses, that is, recruitment rate under the nurse was maintained during periods of low recruitment in open microsites. The deciduous oak had lower recruitment than the evergreen oak during the first 30 years of forest succession and we could not document a positive effect of *R. sphaerocarpa* on recruitment of the deciduous oak. The nurse shrub population developed for 20 years before facilitating oak establishment and oak recruits appeared preferably in medium and big shrubs, always older than 7 years. The facilitative effect of *R. sphaerocarpa* increased with summer aridity. These results suggest a potential dominance of evergreen oaks in restored secondary forests in the mid-term under more arid climate scenarios and show that less stress tolerant species do not necessarily benefit more from facilitative interactions.

## ACKNOWLEDGEMENTS

This study was supported by the Madrid Government projects REMEDINAL-3 S2013/MAE-2719 and REMEDINAL TE-CM S2018/EMT-4338, and a contract between the FIRE (Fundación Internacional para la Restauración de Ecosistemas; <https://www.fundacionfire.org/>) and the University of Alcalá (FIRE-UAH 127/2017). VCA was supported by the Environmental Fellowship Programme of 'Tatiana Pérez de Guzmán el Bueno' Foundation (2015; <http://fundaciontatianapgb.org/>) and the REMEDINAL Postdoctoral Fellowship Programme (TE-CM S2018/EMT-4338). PRB was supported by the TALENTO Fellow Programme (Madrid Government, 2016-T2/AMB-1665). We thank the Spanish Ministry of Defence, especially to Inés Cava for assisting in access to field sites. We thank Daniel Gómez, Adrián Eceolaza, Fernando Viñeola, Pablo Quiles, Paloma Díaz and Julen Astigarraga their help with

field work, sample processing and plant dating; Asun Rodríguez for the help during the early stages of the work; Guillermo Bodega for kindly sharing his knowledge and lab equipment for histology processing; M. Luisa Aranda for facilitating the use of her laboratory microscope; Claudia and Cristina Miguel from the Cell Cultures Centre of the University of Alcalá, and Manuel Báez for their collaboration during the sample processing. Finally, we are very grateful to Jaime Madrigal, Enrique Andivia, Patricia González and Laura Fernández for their disinterested availability to solve any doubt about sample processing and dendrochronology. We acknowledge the input from three anonymous reviewers that improved the final version of this manuscript.

## AUTHORS' CONTRIBUTION

V.C.-A., P.V.-S., P.R.-B. and J.M.R.-B. conceived the ideas and designed the methodology; V.C.-A. and P.V.-S. collected the data; V.C.-A., II and P.R.-B. analysed the data; V.C.-A. and P.V.-S. led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

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## DATA AVAILABILITY STATEMENT

Data available via Figshare: <https://doi.org/10.6084/m9.figshare.9744953.v1> (Cruz-Alonso, Villar-Salvador, Ruiz-Benito, Ibañez, & Rey Benayas, 2019).

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## SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

**How to cite this article:** Cruz-Alonso V, Villar-Salvador P, Ruiz-Benito P, Ibañez I, Rey-Benayas JM. Long-term dynamics of shrub facilitation shape the mixing of evergreen and deciduous oaks in Mediterranean abandoned fields. *J Ecol.* 2020;108:1125–1137. <https://doi.org/10.1111/1365-2745.13309>