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> DOCTORAL DISSERTATION/ TESIS DOCTORAL

# EVOLUTIONARY ECOLOGY OF REPRODUCTION IN TWO MEDITERRANEAN PINE SPECIES: *PINUS PINASTER* AIT. AND *PINUS HALEPENSIS* MILL.

# ECOLOGÍA EVOLUTIVA DE LA REPRODUCCIÓN EN DOS PINOS MEDITERRÁNEOS: PINUS PINASTER AIT. Y PINUS HALEPENSIS MILL.

Presentada por D. Luis Santos del Blanco para optar al grado de Doctor por la Universidad de Valladolid

> Dirigida por: Dr. José M. Climent Maldonado

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### **SUMMARY**

#### -introduction-

The study of life-history traits is central to evolutionary biology and ecology, as lifehistory traits are closely linked to the environment where organisms thrive. Among them, traits related to reproduction such as the threshold size for reproduction, fecundity and the schedule of reproductive investment along life are particularly relevant. Their study helps to understand past adaptive processes as well as to infer future ones. For forest trees, this knowledge is particularly valuable and urgent, given their founding role in ecosystems all over the world.

Life-history traits can be addressed at the species level but usually, intraspecific withinand among-population variation also exists. This is the case for numerous plant species, as flowering time is commonly very variable and found to be correlated with natural selective pressures. This correlation may be due to plastic or genetic causes. A requirement for genetic responses to take place is the existence of additive genetic variation, which can be revealed by quantitative genetic studies or artificial selection experiments. Moreover, according to life-history theory, the expression of particular fitness-related traits is limited by costs or trade-offs with other traits also related to fitness.

*Pinus pinaster* Ait. and *P. halepensis* Mill. are two Mediterranean pine species showing numerous advantages for the study of tree reproductive traits from an evolutionary-ecological standpoint. Using these species as a model, it is possible to integrate aspects such as population differentiation, past and future local adaptation, plasticity, genetic architecture and multi-trait adaptive relationships.

#### -aims-

The aims pursued in the present study were, first, to document population differentiation in reproductive life-history traits. Second, to test the hypothesis that life-history traits are related to the environment by means of phenotypic plasticity and genetic change processes. Third, to estimate reproductive quantitative genetic parameters of key life-history traits. Fourth, to test the hypothesis that investment in reproduction is bound to costs or trade-offs with other adaptive traits, both at phenotypic and genetic levels.

#### -material and methods-

*P. pinaster* and *P. halepensis* are naturally distributed along extensive areas of the Mediterranean basin. *P. pinaster* has a particularly broad ecological niche, and there is wide genetic differentiation among populations, meanwhile *P. halepensis*, very tolerant to drought, shows smaller population differentiation, at least in neutral molecular markers. Data used in the present study were obtained from common garden experiments of three kinds: provenance, provenance-progeny and progeny common gardens, containing genetic material originated in natural populations or derived from breeding programs. Common garden data allowed to estimate to which extent phenotypic expression is genetically based and, when experiments were replicated under contrasting environmental conditions, to know whether this expression was also plastic. Furthermore, climatic data describing the areas where natural populations originated were obtained. Climatic data were then used to check their relationship with phenotypic traits. For *P. pinaster*, this relationship was checked after correcting for strong neutral genetic structure with nuclear microsatellite data.

Traits measured in the common gardens were related either to vegetative growth (total height, diameter at breast height, stem straightness) or with reproduction (presence/absence and quantification of reproduction, both male and female). For *P. halepensis*, reproduction was also manipulated experimentally seeking for the expression of somatic costs of reproduction. Data analysis was accomplished by linear (Gaussian data) and generalised (non-Gaussian data) mixed model analysis.

#### -results and discussion-

In both species, large population differentiation for reproductive life-history traits was a general finding. Moreover, mean population values –most notably regarding female function- covaried according to the environment where populations had evolved. A precocious and intense investment in reproduction was related to unfavorable environments for vegetative growth (low site index). This correlation is in agreement with the adaptive value of reproductive traits in forest trees; just as well as it has been described in other clinal traits such as timing of grow cessation. However, clinal trends were not found for the threshold size for male reproduction in *P. pinaster*. This could be related to sex allocation patterns strongly linked to tree height, as height is related to pollen dispersal conditions.

In *P. halepensis*, environmental effects (plasticity) on the expression of reproductive traits were also observed. In pararell to patterns of population genetic differentiation, a more growth-limiting environment was related to advanced ontogenetic development. In both species, it was noteworthy the absence of genotype by environment interaction for reproductive traits, contrasting with results for vegetative growth.

Plastic adaptive responses triggered in response to environmental change can reduce the need for genetic changes. On the other hand, high genetic variance and moderate to high heritabilities of reproductive life-history traits enable steady local adaptation processes. This possibility was backed by a neat response to artificial selection in *P. pinaster*.

At the somatic level, female costs of reproduction were confirmed in *P. halepensis* by means of a manipulation experiment, but not by correlational studies. In this species, investment in reproduction had a greater effect on future reproduction than on vegetative growth. At the genetic level, costs of reproduction in terms of growth showed a variable pattern depending on the species and the common garden environment. Nonetheless, several reproductive traits displayed a strong correlated response to selection for growth in *P. pinaster* i.e. evidence for the existence of a genetic trade-off between reproduction and growth. From the point of view of forest management, it is advised against the use of size as a single surrogate of fitness. Even though it is true that large trees tend to produce a greater number of offspring, this relationship is not tight and might be shadowed by trade-offs. On the contrary, under a new paradigm of adaptive forest management, the inclusion of reproduction to better define fitness is recommended.

**Keywords:** forest genetics, genetic variation, reproductive life-history traits, allometry of reproduction, costs of reproduction, local adaptation, fitness, evolution

### RESUMEN

#### -introducción-

El estudio de los caracteres de historia vital es central en biología evolutiva y ecología, pues están íntimamente relacionados con el entorno vital de los organismos. Entre los caracteres de historia vital destacan los reproductivos, tales como el tamaño umbral de reproducción, la fecundidad y el reparto del esfuerzo reproductor a lo largo de la vida. Su estudio ayuda a entender procesos adaptativos pasados y a inferir los futuros. En el caso de los árboles forestales, este conocimiento es particularmente valioso y urgente dado el papel básico de los árboles forestales en ecosistemas de todo el mundo.

Los caracteres de historia vital pueden ser estudiados a nivel de especie, pero comúnmente también existe variación intraespecífica entre y dentro de poblaciones. En el caso de numerosas especies vegetales es común observar cómo el momento de floración es variable y está relacionado con presiones selectivas naturales. Esta relación puede deberse a causas plásticas o genéticas. Un requisito para que se den respuestas a nivel genético es la existencia de variación genética aditiva. Ésta puede ser revelada mediante estudios de genética cuantitativa o estudios de selección artificial. Además, según la teoría de historia vital, la expresión de caracteres relacionados con la aptitud biológica (*fitness*) está limitada por la existencia de costes o compensaciones entre funciones.

*Pinus pinaster* Ait. y *P. halepensis* Mill. son dos especies de pinos mediterráneos que presentan numerosas ventajas para el estudio de caracteres reproductivos en árboles desde un enfoque ecológico-evolutivo. Usando estas especies como modelo es posible integrar aspectos como la diferenciación entre poblaciones, adaptación local pasada y futura, plasticidad, arquitectura genética y relación con otros caracteres adaptativos.

#### -objetivos-

Los objetivos del presente estudio fueron, en primer lugar, documentar la diferenciación entre poblaciones para caracteres de historia vital reproductivos. Segundo, evaluar la hipótesis de que los caracteres de historia vital están relacionados con el ambiente mediante procesos de plasticidad fenotípica y de cambio genético. Tercero, estimar parámetros de genética cuantitativa para caracteres reproductivos de historia vital. Cuarto, comprobar la hipótesis de que la inversión en reproducción conlleva costes o compensaciones en otros caracteres adaptativos, tanto a nivel fenotípico como genético.

#### -material y métodos-

Las especies objeto de estudio se distribuyen de forma natural en la cuenca mediterránea, ocupando grandes extensiones. *P. pinaster* tiene un nicho ecológico particularmente amplio existiendo una notable diferenciación genética entre poblaciones, mientras que *P. halepensis* es muy tolerante a la sequía y presenta una menor diferenciación entre poblaciones, al menos en caracteres moleculares neutros. Los datos utilizados en este estudio provienen de ensayos de ambiente común (*common gardens*) de tres tipos: ensayos de procedencias, ensayos de progenies y ensayos de procedencias-progenies, que contienen material genético procedente de poblaciones naturales y de programas de mejora. Esto permite estimar hasta qué punto la expresión fenotípica de los caracteres tiene una base genética, y cuando los ensayos están replicados en ambientes contrastados, saber si esa expresión puede ser plástica. Además se recavó información sobre parámetros climáticos de las zonas de origen de las poblaciones naturales para conocer su relación con caracteres fenotípicos. En el caso de *P. pinaster*, comprobó esta relación tras controlar mediante datos de marcadores microsatélite las posibles interferencias debidas a la existencia de una fuerte estructura genética en la especie.

Los caracteres obtenidos en los ensayos de ambiente común estuvieron relacionados con el crecimiento vegetativo (altura total, diámetro normal, rectitud) y con la reproducción (presencia o ausencia y cuantificación de la reproducción masculina y femenina). Para *P. halepensis*, también se llevó a cabo un experimento de eliminación de conos femeninos para revelar la existencia de costes somáticos de reproducción. El análisis de datos se realizó aplicando modelos mixtos lineales (variables gaussianas) y generalizados (variables no gaussianas).

#### -resultados y discusión-

En ambas especies, la existencia de diferenciación entre poblaciones para caracteres de historia vital reproductivos fue un hallazgo generalizado. Además los valores medios por población –particularmente en lo relativo a la función femenina- covariaron con el ambiente originario de las poblaciones, de forma que una inversión temprana e intensa en reproducción se asoció con valores ambientales poco favorables para el crecimiento (baja calidad de estación). Esta correlación es coherente con la importante función adaptativa de los caracteres reproductivos en árboles forestales, tal como se ha descrito en otros caracteres con variación clinal como puede ser la fecha de cese de crecimiento. Sin embargo, en *P. pinaster* no se hayaron tendencias clinales de variación en el tamaño

umbral de reproducción masculino. Ello puede estar relacionado con patrones de asignación sexual, fuertemente dependientes de la altura del árbol, ya que ésta determina las condiciones de dispersión de polen.

La influencia del ambiente externo (plasticidad) en la expresión de caracteres reproductivos fue constatada en *P. halepensis*. De forma paralela a los patrones observados en diferenciación genética, un ambiente más limitante para el crecimiento se relacionó con una aceleración en el desarrollo ontogénico. En ambas especies es destacable la ausencia de interacción genotipo x ambiente para caracteres reproductivos, contrastando con los resultados en crecimiento vegetativo.

Las respuestas adaptativas plásticas frente a cambios ambientales pueden atenuar la necesidad de un cambio genético. Por otro lado, la existencia de valores altos de varianza genética y heredabilidades de moderadas a altas para caracteres de historia vital reproductivos facilitan los procesos continuos de adaptación local. Esta posibilidad está respaldada por la existencia de una marcada respuesta a la selección artificial en *P. pinaster*.

A nivel somático, se constató la existencia de costes de reproducción femenina en *P. halepensis* mediante un experimento de manipulación, pero no mediante estudios correlacionales. En esta especie, la inversión en reproducción tuvo un mayor efecto en la reproducción futura que en el crecimiento. A nivel genético, los costes de reproducción en términos de crecimiento presentaron patrones variables, en función de la especie y el ambiente considerados. Sin embargo, varios caracteres reproductivos mostraron una fuerte respuesta correlacionada a la selección para crecimiento en *P. pinaster*, lo que constituye evidencia de la existencia de compensaciones a nivel genético entre reproducción y crecimiento. Desde el punto de vista del manejo forestal, se desaconseja utilizar el tamaño como único indicador de aptitud biológica. Aunque es cierto que los árboles grandes tienden a producir un mayor número de descendientes, esta relación es difusa y en gran medida puede estar influida por compensaciones (*trade-offs*). Al contrario, bajo un nuevo paradigma de gestión forestal adaptativa frente al cambio global, se propone incluir los caracteres reproductivos a la hora de definir de forma más precisa la aptitud biológica.

**Palabras clave:** genética forestal, diversidad genética, caracteres de historia vital reproductivos, alometría de la reproducción, costes de reproducción, adaptación local, aptitud biológica, evolución

# **INTRODUCTION**

## 1. INTRODUCTION

The concept of biological adaptation bears different meanings depending on the time scale considered. In forestry or agriculture, a certain genetic material is said to be well adapted, or to adapt itself well to particular environmental conditions if plants thrive well in terms of vegetative growth and survival. That is, if the combination of genotype and environment produces a phenotype that is considered suitable. In forestry, but also in everyday life, the term adaptation is thus understood as a plastic process along the plant's life.

In evolutionary terms, adaptation refers to a genetic process along generations. By adaptive processes, genetically superior individuals contribute disproportionally to the following generation, promoting genetic changes along generations (Darwin, 1859). Under the evolutionary concept of adaptation, a genetic material is considered to be adapted to a particular environment if outcompetes other genotypes. Competition here is not referred to vegetative growth, but to biological fitness i.e. number of gene copies in the next generation. The concept of biological fitness is thus central to understanding adaptation in evolutionary terms, but it is not needed, and often not used, to understand "static" adaptation.

#### 1.1 LIFE-HISTORY TRAITS

Life-history traits define developmental patterns in terms of growth, reproduction and survival in all kind of organisms (Roff, 1992a). The most important life-history traits are size at reproduction, lifetime offspring number, distribution of reproductive effort along time and the interaction between reproduction and adult mortality (Stearns, 1976). There is abundant evidence for numerous organisms on the close relationship between environmental conditions, either natural or induced by man, and life-history reproductive traits (Grime, 1977; Roff, 1992a; Stearns, 1992). Stable environments generally favour selection on large size at maturity, iteroparity (i.e. repeated reproductive events along life), low reproductive effort and high investment in maintenance. Conversely, fluctuating environments promote the opposite constellation of traits: low sizes at maturity, semelparity (i.e. a single reproductive event before death), high reproductive effort and low investment in growth and defence (Stearns, 1976; Grime, 1977).

Reproductive life-history traits provide a fecund and comprehensive research field in genetic, evolutionary and ecology studies and help understand the interactions between constantly adapting populations and their environment. Knowledge on reproductive life-history traits is particularly scarce in forest tree species, but this knowledge is relevant and urgent as they are founding species in their ecosystems, occupying great areas on the surface of the Earth (Petit & Hampe, 2006; Neale & Kremer, 2011).

#### 1.1.1 Life-history traits and local adaptation in plant populations

In the Plant kingdom, forest trees and annual plants illustrate well two extreme life-history strategies. Forest trees share a common pattern consisting of high juvenile mortality, delayed reproduction, large vegetative sizes and extended lifespan. And within woody species, angiosperms tend to have shorter times to reproduction than gymnosperms (Verdú, 2002). Yet, within this general pattern a wide range of variation exists, closely linked to environmental conditions (Thomas, 2011). Studies on within-species variability of life-history traits are especially valuable to illustrate local adaptation, as confounding phylogenetic factors can be minimised.

One of the first studies on genetic differentiation for life-history traits was authored by EP Lacey (1988), reporting on clinal variation for reproductive timing in *Daucus carota*. Remarkably, the studied species was artificially introduced by the 17<sup>th</sup> century in North America, where the study was performed, so that the described pattern of variation likely appeared in a short period of time. This rapid adaptive genetic change in an introduced species further stresses the relevance of life-history evolution research. Several other studies on introduced species have added to Lacey's seminal paper, also with similar results, describing a link between early reproduction and northern latitudes or harsher environments (Kollmann & Bañuelos, 2004; Alexander *et al.*, 2009; Colautti *et al.*, 2010; Lachmuth *et al.*, 2011). Interestingly, this pattern is also common in native species (Van Dijk *et al.*, 1997; Wesselingh *et al.*, 1997; Rees *et al.*, 1999; Callahan & Pigliucci, 2002; Mendez & Karlsson, 2004; Kuss *et al.*, 2008; Kagaya *et al.*, 2009; Brys *et al.*, 2011; Guo *et al.*, 2012).

Numerous studies have not only described extensive ecotypic genetic variation in reproductive life-history traits -most notably size at first reproduction- but also confirmed the existence of trade-offs between reproduction and growth. In addition to differentiation

among populations, high within-population variability is a common finding in life-history studies (Wesselingh & de Jong, 1995; de Jong & Klinkhamer, 2005; Childs *et al.*, 2010). The reference list is however biased towards short-lived species, and studies on life-history traits in trees are few, commonly focused at inter-specific variation (Niklas & Enquist, 2003; Wright *et al.*, 2005) and usually confounding genetic and environmental effects (Dodd & Silvertown, 2000; Thomas, 2011).

#### **1.1.2** Sex allocation theory

Dioecious species or monoecious species with separate male and female reproductive structures, are convenient for addressing more specific questions regarding each separate sex and their interplay (Charnov, 1982; Hardy, 2002; West, 2009). A well-developed field of theory has led to testable hypothesis and, in plants, researchers have posed relevant questions about the occurrence of dioecy (Ashman, 2006), sex ratios, i.e. relative abundance of males to females (Barrett & Yakimowski, 2010) and sex allocation, i.e. the within-individual ratio of male to total reproduction (Goldman & Willson, 1986).

Studies on wind-pollinated plants in reproductive ecology are less frequent than for animal-pollinated plants (Barrett, 2002), but have led to important hypotheses about the relative importance of male to female reproduction depending on size, i.e. size-dependent sex allocation. Fitness gain curves relate investment in a fitness function (usually investment in male or female functions) with fitness returns (Klinkhamer *et al.*, 1997; Zhang, 2006). In wind-pollinated plants, larger individuals are predicted to have relatively higher male investment as a result of the shape of both male and female fitness gain curves (Burd & Allen, 1988). As pollen can travel long distances and it is dispersed more efficiently from taller plants, male fitness gain curves are thought not to saturate (de Jong & Klinkhamer, 1994). While seeds, even though they might be also wind-dispersed, tend to remain closer to the mother plant and when produced in increasingly high numbers compete for limiting space so that fitness returns per unit resource invested in female function tend to flatten (Burd & Allen, 1988; Klinkhamer *et al.*, 1997).

There is considerable overall support for this prediction (Klinkhamer *et al.*, 1997) although conifers with segregated functions seem to be a particular case (Fox, 1993). Many conifers bear female cones on the upper part of their crown as a mechanism to enhance

crossed pollination and to avoid selfing, and it is size and not height what has been related to a plastic response towards increased male allocation (Fox, 1993; Ne'eman *et al.*, 2011). Other aspects like the ecological correlates of sex allocation and gender variation (Case & Ashman, 2005; Delph & Wolf, 2005), within species genetic variation (Koelewijn & Hunscheid, 2000; Friedman & Barrett, 2011) or the existence of plastic responses triggered by cues other than size (Paquin & Aarssen, 2004; Friedman & Barrett, 2011) remain less explored.

#### **1.2 PLASTICITY IN REPRODUCTIVE LIFE-HISTORY TRAITS**

The extent to which reproductive life-history traits are influenced by environmental conditions (plasticity) has also been explored mainly in herbaceous plants (Sultan, 2000; Matesanz *et al.*, 2010). Plant reproduction has a marked allometric pattern, such that larger plant sizes are related to higher reproductive output (Niklas & Enquist, 2003). As vegetative growth is known to be highly plastic, plasticity of reproduction is typically influenced by environmentally-induced changes in vegetative size.

Even though significant, plasticity in reproduction after accounting for size effects has been considered to be minor (reviewed in Weiner *et al.*, 2009). This claim is likely to be influenced by studies on annual plants as they have simpler allocation patterns. Under predictable environments, the optimal allocation strategy for annual plants is to invest all available resources in reproduction just before death, and size usually predicts accurately reproductive allocation (Weiner *et al.*, 2009). Opposed to that, biannual and longer-lived species must also invest a certain amount of resources in maintenance and growth along at least two seasons (de Jong & Klinkhamer, 2005). Thus, reproductive allometry is likely to be determined also by the fitness returns of investment in maintenance and future growth (Zhang, 2006).

The existence of a threshold size for reproduction in many plant species is considered as evidence for such interplay between investment in several fitness-related traits, which may be environment-dependent (Kozłowski, 1992; Burd *et al.*, 2006). As a result, plasticity in this key life-history trait has been documented in several studies in short-lived plants, usually responding to resource availability and competition (de Jong *et al.*, 1998; Sultan, 2000; Bonser *et al.*, 2010; Nicholls, 2011; Anderson *et al.*, 2012). Following expectations derived from life-history theory, and backing the results found in populationdifferentiation studies, harsh environments are found to promote reproduction at low sizes, while competitive environments tend to reduce it. The extent to which these results similarly apply to both male and female sexual functions is currently not so well developed, although female function is considered to be more costly ant thus, perhaps, more dependent on the environment (Obeso, 2002; Case & Ashman, 2005).

## 1.3 ADDITIVE GENETIC VARIATION AND SHORT TERM GENETIC CHANGE IN REPRODUCTIVE LIFE-HISTORY TRAITS

Given the forecasted climate change scenarios and the sheer importance of life-history traits in adaptation, it is relevant to ask whether plants will be able to evolve at a pace fast enough as to track environmental changes. For a trait to be able to evolve in response to natural or artificial selection, first there must be an underlying heritable additive genetic basis upon which to select, and then the selective agent must be able to efficiently select genotypes based on phenotypes (Lynch & Walsh, 1998). Life-history traits were first thought to be genetically depleted, as they are closely related to fitness and are under strong natural selection (Merilä & Sheldon, 1999). It has only later been accepted that although heritabilities for fitness traits might be low, they do have a wide genetic basis (Ellegren & Sheldon, 2008). This apparent contradiction is clarified by the existence of high error variance affecting statistical inference of heritability (Houle, 1992) and multivariate genetic constraints (genetic trade-offs) (Walsh & Blows, 2009).

Rapid formation of adaptive clines in several introduced species suggests that lifehistory traits can evolve readily. This is even more remarkable if we consider that introduction into non-native ranges usually occurs through strong bottlenecks that narrow genetic diversity (Sakai *et al.*, 2001). There is also mounting evidence on the rapid evolution of flowering time as a consequence of climate change (Franks & Weis, 2008; Munguía-Rosas *et al.*, 2011), highlighting once again the adaptive relevance of reproductive traits. Quantitative genetic and artificial selection experiments have also documented high heritabilities and clear responses to selection under controlled conditions (Wesselingh & de Jong, 1995; Wesselingh & Klinkhamer, 1996; Matziris, 1998; Burgess *et al.*, 2007). Therefore it seems clear that plant populations can indeed cope to some extent with novel environmental conditions. But it remains still critical to document correlated responses to selection on life-history traits, most notably on forest trees. Long life cycle of forest trees makes unfeasible for most researchers to accomplish artificial selection experiments and gather results within the duration of a research project. Forest trees are thus largely missing from the literature of artificial selection experiments although this kind of experiments constitute a powerful tool to test ecological hypothesis (Conner, 2003; Kawecki *et al.*, 2012). Collaboration between researchers and forest tree breeders is then perhaps a promising avenue to study responses to selection in trees from an evolutionary perspective.

#### **1.3.1** Evolutionary quantitative genetics in forest trees

Statistical models commonly used to analyse data from quantitative genetic data rely on the assumption that the studied trait is expressed as the result of the additive effect of a large number of loci, and therefore follows a Gaussian distribution. Continuous traits like growth are typically normally-distributed but some key life-history traits have a discontinuous distribution. For example, reproduction itself can be considered a binary trait when estimating precocity or the threshold size for reproduction. But it can also be considered to follow a Poisson distribution when dealing with count data. Direct application of conventional quantitative genetic formulae for analysing non-Gaussian data violates model assumptions and might imply serious bias. Even though phenotypically expressed in a discontinuous fashion, several traits have been shown to be governed by an underlying additive genetic basis. It is only above a certain value of a non-observed trait, directly dependent on that additive genetic basis and known as liability, that the trait is expressed. That is why discontinuous traits are also known as threshold traits.

Generalised linear statistical models (GLZ) implicitly transform non-Gaussian data into continuous normally distributed variables by means of a link scale. GLZ are now widely used in biological sciences in the analysis on non-Gaussian data, but their use in evolutionary quantitative genetics is still scarce. A recent paper by Nakagawa and Schielzeth (2010) described statistical formulae that can be used to estimate repeatability in non-Gaussian data, of which heritability is a particular case (within-family repeatability). Even more recently, the issue of estimating quantitative genetic parameters of non-Gaussian traits by means of animal models has been addressed specifically (Holand *et al.*, 2013). When applied to non-Gaussian life-history traits, those methods can greatly help in advancing our knowledge on relevant basic and applied research questions. See appendix VI for a more comprehensive description and application of quantitative genetic parameter estimation applied to non-Gaussian data.

#### 1.4 COSTS OF REPRODUCTION

Costs or trade-offs are defined as compromises between fitness functions, and have been fundamental in the development of life-history theory (Stearns, 1992). Among all, the costs of reproduction have been described as the *most prominent life-history trade-off* (Stearns, 1989). Their study in plants has been pursued by numerous scholars (reviewed by Obeso 2002 and by Thomas, 2011 in tree species). Overall but not overwhelming evidence confirms the existence of such costs, mainly described in terms of reduced vegetative growth (Obeso, 2002). However, depending on the methods used to describe costs of reproduction, the derived conclusions can be quite different.

Four methods have been proposed for the study of costs of reproduction (Reznick, 1992): correlational studies, manipulation experiments, genetic correlations and artificial selection experiments. The two first methods describe somatic or physiological costs of reproduction while the two latter ones describe genetic costs, and are thus the only relevant for evolutionary studies. Reviewed results, nonetheless, do not seem to greatly differ between phenotypic and genetic studies, or between correlational and experimental ones (Obeso, 2002). A non-trivial amount of non-significant or negative results should still stimulate research in the field, controlling for potentially confounding or compensatory factors and exploring the influence of additional variables. For example, studies considering the influence of additional factors have found evidence for higher costs of reproduction in harsh environments (Hansen *et al.*, 2013), for female sexual function (Montesinos *et al.*, 2006) and at small sizes (Climent *et al.*, 2008).

## 1.5 MEDITERRANEAN PINES AS MODEL TREE SPECIES TO STUDY REPRODUCTIVE LIFE-HISTORY TRAITS

Mediterranean pines, and specially *Pinus halepensis* Mill. And *P. pinaster* Ait., provide an excellent model to study past and future local adaptation processes, both from a basic and applied point of view (Lev-Yadun & Sederoff, 2000). This is due to several life-cycle

characteristics (Ne'eman *et al.*, 2004, 2011; Tapias *et al.*, 2004): (1) short age at first reproduction that enables early measurement of reproductive traits; (2) high reproductive allocation that enhances costs of reproduction; (3) low inter-annual variability in reproduction, as opposed to masting species, which makes easier recording continuous data series with better statistical properties; (4) non-shedding of mature cones (strictly serotinous cones are only found in some *P. pinaster* populations) facilitates reconstruction of female reproduction records in past years; (5) segregation of male and female reproductive structures within a tree crown allows testing specific hypothesis about the role of each reproductive function; (6) large within-population range of sizes, adequate for testing size-dependent sex expression and sex allocation hypotheses; (7) as obligate seeder species, connections between tree parts and functions are more straightforward, increasing the adaptive value of sexual reproduction; (8) their crown architecture is relatively simple and (9) they have a relatively low life span compared to other forest species; this improves the correlation between lifetime reproductive investment and that estimated in a shorter period.

Also relevant for evolutionary ecology and sustainable forest management is the species' wide ecological niche, especially that of *P. pinaster* (Fady, 2012), that translates into a wide distribution range. This allows testing the existence of correlations between phenotypic traits and environmental conditions, indicative of local adaptation. Another important ecological factor common to both species is their relationship with disturbance regimes, namely forest fires, which act as a strong selective agent for traits like the threshold size for reproduction, bark thickness and serotiny (Tapias *et al.*, 2004; Keeley, 2012).

Finally, the vast areas occupied either by natural or planted stands along their distribution range, but notably in Spain, highlight their environmental relevance. Mediterranean pine forest stands, are suffering increasingly frequent and virulent forest fires (Pausas *et al.*, 2008), which has spurred research on their reproductive ecology, most notable in *P. halepensis* (Iraima & Espelta, 2004; Gonzalez-Ochoa *et al.*, 2004; Verkaik & Espelta, 2006; De Las Heras *et al.*, 2007; Moya *et al.*, 2007; Espelta *et al.*, 2008; Ortiz *et al.*, 2011). From a forest management point of view, these studies seek for silvicultural treatments and conditions that enhance precocious seed production. This way, assuming a high prevalence of serotinous cones in young trees (Goubitz *et al.*, 2004), a canopy seed bank is built as soon as possible, thus enabling natural regeneration after recurrent forest fires. The main conclusions drawn from these studies are that treatments that promote tree

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vegetative growth such as pruning or thinning (De Las Heras *et al.*, 2007; Moya *et al.*, 2007; Ruano *et al.*, 2013), fertilization (Ortiz *et al.*, 2011) or site quality (Moya *et al.*, 2007) also promote precocious reproduction. However, a higher individual reproduction in sparse stands might not be enough to achieve higher yields per hectare (Matyas & Varga, 2000; Deng *et al.*, 2012).

Despite the notable relevance of the mentioned studies for applied forest management, their discussion in the broader field of ecology is limited by two main issues. First, cone production is considered as an absolute number per tree o per surface unit, irrespective of tree size. Therefore, larger trees also commonly produce a higher number of cones, despite reproduction per unit biomass is larger in smaller trees (Ortiz *et al.*, 2011). Second, some studies directly compare natural regeneration in different populations (De Las Heras *et al.*, 2007; Moya *et al.*, 2007). This confounds genetic and environmental causes of variation when trying to explain the common link between reproduction and climatic variables. Instead, in order to test systematically whether there exists among-population genetic variation for phenotypic traits, an unbiased sample of genotypes from each study population has to be grown in environmental conditions as homogeneous as possible (White *et al.*, 2007; Salmela *et al.*, 2010).

Forest genetic trials, or common garden studies, are specifically designed to provide uniform environmental conditions under which several genetic entries can be compared. They can be classified into provenance, progeny and/or clonal trials, depending on the genetic material used to set up the common garden, and they can be carried out under greenhouse or field conditions. Progeny and clonal trials can also be used to provide estimates of quantitative genetic parameters describing the genetic architecture and genetic determination of studied phenotypic traits.

In order to be adaptive, a phenotypic trait must be expressed according to a genetic basis and must be related to fitness, such that the trait correlates with higher reproductive success (Barrett, 2010). In forest tree research, many technical handicaps hamper a direct measurement of fitness under controlled conditions such as those found in a common garden. Researchers have thus focused on traits putatively linked to fitness that are more amenable to measurement, like vegetative growth (Alía *et al.*, 1995; Chambel *et al.*, 2007), phenology (Codesido & Fernandez-Lopez, 2009), water use (Voltas *et al.*, 2008; Aranda *et al.*, 2010) and cold stress (Climent *et al.*, 2009) in order to illustrate patterns of local adaptation and population differentiation in several tree species. Life-history trait variation and evolution in Mediterranean pines has been recently studied (Grivet *et al.*, 2013) but

basic account of within-species variation in reproductive traits is still largely missing, despite their relevance for understanding adaptation (but see Tapias *et al.*, 2001, 2004; Climent *et al.*, 2008).

AIMS

# 2. AIMS

Throughout my work I aimed to test basic evolutionary and ecological hypothesis derived from life-history theory, applied to the case of forest Mediterranean species such as *Pinus pinaster* and *P. halepensis*. I also intended that the research questions posed here could be related to practical issues regarding forest management and conservation of genetic resources. Specifically, my aims were:

- Document population differentiation in reproductive life-history traits such as the threshold size for reproduction and reproductive allometry (I, II, III, VI, VII)
- Test the hypothesis that reproductive life-history traits, as closely linked to fitness, are related to and modulated by the environment by means of plasticity (III) and natural selection (II, III, VII)
- Estimate quantitative genetic parameters (additive genetic variance, heritability, Qst) of reproductive life-history traits (I, IV, VII)
- Test the hypothesis that fitness components are negatively correlated by means of trade-offs, specifically between reproductive and vegetative growth traits by means of phenotypic correlations (I, V, VII), manipulation experiments (V), genetic correlations (I, II, III, IV, V, VII) and selection experiments (IV)

# **MATERIAL AND METHODS**

## 3. MATERIAL AND METHODS

#### 3.1 STUDY SPECIES

*Pinus pinaster* Ait. (maritime pine) and *Pinus halepensis* Mill. (Aleppo pine) are two Mediterranean pine species belonging to section *Pinus*, subsection *Pinaster*. *P. pinaster* occurs in Southern France, Northern Africa and the Iberian Peninsula, where glacial refugia have been suggested (Bucci *et al.*, 2007; de Lucas *et al.*, 2009). It has a very broad ecological niche, spanning from Atlantic to continental and semi-arid climates, and occurs at elevations ranging from sea level to 2000 m a.s.l. in Morocco.

*P. halepensis* has a broader geographical distribution, from Syria and Turkey in the East to the Eastern Iberian Peninsula in the West. Despite its common name, Aleppo pine, populations in the Eastern-most range are scarce and isolated, being more frequent a closely related species, *P. brutia* Ten.. *P. halepensis* also occurs around the Mediterranean basin in Greece, the Balkans, Italy, France, Israel, Tunisia, Libya, Algeria and Morocco (Chambel *et al.*, 2013). Present-day distribution is thought to be derived from a range expansion from Eastern populations towards the West (Grivet *et al.*, 2009b). Its ecological niche, though also broad, is more restricted to drier and warmer areas and has a greater tolerance to drought stress than *P. pinaster*. Both species coexist in Eastern and Southern Iberia along a coastal cline as well as in Northern Morocco (Soto *et al.*, 2010).

In Spain, the two species are found in vast areas in both natural and planted stands, as they have been extensively used in afforestation (Alía *et al.*, 1996; Gil *et al.*, 1996). *P. pinaster* is harvested for timber production mostly in the Atlantic region, and used to be an important species for resin production in Portugal and Central Spain (Perez *et al.*, 2013), an economic activity that is regaining importance. Recently, *P. pinaster* forests have also received attention as important producers of edible wild mushrooms (Gassibe *et al.*, 2011). Apart from this economic interest, both species are widely used for ecological restoration after forest fires and for former agricultural land conversion into forests, although their suitability for this objective has been largely debated during the last decades (Bellot *et al.*, 2004; Maestre & Cortina, 2004).

With such a high relevance, a comprehensive series of common garden provenance and provenance-progeny experiments of both species was implemented in Spain as well as in other Mediterranean countries (Chambel *et al.*, 2013). Rather than breeding, the main purpose of those experiments was to determine among-population ecotypic variation and to estimate quantitative genetic parameters for adaptive traits. This information is key to guide the use of forest reproductive materials and to infer the possible sources of adaptation to climate change. When replicated at contrasted sites, these series of common garden experiments are valuable to estimate phenotypic plasticity and genotype x environment interaction.

#### 3.2 COMMON GARDEN EXPERIMENTS

Data presented in this study are derived from several common garden experiments installed in Spain and, in the most part, belonging to the Spanish National Forest Genetic Trial Network (www.genfored.es) (Figure 1). Plantation, maintenance and measurement of the common gardens has been mostly accomplished by the Centro de Investigación Forestal belonging to the Instituto Nacional de Investigación Agraria (INIA-CIFOR), in collaboration with the Spanish Government (former MMA, currently MMARM) as well as other regional institutions (CITA-Aragón, Junta de Castilla y León, Banc de Llavors). Collaboration from local forest services has been fundamental for the maintenance of such long-lasting experiments.

Except for trials F26MON (II), PREB (I) and PRIA (I), which are progeny tests from breeding programs, sampling was done randomly in wild populations. For each wild population, seeds were collected from about 30 trees at least 100 m apart from each other. Then, seeds were either bulked (provenance trials) or kept separate in families (progeny trials). Seedlings derived from those seeds were produced under identical conditions in a nursery and planted into the final common garden site after one or two years.

For *Pinus halepensis*, I used both provenance and provenance-progeny common garden experiments. The provenance common garden was installed in 1997, comprising 52 native range-wide provenances, originated in continental Spain, Balearic Islands (Spain), Italy, France, Tunisia and Greece. It consists of six replicates placed in contrasting environments. In the present study (III) I used data from only two of those sites (Valdeolmos, Madrid –P24OLM-, and Rincón de Ademuz, Valencia –P24ADE-), selected

according to data availability and contrasting environmental conditions. P24OLM study site has a deep alluvial soil and the area has warm summers and mild winters. By contrast, P24ADE has a rocky and shallow soil at a windy slope with colder winters. This combination of soil and climate factors makes P24ADE site much more limiting for vegetative growth than P24OLM. Further details can be accessed in (III).

The *P. halepensis* provenance-progeny common garden was installed in 1995, being replicated in two sites, both included in the present study (I) (Megeces, Valladolid – F24MEG- and Montañana, Zaragoza, F24MON-). They comprise open pollinated families from 28 populations spanning the species distribution range in continental Spain and Balearic Islands, as well as progeny from three planted stands from inland Spain. Both trial sites also show a sharp environmental contrast. F24MEG is placed on a calcareous soil on a gentle slope, while F24MON is placed on a fertile alluvial soil with comparatively lower summer drought stress. Further details can be accessed in (I). In 2010, a subset of 110 trees in P24MEG common garden was also used in a manipulative experiment aiming at detecting somatic costs of reproduction (IV). First and second year developing female cones were carefully removed from half of the trees, remaining the other half as a control group. Vegetative growth and female reproduction were recorded during two years after treatment, and values for trees subject to treatment and control trees were compared. See (IV) for further details.

For *Pinus pinaster*, I used one provenance-progeny and three progeny common garden experiments, two of them not included in Genfored, but carried out by the Lourizán Forest Research Centre (Xunta de Galicia) (I). The provenance-progeny trial was installed in 2005, being replicated at five different sites, only two of them used in the present study (I,II) (A Merca, Orense –F26MER- and Cavada, Asturias –F26CAV-). They comprise open-pollinated families from 23 native populations spanning most of the species distribution range, including Atlantic Iberian Peninsula, Atlantic France, Corsica, Mediterranean Spain and Morocco. The ecological conditions of F26MER trial site can be considered intermediate respect to the species ecological niche, as it is placed in inland north-western Spain at a transition zone between Atlantic and Mediterranean climates. Further details can be accessed in (II). Ecological conditions of F26CAV are typically Atlantic, with higher summer rainfall than F26MER (I).



Figure 1. Location of *Pinus pinaster* and *P. halepensis* common gardens used in the present study. Red icons, *P. pinaster*; blue icons, *P. halepensis*. Check common garden code for a more detailed description in the text.

The two *P. pinaster* progeny trials reported in (I) (PREB and PRIA) were installed in 2003 with 28 progenies from selected plus trees from the Atlantic coast of Galicia (NW Spain). The *P. pinaster* progeny trial (F26MON) reported in (IV) was installed in 2001 at a highly continental area in the northern Iberian Range (Central-North Spain) as part of a breeding program in this Region of Provenance. It comprises open-pollinated families from phenotypically selected trees in natural stands neighbouring the trial site. It also contains the bulked progeny of a group of randomly selected trees from the same stands. The trial site has limiting conditions for vegetative growth due to cold winters and short summers plus a shallow and unfertile soil. Further details can be accessed in (IV).

#### 3.3 MEASUREMENTS

All common gardens were measured periodically since their establishment so that wide data series were already available prior to this work. This was possible thanks to the above-mentioned cooperative work in the framework of Genfored. Recorded variables were related to vegetative growth and more recently to reproduction (Table 1). Total height was measured with a telescopic pole to the nearest cm and diameter at breast height with a calliper to the nearest mm. When possible, female cones were counted according to cohorts, attending to their size and colour. This was possible for most of the common gardens (F26MER, F26CAV, PREB, PRIA, F26MON, F24MEG, P24OLM, P24ADE), given the relatively small size of the trees. Female strobili are formed during spring and are red and small sized (approx. 1 cm long) (Figure 2). By the end of the first season their colour turns green and their size increases (2-3 cm long). During the second season they attain their final size (10-20 cm long P. pinaster, 10 cm P. halepensis) but remain green. Then they gradually change their colour into bright brown between the autumn of the second season and the beginning of the third season. From that moment, the colour of the zones of the cone most exposed to the sun starts fading, turning into pale grey (Gil et al., 2009) (Figure 2). In F24MON, trees were large and crowns were close to each other, making it difficult to discriminate between cohorts. Instead, total female cones per tree were estimated from counts during 15 seconds (Knops & Koenig, 2012). This time interval was chosen after several tentative counts by different observers, aiming at combining a reasonable correlation with full cone count and a fast speed needed for measuring hundreds of trees with affordable field work schedules.



Figure 2. *Pinus pinaster* (a-e) and *P. halepensis* (f-j) female cone developmental stages and male cone clusters. a,f, female strobili emerged in spring; b,g one-year old female conelets; c, h, two-year old female cones; d, i, serotinous cones; e, j, male cone clusters.

Along ontogeny, male reproduction follows female reproduction in *P. halepensis* (Ne'eman *et al.*, 2004) but not in *P. pinaster*, indicating a more prominent role of early male reproduction in this species. Semi-quantitative data on male reproduction was recorded in F26MER common garden. Male pollen cone clusters were counted in 15 seconds and a qualitative assessment was used to estimate their size and density. Then, quantitative count data were weighed according to qualitative estimates.

As derived variables, stem volume over bark was used as an indicator of available resources in (I). Total biomass was estimated from allometric formulae, using diameter at breast height data (Montero *et al.*, 2005). The threshold size for reproduction was derived from binary models combining vegetative growth data and male or female binary reproduction data (presence / absence) (Wesselingh *et al.*, 1997) (I, II, III, IV, VII). Correlations between several directly measured and derived traits, indicative of phenotypic or genetic trade-offs were performed at several levels (individual, family or population) (I, II, III, IV, VII).

Species	Trial	Appendix	Recorded trait
	F26MER	I, II, VI, VII	Total height, diameter at breast height, quantitative female reproduction by cohorts and semi-
P. pinaster	F26CAV	Ι	quantitative male reproduction
- · <i>P</i> ·······	PREB	Ι	Total height, basal diameter and quantitative female
	PRIA		reproduction by cohorts
	F26MON	IV	Total height, diameter at breast, qualitative stem form, quantitative female reproduction by cohorts and qualitative male reproduction
	F24MEG	I, V, VII	Total height, diameter at breast height, and female quantitative female by cohorts
P. halepensis	F24MON	I, VII	Total height, diameter at breast height, and total quantitative female reproduction
	P24OLM	III, VII	Total height, diameter at breast height, and quantitative female reproduction by cohorts
	P24ADE	III, VII	Total height, diameter at breast height, and quantitative female reproduction by cohorts

Table 1. Summary of reported *Pinus pinaster* and *P. halepensis* traits recorded in several common garden experiments (see corresponding appendix).

Environmental conditions (temperatures and rainfall) from the areas where sampled populations had evolved, as well as from the common garden sites, were obtained from the climatic models by Gonzalo-Jiménez (2010) for Iberian populations and by Hijmans *et al.* (2005) for non-Iberian populations (II, III). Two models were used because Gonzalo-Jiménez's model provides better estimates than others mostly due to the use of a denser data network, but it is only available for the Iberian Peninsula. Otherwise, both models are highly compatible. These data were used to test for the existence of ecotypic trends in phenotypic traits measured in the common gardens. Environmental conditions in the common garden were also used to compute ecological distances (Gower's distance) between the conditions of origin of the sampled populations and those at the common garden (II, III).

#### 3.4 MOLECULAR DATA

Molecular marker data was used in (II) in order to control for strong neutral population genetic structure in *P. pinaster*, when testing for local adaptation patterns. Twelve nuclear microsatellite markers were genotyped in an average of 16 trees per population (range 6 - 30). Genotypic data was used to compute neutral genetic structure with the software STRUCTURE v2.2 (Pritchard *et al.*, 2000) (Figure 3). More details can be accessed in (II).



Figure 3. Barplot of individual-tree assignment probability to each of the optimal K = 6 clusters representing five geographical genetic groups in *Pinus pinaster* as produced by STRUCTURE 2.2 software. Each tree is represented as a line segment and vertically partitioned into K-coloured components, representing the individual's estimated proportions of ancestry in the K clusters. Population abbreviations are given in appendix II.

#### 3.5 DATA ANALISYS

Gaussian variables were fitted with Linear Mixed Models (LMM) while Generalised Linear Mixed Models (GLMM) were used for non-Gaussian variables. Population and/or family within population were included as random factors and informed about population differentiation or additive genetic variance at the species level. In IV, a two-level fixed factor (selected and non-selected) was included in the models in order to test for selection effects. Adjusted population and family estimates were derived from those models and used in correlations with other traits and climatic variables.

Block structure from the experimental design was included in the models when analysing growth variables in order to minimise environmental noise. In some cases, block structure was not considered when estimating quantitative genetic parameters. This yields results closer to those expected in nature, which might be more meaningful for evolutionary studies (Hadfield *et al.*, 2010). Models fitted to reproductive trait data (threshold size for reproduction, cumulative reproductive investment and reproductive allometry) could include tree height, volume over bark or biomass as a covariate. Thus, results reflect variation in reproductive traits not merely due to size effects (Weiner *et al.*, 2009). Several quantitative genetic parameters were computed from variance components derived from mixed models in provenance-progenies or progenies trials, namely narrowsense heritability ( $h^2$ ), population differentiation ( $Q_{ST}$ ) and coefficient of additive genetic variance ( $CV_A$ ) (Falconer, 1989). Specific formulae to compute quantitative genetic parameters derived from Generalised Linear Mixed Models are described in IV and VI after (Nakagawa & Schielzeth, 2010).
RESULTS

# 4. **RESULTS**

## 4.1 POPULATION DIFFERENTIATION AND LOCAL ADAPTATION

Significant population differentiation for vegetative and reproductive traits both in *P. halepensis* and *P. pinaster* has been a common finding throughout this work (I, II, III, VI, VII), as indicated by significant population effects in statistical analyses (Table 2). In F26MER common garden, *P. pinaster* male and female sexual functions were analysed separately in the same individuals, allowing the study of early sex allocation patterns. Despite significant genetic differentiation for both sexual functions, male threshold size for reproduction was much less variable than female threshold size for reproduction, leading to variation in early sex allocation mainly due to variation in female function (Fig. 4).

Significant correlations between environmental variables and phenotypic traits, likely indicative of local adaptation were found for *P. pinaster* (II) and *P. halepensis* (III). In the case of *P. pinaster*, correlations with environmental variables were still significant after correcting for neutral genetic structure, a potentially important confounding factor. Equivalent molecular data for *P. halepensis* were not available and correlations were not corrected, but in this species neutral genetic structure is much weaker than in *P. pinaster*. For both species, results were remarkably similar, with variables representing enhanced and early reproduction being negatively correlated with environmental variables indicative of favourable growth conditions, mostly higher winter temperatures and lower summer temperatures. Surprisingly, rainfall regimes were little correlated with the variation patterns found.

	Trait	Trial	Species	Appendix
Vegetative	Height	F24MEG, F24MON, P24ADE, P24OLM F26MER	P. halepensis, P. pinaster	III, VII, II, VII
	Volume over bark	F24MEG, F24MON, PREB, PRIA	P. halepensis, P. pinaster	Ι
	Biomass	F24MEG, F24MON, P24ADE, P24OLM, F26MER	P. halepensis, P. pinaster	III, VII, VII
Reproductive	Female cumulative repr. investment	F24MEG, F24MON, P24ADE, P24OLM, F26MER	P. halepensis, P. pinaster	III, VII, VII
	Female repr. allocation	F24MEG, F24MON, P24ADE, P24AOLM PREB, PRIA	P. halepensis, P. pinaster	I, III
	Repr.allometry	P24ADE, P24OLM, F26MER	P. halepensis, P. pinaster	III, VII
	Female threshold size for reproduction	F24MEG, F24MON, P24ADE, P24OLM, F26MER, F26CAV	P. halepensis, P. pinaster	I, III, VII, VII
	Male threshold size for reproduction	F24MEG, F26MER	P. halepensis P. pinaster	II, VI, VII

Table 2. Vegetative and reproductive traits for which evidence of population differentiation was found in several *Pinus pinaster* and *P. halepensis* common gardens.



Figure 4. Median threshold size for first reproduction through male (white bars) and female (grey bars) functions in *Pinus pinaster* populations grown in F26MER common garden in north-west Spain. Bars represent the posterior mode of Bayesian estimates, with lower and upper 95% credible intervals. Population abbreviations are given in appendix II.

#### 4.2 PHENOTYPIC PLASTICITY

The study of phenotypic plasticity and genotype by environment patterns for reproductive traits was first addressed in (I) and then more explicitly in (III). The allometry of reproduction in *P. halepensis* responded plastically to environmental conditions, expressing an advanced ontogeny (enhanced reproduction) when grown under environmental stress (III) (Figure 5). Interestingly, cumulative reproductive investment was similar between two contrasted sites (III, Figure 5). A consistent finding across species and sites was the absence of relevant genotype by environment interaction for reproductive traits (threshold size for reproduction, reproductive allometry or cumulative reproductive investment) indicated either by high site-to-site correlations (I, III) or non-significant site by populations terms (III). This result contrasted with that found for vegetative traits (height, biomass, volume over bark) where site-to-site correlations were low and site by population terms were highly significant.



Figure 5. Reaction norms for vegetative and reproductive traits measured in two *Pinus halepensis* provenance common gardens subject to contrasting environmental conditions (P24OLM low and P24ADE high environmental stress). Lines connect mean values per source population at each common garden. Variables and units are as defined in appendix III.

# 4.3 ADDITIVE GENETIC VARIATION AND SHORT TERM GENETIC CHANGE IN REPRODUCTIVE LIFE-HISTORY TRAITS

Heritabilities for vegetative and reproductive traits are reported in (I, IV, VI and VII). Heritabilities for vegetative traits were lower than those for vegetative traits in both species, except for male function in IV, which was bound to cero. When comparing male and female function, no patterns emerged for range-wide populations in *P. pinaster*, with moderate to high heritabilities (0.27-0.73) (VI).

A single event of artificial selection was sufficient to produce genetic changes in all recorded traits at a young age, except diameter at breast height and biomass (IV). Meaningfully, direct response to selection in vegetative traits was much weaker than the indirect response for both male and female reproductive traits.

#### 4.4 COSTS OF REPRODUCTION

The study of costs of reproduction, either phenotypic or genetic, was present all along this work (I, II, III, IV, VII) and was specifically addressed in V. Altogether, all four approaches to the description of costs of reproduction were explored: phenotypic correlations (I, V), manipulation experiments (V), genetic correlations (I, V) and selection experiments (IV).

Phenotypic correlations between variables relating vegetative size and absolute female reproductive size were invariably positive, and ranged from a minimum of 0.12 in *Pinus pinaster* (I) to a maximum of 0.56 in *P. halepensis* (I), indicating that larger trees tended to produce more seed cones (Figure 6). When vegetative size was compared with reproductive allocation, correlations tended to be negative [-0.12 – -0.18 *P. pinaster* (I), -0.40 – 0.11 *P. halepensis* (I, V)], indicating that small trees tended to produce a proportionally higher number of seed cones. In *P. pinaster*, trees that started their reproductive phase producing only females cones were smaller than those that started producing only male cones, while cosexuals were the largest reproductive class (I, II, IV). In *P. halepensis*, there were virtually no trees producing only male cones (I).



Figure 6. Scatterplot of the number of developing female cones (emerged in the 14<sup>th</sup> and 15<sup>th</sup> year, respectively) versus tree size at 15 years, as estimated by the stem volume over bark, in several provenances of *Pinus halepensis* trees grown in F24MEG common garden. Blue dots, Cabanellas population; yellow dots, Santany population; pink dots, Cazorla population; green dots, Villavieja de Tordesillas population. r = 0.43; 95% confidence interval 0.39-0.47; p-value <0.001; n = 1385.

The results from the manipulation experiment (V) indicated that removal of female cones prior to their development resulted in a marginally significant (P < 0.07) increment of growth the same year after treatment, but the effect vanished one year after P = 0.77). In contrast, there was a more significant increase of female reproduction one year after cone removal (P < 0.01) (Figure 7).

Genetic correlations between vegetative size and absolute female reproductive size yielded contrasting results. Strong negative genetic correlations were found for *P. halepensis* in F24MON (I) and weak negative for *P. pinaster* in PREB and PRIA, opposing phenotypic correlations in those three cases and thus implying strong positive environmental correlations. Genetic correlations in F24MEG (I, V) were also lower than phenotypic correlations, but remained positive. Genetic correlations between vegetative size and reproductive allocation were all negative and commonly strong in both species (I, II). Among-population correlations, indicative of genetically-based trade-offs were also reported in II and III for *P. pinaster* and *P. halepensis*, respectively. A positive correlation between size and threshold size for female reproduction (i.e. delayed reproduction at bigger sizes) was found, although only under a stressful environment for *P. halepensis*.

Results from the selection experiment (IV) provided stronger evidence for genetically-based costs of reproduction in *P. pinaster*, as progeny from trees selected for increased growth also showed a reduced investment in both male and female reproduction.

This was indicated by increased threshold sizes and enhanced size-corrected reproduction for male and female reproductive functions (Figure 8).



Figure 7. Boxplots of a) basal area in 2010, one growing season after experimental manipulation, and b) number of female cones in 2011, one reproductive season after experimental manipulation comparing control *Pinus halepensis* individuals (CTR) with others subjected to experimental removal of developing female cones (FCR). Appendix V.



Figure 8. Comparison of height density distribution and threshold size (height) for first reproduction between *Pinus pinaster* progenies of trees either selected (S) or not selected (C, control) for timber production after one generation. F26MON common garden. Bell-shaped lines represent height probability distribution at age 10 yrs. Control group, orange lines; selected group, blue lines. Vertical lines represent threshold sizes for reproduction. Solid line, female function; broken line, male function. 95% Credible intervals for threshold sizes are represented by horizontal lines. Arrows show changes in threshold size for reproduction in male and female function due to selection. Control group, left; selected group, right. Appendix VII

DISCUSSION

# 5. **DISCUSSION**

Results presented throughout this study based on data from range-wide *P. pinaster* and *P. halepensis* common garden experiments have revealed significant population variation for several key reproductive traits. These findings, that build up on previous work in *P.halepensis* (Climent *et al.*, 2008) and *P. pinaster* (Tapias *et al.*, 2004), are among the first reports in the forestry literature on this topic. Results also highlight a close relationship between reproductive traits and provenance environmental variables i.e. those found where populations had evolved, as well as between reproductive and growth traits.

### 5.1 POPULATION DIFFERENTIATION AND LOCAL ADAPTATION

The existence of within-species population variation for adaptive traits is a wellestablished fact for species with extensive distribution ranges e.g. *P. contorta* (Rehfeldt *et al.*, 1999) and *P. sylvestris* (Rehfeldt *et al.*, 2002), or for those showing a wide neutral genetic differentiation between populations e.g. *P. pinaster* (Alía *et al.*, 1995). This is less acknowledged for species that do not meet those conditions, like *P. halepensis*. But even within the Iberian Peninsula, where neutral genetic diversity is lowest due to recent migration (Grivet *et al.*, 2009b), population differentiation is also present in this species. So, barring some exceptional cases like *P. pinea* (Mutke *et al.*, 2010), within-species variation is a frequent finding (Alberto *et al.*, 2013).

Local adaptation in plants is commonly tested by comparing fitness of local versus non-local populations grown in reciprocal transplant experiments (Leimu & Fischer, 2008). Unfortunately, in tree studies such comprehensive tests are often unaffordable and usually, many populations are tested in only one or few common gardens. Evidence of local adaptation can nonetheless be inferred from the relationship between phenotypic trait values measured at the common garden, and environmental descriptors of the areas where populations have evolved. Also, under the hypothesis of local adaptation, it is expected a positive correlation between fitness and a measure of how similar is the common garden environmental distance. In the present study, the aforementioned population variation found both for vegetative and reproductive traits in *P. pinaster* (I, II, VII) and *P.* 

*halepensis* (I, III, VII), did not occur haphazardly. Instead, mounting evidence suggests that indeed adaptive phenotypic variation is related to environmental conditions where populations have evolved (reviewed in Alberto *et al.*, 2013) suggesting that studied traits have adaptive value and that local adaptation processes have occurred and might still be occurring. Additionally, when phenotypic traits are measured directly in native populations rather than in common gardens, population differentiation tends to be much larger, given synergistic phenotypic and genetic variation i.e. co-gradient variation (Kremer *et al.*, 2013). Phenological traits related to cold damage like bud burst and growth cessation, are commonly used as examples of traits with adaptive clinal variation (Savolainen *et al.*, 2007; Neale & Kremer, 2011). The results presented in this study (II, III), add reproductive traits to that list.

Drought is thought to be an important limiting factor for Mediterranean species (Baquedano et al., 2008), and thus a correlation between rainfall and life-history traits was expected. Also, summer rainfall is correlated with area burnt in forest fires (Pausas, 2004; Pausas & Paula, 2012), and can potentially be linked to fire-related traits such as serotiny (Hernández-Serrano et al. in prep). However, it was the continentality index (annual temperature range) that lied among the climatic variables that best correlated with reproductive traits. Precocious populations tended to originate in areas with low winter and high summer temperatures, i.e. large continentality index, that define short growing seasons. Populations with delayed reproductive ontogeny tended to originate in areas with mild winter and summer temperatures, i.e. low continentality index that define long growing seasons. This pattern was stronger in *P. pinaster* (II) than in *P. halepensis* (III). These results largely agree with assumptions derived from life history theory, predicting a delayed reproduction (both in age and size terms) in those environments that favour vegetative growth (Roff, 1992b). The extent to which genetically-based patterns are enhanced or counteracted by phenotypic plasticity in natural P. pinaster or P. halepensis populations, resulting in co-gradient or counter-gradient variation remains unexplored. Results obtained in a replicated common garden experiment (III, see below), point towards co-gradient variation for genetic and environmental effects.

The study of population variation in early male reproductive function, addressed for *P. pinaster*, provided a different picture. Among-population differences in male threshold size for reproduction (TSR), despite significant, varied much less than those for female TSR, and they were not correlated with any environmental variable. A likely explanation for these results can be derived from size-dependent sex allocation theory applied to wind-

pollinated plants (Klinkhamer et al., 1997), and specifically to monoecious trees (Fox, 1993). It is believed that fitness gain curves in wind pollinated species will be saturating for female function, but not or to a much lesser extent for male function. Also, male function is likely to benefit more from indirect size effects (de Jong & Klinkhamer, 1994). This seems to be corroborated by parentage studies in wind pollinated trees, as the number of sires and male migrants contributing to the next generation is much higher than that of females (Bittencourt & Sebbenn, 2007; Grivet et al., 2009a; Robledo-Arnuncio, 2011). Such combination of male and female fitness gain curves theoretically leads to increased male sex allocation with size, a pattern confirmed in this study in P. pinaster (VI) and independently also in *P. halepensis* (Ne'eman *et al.*, 2011). Here, the role of height, although highly correlated with total size, seems to be indirect for the explanation of sizedependent sex allocation. This is because in conifer species like P. pinaster (II) and P. halepensis (Shmida et al., 2000) gender is segregated in tree crowns, with female cones placed on top of the tree. The existence of low variability in the male TSR can then be related to uniform conditions across populations for pollen dispersal, depending on tree height. This would set a relatively narrow threshold from which release of pollen is efficient, but above which no major increases in efficiency are gained (II). These results will also need to be contrasted with quantitative male reproduction at a more advanced age.

### 5.2 ADAPTIVE PROCESSES WITHIN POPULATIONS

The vision that reproductive traits are important targets of natural selection is widely supported in plants by increasing evidence (Munguía-Rosas *et al.*, 2011; Shaw & Etterson, 2012): first, genetically-based clinal variation in reproductive traits has been commonly reported for short lived plants, interestingly, also for species in introduced ranges (Colautti *et al.*, 2010; Lachmuth *et al.*, 2011). Second, variation in reproductive traits has been related to species persistence in native plant communities (Willis *et al.*, 2008; Dante *et al.*, 2013). And third, microevolutionary processes in annual species can produce significant changes in reproductive traits in only few generations (Franks *et al.*, 2007; Sultan *et al.*, 2013). The question seems solved for short lived plants species, but will forest trees be able to respond to selective pressures if reproductive traits are targeted? (Davis *et al.*, 2005)

Selection acts on phenotypes, which are the result of the interaction between particular genotypes and particular environments. Local-scale environmental influence varies on different traits, being the expression of vegetative growth commonly affected by small-scale spatial heterogeneity (Zas, 2006) as opposed to the allometry of reproduction (II, III). Similarly to phenotypic variation, genetic variance of plastic traits is also dependent on the environment (Sgrò & Hoffmann, 2004; Matesanz *et al.*, 2013). The present study suggests that the expression of growth traits under unfavourable conditions can be more constrained than that of reproductive traits, despite both exhibit plasticity (III). Also, significant genotype by environment interaction for growth traits but not for reproductive traits (I, III) may be related to the spatial scale at which the environment influences those traits (van Kleunen & Fischer, 2005; Rodríguez, 2012).

A factor known to lower the effectiveness of selective processes is the expression of adaptive phenotypic plasticity (static adaptation) (van Kleunen & Fischer, 2005). Reproductive allocation in short lived plants had been thought to be one of the less plastic traits (Weiner *et al.*, 2009). Nonetheless, it has been suggested that plasticity of reproductive allocation in long-lived sessile species like trees should be relevant (Burd *et al.*, 2006), but there is a scarcity of studies reporting on this subject. The case of adaptive plasticity of reproductive allocation in range-wide *P. halepensis* populations described in this study is therefore particularly important (III). Trees grown under a more stressful environment reached sexual maturity earlier, both in time and in size, than under a favourable environment. Notably, this response matches genetic responses to selection in climatic clines (II, VII).

Given that only genetic differences can promote evolutionary change, and considering single traits, a higher correlation between genotype and phenotype implies a higher efficiency in selective processes, as defined in the classical breeder's equation (Cornelius, 1994; Lefèvre *et al.*, 2013). In the present study, higher heritability values for reproductive versus growth traits (I, VI) suggest a more efficient selection for reproductive traits. A greater change due to artificial selection in reproductive traits compared to growth traits (IV) supports this idea. Interestingly, other phenotypic traits that also show clinal variation like bud burst and bud set also have high heritabilities (>0.5) (Neale & Kremer, 2011). Still, the only currency of natural selection is fitness, and only those phenotypic traits genetically correlated with fitness (either positively or negatively) will experience genetic changes after selection. This is known as the Robertson-Price identity (Robertson, 1966; Price, 1970), and currently there is a lack of studies applying it to trees. High within-

population genetic variation and its phenotypic expression, likely indicate that reproductive life-history traits will indeed respond to natural selective forces. This response might be delayed by the expression of adaptive phenotypic plasticity (Matesanz *et al.*, 2012; Anderson *et al.*, 2012), although considering large time scales, early reproduction has also been related to increased evolutionary rates (Verdú, 2002).

### 5.3 COSTS OF REPRODUCTION AND TRADE-OFFS

Considering the principles of allocation and life-history theory (Stearns, 1989), for a given amount of resources an increase in reproduction is typically related to a decrease in growth (Roff, 2000; Obeso, 2002; Thomas, 2011). At the individual level, trade-offs might not be evident given that not all trees have access to the same level of resources (Reznick *et al.*, 2000). Within a population, a tree with access to more resources e.g. deeper soil, can attain a larger size and at the same time produce a higher number of seeds than other i.e. a positive environmental correlation. This, in turn, creates a positive relationship between growth and reproduction for the whole population (de Jong & Klinkhamer, 2005). Forest management practices aimed at increasing cone production in a given forest stand (thinning, pruning, fertilization), exploit this phenotypic relationship between large size and large seed production (Gonzalez-Ochoa *et al.*, 2004; De Las Heras *et al.*, 2007; Ortiz *et al.*, 2011; Ruano *et al.*, 2013).

Potentially confounding factors can be controlled, for example by submitting to experimental manipulation a given group of trees and comparing the response to an equivalent control group (Reznick, 1992). The results of such an experiment in young *P*. *halepensis* trees provided some insight into the expression of somatic costs of reproduction in this species (III). Removal of developing female reproductive structures was related to a 70% increase in reproduction the following season, compared to the control group. Contrarily, the response in terms of vegetative growth was weak. This suggests that costs of reproduction in this species might be more relevant in terms of future reproduction, rather than just considering vegetative growth. A cost of current reproduction paid in future reproduction has indeed a deeper evolutionary significance (Reznick, 1985; Stearns, 1989) but evidence in plants is not overwhelming (Obeso, 2002) and has been rarely considered in tree species (Thomas, 2011).

Evidence of somatic costs of reproduction is also inferred from differences in average size for trees starting their reproductive phase as males, females or cosexuals (I, II and VI). Early females were smaller than early males and cosexuals, suggesting a higher cost of reproduction for female function (Montesinos *et al.*, 2006; Verdú *et al.*, 2007). Nonetheless, the pattern is somewhat blurred given that cosexuals, bearing both female and male reproductive structures, are among the largest individuals. It is therefore difficult to separate to what extent reproduction is dragging growth, and in turn, to what extent vigour can help overcome those costs of reproduction. Perhaps, a more direct approach of female versus male costs of reproduction could be achieved in dioecious species (Montesinos *et al.*, 2012), which in conifers tend to be dispersed by animals (Givnish, 1980). Also, in mature cosexual pine trees investment in male function is typically very high, and likely to be costly. The hypothesis that male sexual function in conifers has been enhanced by sexual selection remains unexplored (Moore & Pannell, 2011; Pannell & Labouche, 2013).

Phenotypic correlations between size and reproduction where invariably positive for all datasets analysed in this study. In fact, positive phenotypic vegetative-reproductive size correlations, are the base for general allometric models (Niklas & Enquist, 2003), and have been rightly considered as proof for the adaptive value of size. A few studies on parentage analysis have also been able to find a positive correlation between tree size and the number of seedlings established in a forest stand (González-Martínez et al., 2006; Moran & Clark, 2012). But this does not necessarily mean that size is an accurate predictor of reproduction. Taking the relationship one step further, and considering it strong and invariable, growth traits have been repeatedly used as surrogates of fitness in forest trees (Wu & Ying, 2004; Savolainen et al., 2007; Oddou-Muratorio & Klein, 2008; Ramírez-Valiente et al., 2010). The allometry of reproduction has been studied most extensively in short lived plants, where statistical models with plant size as an explanatory variable typically explain 90% of reproductive variance (Samson & Werk, 1986; Weiner et al., 2009). Instead, the relationship between vegetative and reproductive size in trees is far from being tight, and models with highly significant correlations usually only explain a low proportion of variance e.g. (House, 1992; Haymes & Fox, 2012; Granado-Yela et al., 2013). For long lived species, relevant variation of reproduction after accounting for size effects is left to be explained by additional factors like the environment or the genetic background (III).

An interesting case of loose allometric relationships between vegetative and reproductive size is described in *P. halepensis*, where some large trees, well beyond their

threshold size for reproduction produce none or very few cones (Climent *et al.*, 2008 and III). Under the assumption of a deterministic allometric model, those trees will eventually outperform smaller trees in terms of reproductive output. This is known as a bang-bang strategy (Falster & Westoby, 2003; Weiner *et al.*, 2009). Despite this assumptions seems reasonable, fuzzy vegetative-reproductive size relationships are common, and the fate of those trees is nonetheless uncertain. Could it be possible that trees with low reproductive allocation are also over-represented in the largest tree classes in forest stands? In such case, this could be the outcome of senescence (Genet *et al.*, 2010; Moran & Clark, 2012), an invalid explanation for young trees. Another option is that large trees with low or no reproductive investment could also arise merely due to (lack of) phenotypic costs of reproduction, as attaining large sizes may only be possible if investment in reproduction is low (Moran & Clark, 2012). Also, besides environmental plastic effects (III), and given within-population genetic variability for the allometry of reproduction (I, VI), the occurrence of such phenotypes can have a genetic basis.

As mentioned above, a genetic trade-off between investment in growth and reproduction is in fact a general expectation derived from life-history theory (Roff, 1992a), provided that no third variables interact (Roff, 2000). Under this assumption, a high genetic breeding value for growth implies a lower investment in reproduction, a possibility already mentioned in the forestry literature (Loehle et al., 1987). Genetic correlations between reproduction and growth reported in this study did not support this view unambiguously (I, III, V, VI and VII). It seems clear however, that both phenotypic and environmental correlations are positive, and that genetic correlations are smaller even though not always negative. As genetic correlations are known to be environmentaldependent (Sgrò & Hoffmann, 2004 and III), a meta-analytical approach will be useful to shed light on the issue. Also, studies that integrate a comprehensive set of fitness-related traits will provide a more realistic picture of adaptive strategies (Pigliucci, 2004). An integrated analysis of further traits such as constitutive and induced defences (Sampedro et al., 2011), drought resistance (Voltas et al., 2008), cold tolerance (Climent et al., 2009), phenology (Salmela et al., 2013), serotiny (Hernández-Serrano et al. in prep), stem architecture (Sierra-de-Grado et al., 1999), ontogenetic heteroblasty (Climent et al., 2013), bark thickness (Tapias et al., 2004) and seed size and dispersal ability (Santos-del-Blanco & Climent, 2011) will be highly valuable.

Despite inconsistencies in genetic correlations, empirical evidence for withinpopulation genetic variation in allocation to reproduction trading-off with variation in size is provided in (IV). Progeny of trees selected under natural conditions for good growth and stem form did also show reduced allocation to reproduction during their early developmental stages. It can be argued that those trees will also end up producing large amounts of cones later in development (bang-bang strategy), but the example at least provides evidence for the relevance of the genetic basis of reproduction, even under natural conditions where environmental variation is high (Wilson *et al.*, 2010). Other studies have also revealed negative genetic correlations between growth and other adaptive traits like drought resistance (Kaya *et al.*, 1994) and cold hardiness (Howe & Aitken, 2003), which altogether add to the importance of recording and integrating traits other than vegetative growth in order to define adaptation.

## 5.4 FITNESS TRAITS AND ADAPTIVE FOREST MANAGEMENT

The present study raises concern about considering size solely as a surrogate for fitness (III), even more if size is taken as a criterion to select trees to produce seed for regeneration in managed stands (Lefèvre *et al.*, 2013). Measuring fitness is a gigantic and mostly unreachable task in forest trees (Shaw & Etterson, 2012). Although seedling paternity analysis in forest stands can be used to relate realised fitness with certain phenotypic traits of parent trees (Burczyk *et al.*, 2006), large population sizes, large distribution areas and extensive gene flow distances together with changing selective pressures along development make the results only partially realistic.

Still, this kind of studies is hugely valuable to assign weighed importance to specific phenotypic traits such as size, and justify their use as surrogates of fitness (Moran & Clark, 2012). As expected, the few examples available to date do find a positive relationship between size and reproductive success (Oddou-Muratorio & Klein, 2008; Oddou-Muratorio *et al.*, 2010), but on top of that, also one between reproduction and reproductive success (González-Martínez *et al.*, 2006; Piotti *et al.*, 2009). Several studies in seed orchards have also shown a correlation between pollen production and reproductive success (Schoen & Stewart, 1986; Goto *et al.*, 2005; Moriguchi *et al.*, 2007; Hansen & Nielsen, 2010; Doerksen *et al.*, 2011).

Importantly, under a scenario of increased disturbance events such as forest fires and extreme drought (Lindner *et al.*, 2010), life expectancy of trees might be lower than in the present –particularly in the western Mediterranean basin, a hotspot of predicted climate change. Such scenario is likely to select for early reproduction, with trees investing resources in reproduction early in life (Kozłowski, 1992; Munguía-Rosas *et al.*, 2011) rather than expecting future unlikely rewards in terms of increased reproductive output.

The use of size as a surrogate for fitness is also frequently justified as height and diameter at breast height are much easier to estimate than reproductive output (Wu & Ying, 2004). It seems necessary then to improve and develop methods to estimate reproductive output fast and accurately (Koenig *et al.*, 1994, 2013; LaMontagne *et al.*, 2005) (I, III, VI). Though, for the same reasons stated above, namely negative genetic correlations, it will not be desirable to rely solely on reproductive output as a criterion for selection (Varghese *et al.*, 2009).

Recently a new paradigm of adaptive forest management has emerged, aiming to maintain evolutionary processes and adaptive potential of managed populations (Koskela *et al.*, 2013; Lefèvre *et al.*, 2013). In order to achieve it, a better understanding of evolutionary processes shaping fitness traits will be necessary. Also, as exposed throughout this study, phenotypic data obtained from common garden experiments will be relevant in order to characterise forest reproductive materials and to aid in sustainable forest management.

CONCLUSIONS

# 6. CONCLUSIONS

- I. *Pinus pinaster* and *P. halepensis* range-wide populations vary significantly in terms of reproductive life-history traits like the threshold size for reproduction, reproductive output and reproductive allometry.
- II. Genetically-based population mean values for reproductive life-history traits tend to vary clinally according to environmental factors, particularly those determining the length of the growing season.
- III. In *Pinus pinaster*, clinal variation was not found for the threshold size for male reproduction. This result could be explained by environment-independent male fitness gain curves, as conditions for pollen dispersal in this wind-pollinated species seem to be uniform across populations.
- IV. The allometry of reproduction shows putatively adaptive plasticity in *Pinus halepensis* as trees accelerate their sexual development under stressful conditions. This is a likely scenario to be found under climate change, and may provide tree populations with enhanced resilience.
- V. There is substantial within-population genetic variation for reproductive lifehistory traits, and these show higher heritabilities than growth traits, allowing for steady adaptive processes.
- VI. Somatic costs of reproduction in *Pinus halepensis* were detected by means of a manipulation experiment but not by a correlational study. In this species, reproduction had a greater effect in terms of future reproduction than in terms of growth.

- VII. Genetic costs of reproduction in terms of growth were more elusive to be detected and might be species- and environment-dependent. However, a selection experiment in *Pinus pinaster* aimed at increasing growth yield, resulted in a strong correlated negative response in reproductive traits, indicating a genetically based trade-off between growth and reproduction.
- VIII. Considering the prevalence of a loose relationship between reproduction and vegetative size in trees, and the likely existence of a genetic trade-off between reproduction and growth, it is advised against the use of size as a single surrogate of fitness. Instead, obtaining reproductive data is encouraged if criteria for achieving adaptive forest management are to be met.
- IX. To the extent that the described population variation is relevant for (static) adaptation, population characterization will be valuable for conservation and sustainable management of forest genetic resources. Standing genetic variation and differentiation also provides insight into past adaptive processes and inform about likely future evolutionary responses.

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# 7. **REFERENCES**

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Lausana, Suiza, 18 de septiembre de 2013

## APPENDIX

## 9. APPENDIX

## Articles published in Science Citation Index journals:

- (I) Variation of early reproductive allocation in multi-site genetic trials of Maritime pine and Aleppo pine. Santos-del-Blanco L, Notivol E, Zas R, Chambel MR, Majada J, Climent J. 2010. Forest Systems 19: 381–392.
- (II) Genetic differentiation for size at first reproduction through male versus female functions in the widespread Mediterranean tree *Pinus pinaster*. Santos-del-Blanco L, Climent J, González-Martínez SC, Pannell JR. 2012. Annals of Botany 110: 1449–1460.
- (III) Plasticity in reproduction and growth among 52 range-wide populations of a Mediterranean conifer: adaptive responses to environmental stress. Santos-del-Blanco L, Bonser SP, Valladares F, Chambel MR, Climent J. 2013. Journal of Evolutionary Biology. 26: 1912-1924.

### **Complementary manuscripts and conference proceedings:**

- (IV) Correlated effects on reproductive traits derived from domestication of a forest tree are relevant for future forest resilience. Santos-del-Blanco L, Alía R, González-Martínez S, Sampedro L, Lario F, Climent, J. 2013. Manuscript.
- (V) Somatic and genetic costs of reproduction in a conifer tree. Santos-del-Blanco
   L, Climent J. Manuscript.
- (VI) Guidelines for estimation of quantitative genetic parameters of reproductive traits. Santos-del-Blanco L, Climent J. Manuscript.
- (VII) Ecología evolutiva de la reproducción en dos pinos mediterráneos: *Pinus pinaster* y *Pinus halepensis*. Santos-del-Blanco L, Chambel R, Notivol E. Alía R, Climent J. Proceedings 6º Congreso Forestal Español. Sociedad Española de Ciencias Forestales, Vitoria, 2013.

# APPENDIX I

Variation of early reproductive allocation in multi-site genetic trials of Maritime pine and Aleppo pine. **Santos-del-Blanco L**, Notivol E, Zas R, Chambel MR, Majada J, Climent J. 2010. Forest Systems 19: 381–392.

## Variation of early reproductive allocation in multi-site genetic trials of Maritime pine and Aleppo pine

L. Santos-del-Blanco<sup>1,5</sup>\*, R. Zas<sup>2</sup>, E. Notivol<sup>3</sup>, M. R. Chambel<sup>1,5</sup>,

J. Majada<sup>4</sup> and J. Climent<sup>1,5</sup>

<sup>1</sup> CIFOR-INIA. Madrid. Spain
 <sup>2</sup> CITA. Zaragoza. Spain
 <sup>3</sup> MBF-CSIC. Pontevedra. Spain
 <sup>4</sup> SERIDA. Asturias. Spain
 <sup>5</sup> Sustainable Forest Management Research Institute UVa-INIA. Palencia. Spain

### Abstract

Life histories in Mediterranean pines are well known to be closely related to different fire and disturbance regimes. Variation in these factors is also reflected in reproductive strategies at the specific and intraspecific level. Specifically, the onset of reproduction is a crucial stage for any organism because it has profound implications on fitness. In this paper we focus on the intraspecific variation and plasticity in the threshold size for reproduction and reproductive allocation in two Mediterranean pines, assessed at the onset of reproduction, when trade-offs between reproduction and growth are expected to be greater. Replicated common garden provenance and progeny trials of Aleppo pine and Maritime pine were used to estimate genetic parameters for reproduction and vegetative growth, as well as variation in plasticity in reproductive strategies at the intraspecific level. In both species, high variation among populations was found for both threshold size for reproduction and for reproductive allocation. Reproductive allocation was also highly variable within populations and showed moderate to high values of heritability and high coefficients of additive genetic variation. These results indicate a high genetic control of these reproductive traits, while high additive genetic variation is maintained, allowing to face selective pressures. Moreover, reproductive strategies although plastic, showed low genotype x environment interaction, and intraspecific variation was highly consistent across trial sites both at the population and the family levels. The former data confirm the strong genetic control of reproductive traits in these species. Finally, the fact that Aleppo pine starts its reproductive phase as female while Maritime pine can start reproducing either as male or female deserves further attention.

Key words: *Pinus halepensis*; *Pinus pinaster*; genetic variability; plasticity; trade-offs; early reproductive strategies; threshold size for reproduction.

#### Resumen

#### Variación en asignación reproductiva temprana en ensayos multi-localidad de pino carrasco y pino negral

Las estrategias de historia vital en pinos mediterráneos están estrechamente ligadas a diversos regímenes de incendios y perturbaciones. A su vez, la variabilidad de esos factores también se ve reflejada en las estrategias reproductivas a niveles inter e intraespecíficos. Concretamente, el comienzo de la reproducción es una etapa crucial para cualquier organismo debido a su profunda influencia en su adaptación al medio. Este trabajo se enfoca en el estudio de la variabilidad intraespecífica y la plasticidad en el tamaño umbral de reproducción y la asignación reproductiva en dos pinos mediterráneos. El estudio se realizó durante el comienzo de la fase reproductiva, momento en que se estima que la compensación entre reproducción y crecimiento es de mayor importancia. Se utilizaron ensayos multi-sitio de procedencias y progenies de pino carrasco y pino negral para estimar los parámetros genéticos de caracteres reproductivos y de crecimiento vegetativo, así como para conocer la variabilidad en la plasticidad de las estrategias reproductivas a nivel intraespecífico. En ambas especies se halló una alta variabilidad entre poblaciones para el tamaño umbral de reproducción y para la asignación reproductiva. La asignación reproductiva también fue altamente variable dentro de poblaciones y mostró valores de heredabilidad de moderados a altos y altos coeficientes de varianza genética aditiva. Estos resultados indican un alto control genético de los rasgos reproductivos, mientras se mantiene una alta varianza genética, permitiendo afrontar futuras presiones selectivas. Además, a pesar de la existencia de plasti-

<sup>\*</sup> Corresponding author: santos.luis@inia.es Received: 15-04-10; Accepted: 21-07-10.

cidad en las estrategias reproductivas, éstas mostraron una baja interacción genotipo  $\times$  ambiente, y la variación intraespecífica fue acorde en los diferentes lugares de ensayo tanto a nivel poblacional como familiar. Los datos anteriores confirman el alto control genético de los rasgos reproductivos en estas especies. Finalmente, el hecho de que el pino carrasco comience su fase reproductora como hembra mientras que el pino negral pueda comenzar su reproducción bien como macho o bien como hembra, merece una atención más detallada.

Palabras clave: *Pinus halepensis; Pinus pinaster;* variabilidad genética; plasticidad; trade-offs; estrategias de reproducción temprana; tamaño umbral de reproducción.

## Introduction

Due to its direct link to fitness, information on the intraespecific variation in the time at which organisms reach maturity and the amount of resources allocated to reproduction is central to understand how evolution has shaped these traits in the past and to infer how can they be affected by future evolutionary forces. In long-living plant species, optimal size or age at maturity reflects the balance between the benefits of an early reproduction and its costs reflected in disminished future reproduction and survival (Wesselingh *et al.*, 1997). Theory predicts that mortality risk and its predictability drive the time at first reproduction within and among species (Kozlowski, 1992).

Pines are known to have developed astonishing adaptations to fire such as the grass stage or serotiny, but also a group of other life history traits, being reproductive traits like precocity and intensity of early reproduction among the most important (Keeley and Zedler, 1998). Species adapted to frequent crown fires and unable to resprout, are expected to show an early intense reproduction while species not adapted to fire or adapted to ground fires typically show a delayed reproduction (Agee, 1998). Mediterranean pine ecosystems are frequently affected by forest fires and droughts (Richarson *et al.*, 2007), and thus Mediterranean pines can provide a good example for studying the relationship between early reproductive strategies and disturbance regimes.

General reproductive strategies are known for many pine species. However, there is a lack of information at the intraspecific level. The vast majority of the studies at the intraspecific level are biased towards economically important species, related to breeding programs interested in reducing the time between cycles, increasing seed crops and avoiding unequal contributions from some genotypes in seed orchards (Koenig and Knops, 2000; Kang *et al.*, 2003). Consequently, an evolutionary and ecological discussion on this topic is not common. In widely distributed Mediterranean pines like Aleppo pine (*Pinus halepensis* Mill.) and Maritime pine (*Pinus pinaster* Ait.), information on reproductive trait variation within species holds great interest for understanding adaptation to contrasting local conditions as seen, for example, in the model genus *Arabidopsis* (Bonser and Aarssen, 2001; Rutter and Fenster, 2007). This information can also prove useful for an optimum deployment of genetic materials as a mean to increase forest resilience facing frequent disturbances and to facilitate adaptation to climate change (Thompson *et al.*, 2009). Common garden trials of forest trees, usually planted for genetic breeding, can offer valuable information on these subjects, provided the necessary cooperation between foresters and evolutionists.

Extensive research on the ecology and population genetics regarding P. pinaster and P. halepensis is available, but only few references focus on reproduction (Richardson, 1998), despite its close relation to fitness. Although both species have distribution areas that overlap partially at several points within the Iberian Peninsula, some differences are remarkable. Maritime pine spreads across the western Mediterranean basin from North Morocco with Mediterranean climate to South Western Atlantic coast of France, with a humid Atlantic climate. Three different main gene pools have been differentiated (Bucci et al., 2007). On the other hand, Aleppo pine has a circunmediterranean distribution with genetically diverse populations in Greece and Turkey but more genetically uniform populations towards the west of the basin following a proposed colonization route (Grivet et al., 2009).

According to its genetic diversity, common garden trials have shown a high variability in reproductive strategies for Maritime pine (Tapias *et al.*, 2004) and variable differentiation in Aleppo pine populations depending on the trait and experimental site (Climent *et al.*, 2008). Genetic differentiation among populations for reproductive traits is proposed to reflect local selective pressures, consistent with empirical examples about how fire can act to shape early reproductive allocation (González-Ochoa *et al.*, 2004; Gil *et al.*, 2009).

Genetic parameters such as narrow sense heritability  $(h^2)$ , additive genetic coefficient of variation  $(CV_a)$ , the quantitative differentiation between populations  $(Q_{st})$ , and genetic correlations among traits are relevant to describe the genetic control of reproductive traits, to assess quantitative variation within and among populations and to describe the existence of trade-offs between traits (Roff, 2000). The comparison of the genetic architecture of traits closely linked to fitness with that of other traits is scientifically challenging, since it deals with the interplay between past directional selection (reducing within population variation and promoting differentiation between populations) and the availability of enough additive genetic variation to enable future evolutionary processes (Merilä and Sheldon, 1999). Furthermore, there is growing evidence showing the importance of plasticity in trees affecting the estimation of genetic parameters and the necessity to conduct experiments with genetic entries replicated in a range of environmental conditions (Sgrò and Hoffmann, 2004).

In this work, we focus on describing the intraspecific variability in multi-site genetic trials of *P. halepensis* and *P. pinaster* for reproductive traits, namely threshold size for reproduction and reproductive allocation from an evolutionary quantitative genetic approach. We aim also to determine whether early reproduction in these species entail vegetative fitness costs, which will be reflected as negative genetic correlations between reproductive and vegetative traits.

### Material and methods

#### Field trials and Plant material

For Aleppo pine, we used a provenance-progeny trial replicated in two sites in inland Spain (Megeces —AMEG—, and Montañana —AMON—) comprising 148 open-pollinated families of 32 populations covering the species' natural range in the Iberian Peninsula and Balearic Islands plus three additional sources from planted stands of unknown origin (Table A2). One year old seedlings were planted in 1995 in a randomized complete block design with seven blocks, and two contiguous plants per plot. Spacing was 2.5 × 2 m at AMEG and 5.2 × 1 m at AMEG and AMON. Trial sites were ecologically contrasting (Table A1): AMEG is situated

on a dry shallow calcic soil with < 15% slope in the Castilian Plateau; as a result of the harsher conditions, mortality in this site was high (33%). Despite being outside the species natural range, AMEG lies within an area with extensive Aleppo pine plantations. AMON is sited on a deep fertile alluvial soil, well irrigated during summer, and more favorable for pine growth, although mortality after plantation affected many seedlings.

**Maritime pine** trials are represented by a progeny trial replicated in two sites (Rebordelo —PREB—, and Rianxo —PRIA—) and a provenance-progeny trial, also replicated in two sites (A Merca —PMER—, and Cavada —PCAV—), all located at Northwestern Spain, under temperate Atlantic climate (Table A1). Soil and climate in PREB and PCAV are rather similar, with high annual and summer precipitation, while PMER represents a transition towards Mediterranean conditions, with much lower summer rainfall and PRIA is representative of mild winter coastal conditions.

In PREB and PRIA, 28 open-pollinated (o.p.) families of superior trees selected within the Atlantic coast of Galicia were planted in 2003 under different establishment fertilization treatments. The original experimental design included three unimproved seed lots that were not considered in this study. The experimental layout in both sites was a split-plot design in ten blocks, with nine fertilization treatments acting as the main factor and the genetic entries as the split factor (see details in Martíns et al., 2009). Althoug fertilized plants attained bigger size and more cones, fertilization did not affect the ratio between cones and tree size *i.e.* reproductive allocation (data not shown), therefore fertilization was not further considered in this work. Both progeny trials were thus considered to follow a randomized complete block design with 90 blocks and single-tree plots. Spacing was  $3 \times 2$  m.

In PMER and PCAV, 250 open-pollinated families pertaining to 26 natural populations covering most of the natural range were planted in 2005 (Table A3). Experimental layout was a complete randomized block design with 4 plants per family and block and 4 blocks. Spacing was  $3 \times 2$  m.

#### Assessments

Reproductive and growth-related variables were measured in all sites at young ages, after a significant proportion of the trees started to produce cones. Nevertheless, it was not possible to measure the same variables coding for reproduction or growth in all sites due to differences in development.

In the Aleppo pine trials, basal diameter and total tree height were measured and cones belonging to different cohorts were counted in 2005 and 2009 when trees were 11 and 15 years old, respectively. The small tree size and good visibility in the AMEG site allowed for distinguishing up to three cone cohorts, representing yearly reproduction. In winter 2005 and 2009, first and second year developing conelets were counted. Male reproduction was also recorded through a binary variable, present or absent. Regarding its reproductive status, each individual tree was classified as male (protandrious, with only male cones), female (protogynous, with only female cones), synchronous (with both male and female cones) or juvenile (no cones).

In AMON, only one measurement was carried out in winter 2009, recording basal diameter and tree height. The better growing conditions in this site, compared with AMEG, were reflected in a more vigorous growth and more advanced ontogenic development. Trees were too high to allow a clear visibility of all developing cones and to distinguish them from older ones. Since Aleppo pine cones remain attached to the branches, either opened or as serotinous cones, we used an alternative method to estimate accumulated cone production trough the tree life. Cones counted in 15 seconds were taken as a surrogate of reproduction (Koenig et al., 1994). According to a preliminary sub-sample, counting cones during 15 seconds was considered to provide a reliable estimation of the total cone number per tree, comparing estimations in trees with different cone loads.

Reproductive allocation, RA, (following Karlsson and Méndez, 2005), was calculated for each tree as a ratio between the number of cones (sum of developing first and second year conelets (Cone count, Cc) for AMEG and the total number of cones counted in 15 seconds for AMON) and stem volume over bark (Vob), a surrogate for biomass, and hence, resource availability (Climent *et al.*, 2008). Vob was calculated according to the formula:

$$Vob = \frac{\pi}{12} Db^2 H$$

where Db is basal diameter and H is total tree height.

In the PRIA and PREB maritime pine progeny trials, total tree height and basal diameter were measured in December 2007, when trees were 5 years old. First and second year conelets of the two coexisting cohorts within the crown were counted in all trees, and RA was estimated as described before.

In PMER and PCAV height was measured, first and second year conelets were counted and male reproductive status was recorded as a binary variable in 2009, when trees were 5 years old. This was the time in which a significant proportion reached maturity since in previous years reproduction was almost absent. As in Aleppo pine, individuals were classified as juvenile, male, female or synchronous. We used data from this trial series exclusively to assess the variation and plasticity of the threshold size for male and female reproduction.

#### Data analysis and genetic parameters

Threshold size for reproduction (TSR) was studied with a logistic model similar to that used by other authors (Wesselingh et al., 1997; Méndez and Karlsson, 2004). Reproduction probability was analyzed by adjusting a variable termed CATREP, representing the reproductive status of an individual (0, non reproductive, 1, reproductive, bearing female and/or male cones). Since we were interested mostly in the variation between populations for TSR, we applied this analysis to the provenance-progeny trials at the time of maximum differentiation, that is, when close to 50% individuals were reproductive. Sites AMEG of Aleppo pine in 2005 and PMER and PCAV of Maritime pine fulfilled this requisite. Logistic curves were adjusted with size (stem volume over bark in P. halepensis or height in P. pinaster) as a quantitative factor. A first analysis was made considering all populations per site as a categorical factor in order to test for its significance. Then, a curve was fitted for each provenance (Fig. 1):

$$CATREP = \frac{e^{(a+bx)}}{1+e^{(a+bx)}}$$

being *a* and *b* coefficients adjusted for each regression and *x* was either *Vob* in Aleppo pine or H in Maritime pine.  $V_{50}$  or  $H_{50}$  were defined as the volume or height at which the probability for a tree to have reached sexual maturity is 50% (Méndez and Karlsson, 2004).

#### Genetic parameters

Target variables *Vob, Cc* and *RA* were analyzed by Mixed Linear Models testing for variability at provenance and family within provenance levels as follows:



**Figure 1.** Example of a logistic curve representig probability of reproduction as a function of size (stem volume over bark) in Pinus halepensis from Hijar, NE Spain. Dashed lines indicate the 75% confidence intervals. The vertical line denotes the volume  $V_{50}$  at which the probability for a tree to have reached sexual maturity is 50%.

$$Y_{i,i,k} = \mu + P_r + F_i(Pr_i) + B_k + \varepsilon$$

where Y is the dependent variable,  $\mu$  is the general mean, Pr is the random effect of the provenance, F(Pr) is the random effect of family within provenance, B is the random effect of the block, and  $\varepsilon$  is the error term. The two progeny trials PREB and PRIA, were analyzed with an equivalent linear model without the provenance effect, and excluding the three unimproved seed sources from the analysis.

Narrow sense heritability  $(h^2)$  was calculated assuming the open-pollinated families as true half-sibs:

$$h^2 = \frac{\sigma_A^2}{\sigma_P^2} = 4 \frac{\sigma_f^2}{\sigma_f^2 + \sigma_\epsilon^2}$$

where  $\sigma_A^2$  is the additive variance,  $\sigma_P^2$  is the phenotypic variance,  $\sigma_f^2$  is the familiar variance and  $\sigma_{\epsilon}^2$  is the variance due to error.

Quantitative trait variation  $(Q_{st})$  was calculated as

$$Q_{st} = \frac{\sigma_f^2}{\sigma_f^2 + 2 \cdot 4\sigma_{f(Pr)}^2}$$

where  $\sigma_{f(Pr)}^2$  is the family variance within provenances.

Coefficient of additive genetic variance  $CV_a$  was defined as

$$CV_a = \frac{\sigma_A^2}{\mu}$$

being  $\mu$  the general mean.

Phenotypic  $r_p$  and genetic correlations  $r_A$  were calculated according to

$$r_{A} = \frac{COV_{f}(x,y)}{\sqrt{\sigma_{fx}^{2} \cdot \sigma_{fy}^{2}}}$$
 and  $r_{P} = \frac{COV(x,y)}{\sqrt{\sigma_{x}^{2} \cdot \sigma_{y}^{2}}}$ 

where  $COV_{xy}$  is the covariance between any two variables x and y and  $\sigma_x^2$  and  $\sigma_y^2$  are their corresponding variances.  $COV_{fxy}$  is the family variance, obtained as follows:

$$COV_{fxy} = \frac{\sigma_{fxy}^2 - \sigma_{fx}^2 - \sigma_{fy}^2}{2}$$

where  $\sigma_{fxy}^2$  is the family variance of a composite variable resulting from the sum of any two variables *x* and *y*.

#### Results

#### Threshold size for reproduction

In *Pinus halepensis* trials, mean height was higher and ontogenic development more advanced in AMON than in AMEG as a result of less limiting ecological conditions. In *P. pinaster* PMER and PCAV trials had a similar development, with slightly bigger trees but less sexually developed in PCAV. In PREB and PRIA trials, height was also very similar but in PRIA the number of reproductive trees was lower. Overall, survival in *P halepensis* trials was lower than in *P. pinaster* ones (Table1).

In Aleppo pine (AMEG trial assessed in 2005), both Vob ( $\chi^2 = 561.9$ , p < 0.0001) and population ( $\chi^2 = 70.3$ , p < 0.0001) contributed significatively to the fact of being adult or juvenile (CATREP). There was a 3.6-fold difference among populations for V50 and a 8.5-fold difference for RA. V50, despite large errors, showed a significant correlation with mean RA at the population level (Fig. 2). Trees reached maturity at a mean height of 243 cm, and this was virtually always as females since just 2 out of 1,493 trees bore male cones only (Fig. 3).

Similarly, in the two Maritime pine trials (PMER and PCAV sites) both tree height ( $\chi^2$ =512.9 for PMER, and  $\chi^2$ =93.5 for PCAV, p<0.0001) and population ( $\chi^2$ =114.2 for PMER and  $\chi^2$ =164.5 for PCAV, p<0.0001) had a significant effect on CATREP. There was a 2.4-fold (152 cm) difference between the most and lest precocious populations in PMER, and a 3.3fold (226 cm) difference in PCAV, with strong among provenance correspondence between the two test sites (Fig. 4). Unlike Aleppo pine, Maritime pine trees reached maturity either as males or females (Fig. 3). In both sites, 95% confidence level intervals showed

Trial	Plants	Populations	Families	Assessment age (yrs)	Survival (%)	H (cm)	R (%)
Aleppo pine							
AMEG	2,182	32	148	11/15	67.0	217	73.8
AMON	2,037	32	148	15	67.1	784	99.8
Maritime pine							
PMER	3,152	25	217	5	81.0	161	64.2
PCAV	3,456	25	224	5	83.9	182	52.4
PREB	2,007		28	5	80.9	359	75.4
PRIA	2,098		28	5	83.2	350	64.7

**Table 1.** Summary information of Aleppo pine and Maritime pine genetic trials: code, number of plants, number of populations, number of families, assessment age (years), survival (%), average tree height (cm) and reproductive trees (%)

differences between average heights in juvenile, and female trees respect to male and synchronous ones (Fig. 3).

## Intraespecific variation in reproductive allocation and tree size

In Aleppo pine, both volume and reproductive allocation differed significantly among populations (p < 0.001) and among families within populations (p < 0.001) in both years, although in 2005, variation among families within populations was just marginally significant (p < 0.10) for RA. Because measures in RA were not comparable between the two sites, variation in plasticity was ilustrated by rank Spearman correlation in



**Figure 2.** Relationship across populations between reproductive allocation (RA) in 2005 in AMEG and the corresponding Volume over bark at which the reproduction probability was 50% ( $V_{50}$ ). Each point represents a population. Vertical lines indicate 75% confidence intervals for  $V_{50}$  and horizontal lines standard errors for RA in year 2005.

2009 (Fig. 5). A consistent behaviour was found among populations between both sites for RA ( $\rho = 0.80$ ), indicating limited population × site interaction; *i.e.* low differences for plasticity between populations.

Significant within population variation (p < 0.001) in RA and Vob was also observed in the *P. pinaster* trials, with a very high genetic correlation among sites for RA (Fig. 6). The high correspondance between sites in reproductive traits constrast with the strong genotype × environment interaction observed for growth (r=0.13 for Vob vs r=0.89 for RA).

Heritabilities for RA ranked from 0.27 to 0.63 and they were higher than those for Vob (0.14-0.22). Heritabilities for RA were also more variable than for Vob. The highest value was attained in the Aleppo pine trial AMEG in 2009. The coefficient of quantitative variation (Qst) did not show a consistent difference for RA and Vob. Hence, Qst for RA was much higher than for volume in AMEG in 2005, similar in 2009 but lower in AMON. However, the greatest difference for genetic parameters between reproductive allocation and tree size was that of the coefficients of additive genetic variance (CVa), which were consistently higher (up to 5.6-fold) for RA across species, sites and years (Table 2).

#### Phenotypic and genetic correlations

Phenotypic correlations between RA and Vob were negative in all cases except for AMEG in 2005, being between moderate and low ( $r_P = -0.18$  to 0.11) except in AMON ( $r_P = -0.37$ ). Genotypic correlations were also negative but much stronger ( $r_A = -0.30$  to -0.97) than phenotypic correlations except in AMEG in 2009 ( $r_A = -0.05$ ) (Table 3)



**Figure 3.** Proportions of juvenile (white), male (pale grey), female (dark grey) or synchronous (black) individuals in AMEG (Aleppo pine) and in PMER and PCAV (Maritime pine) trial sites. Numbers indicate average  $(\pm s.e)$  height for each group or site.

## Discussion

Our results show significant intraspecific variation on reproductive traits both in Aleppo pine and in Maritime pine, consistent with previous information. Reproductive allocation was studied here during the onset of reproduction, when it is most relevant in the case of short fire return intervals and when trade-offs with vegetative growth are expected to be greater (Wesselingh *et al.*, 1997).

#### Intraespecific variation in threshold size for reproduction and reproductive allocation

TSR in plants has been assessed in the frame of developmental biology mainly for herbaceous plants, existing however only few examples (but see Méndez and Karlsson, 2004). Estimation of intraspecific TSR in trees is inherently challenging due to their late maturity (up to many years), and their relatively larger size. There is however some information regarding age

**Table 2.** Genetic parameter estimates for early reproductive allocation (*RA*) and Volume over bark (*Vob*) in different Aleppo pine and maritime pine sites and ages.

<u>Carocian</u>	Site	<b>A</b> = -	$h^2$		$Q_{st}$		CVa	
Species	Site	Age -	RA	Vob	RA	Vob	RA	Vob
Aleppo pine	AMEG	11	0.29	0.22	0.48	0.18	82.29	43.31
	AMEG	15	0.63	0.22	0.12	0.12	100.56	33.02
	AMON	15	0.27	0.14	0.21	0.29	40.23	21.91
Maritime pine	PREB	5	0.47	0.14			82.34	11.65
*	PRIA	5	0.32	0.18			71.21	17.51

 $h^2$ : narrow sense heritability.  $Q_{st}$ : coefficient of quantitativ variation. CVa: coefficient of additive genetic variance.



**Figure 4.** Relationship across populations between threshold size for reproduction ( $H_{50}$ , height at which reproduction probability is 50% for a given population) in two Maritime pine trial sites, PMER and PCAV. Vertical and horizontal lines indicate 75% confidence intervals for  $H_{50}$ . Lines were not included if logistic curve adjusting for a given population and site was not significant.

or size at maturity for some tree species, including Mediterranean pines (Schmida *et al.*, 2000), but most studies lack any control of environmental conditions or genetic background.



**Figure 5.** Relationship between the population ranking for early reproductive allocation (RA) estimated in two contrasting *P. halepensis* sites in 2009, AMEG and AMON. Each point represents a population.



**Figure 6.** Family relationship in early reproductive allocation between two *P. pinaster* progeny trials, PREB and PRIA. Each point represents an open-pollinated familiy. Vertical and horizontal lines denote standard errors.

In Aleppo pine three main findings stand out. First, we found large differences between populations in both threshold size for reproduction (TSR) and reproductive allocation (RA), consistent with previous results in this species (Climent et al., 2008) despite Spanish populations were reported to be more genetically uniform (Grivet et al., 2009). Second, a close inverse relationship between reproductive allocation (RA) and TSR was found (Fig. 2) confirming that precocious populations tend also to invest more in reproduction. Third, reproductive allocation at the population level showed a consistent pattern across sites (AMEG and AMON) despite the widely contrasting experimental environments. This evidence supports the idea that the differential reproductive strategies among populations are not affected by the environmental conditions, and remain largely consistent across environments, even though as it was the case in AMON, growth conditions were almost unlimited. This high genetic control for female reproduction in Aleppo pine is in line with results from breeding programs reporting high heritability for flower and cone production and consistent behaviour along time (Matziris, 1997).

Significant variability between populations for TSR was also found in Maritime pine and, as in Aleppo pine, with a high consistency of behaviours across environments (*i.e.* low genotype  $\times$  environment interaction). However, contrasting with Aleppo pine, male reproduction was a highly relevant factor to explain those differences. It has been described an enhanced male reproductive allocation in pines as a result of disturbances such as herbibory or shadow (Schmida *et al.*,

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Species	Site	<b>A</b> 60	Cc-I	Vob	RA-Vob		
Species	Site	Age	r <sub>a</sub>	<b>r</b> <sub>p</sub>	r <sub>a</sub>	$r_p$	
Aleppo pine	AMEG AMEG AMON	11 15 15	0.21 0.42 -0.93	0.56 0.49 0.29	$-0.30 \\ -0.05 \\ -0.97$	$0.11 \\ -0.09 \\ -0.37$	
Maritime pine	PREB PRIA	5 5	$\begin{array}{c} -0.21\\ -0.21\end{array}$	0.16 0.12	$-0.36 \\ -0.42$	$-0.12 \\ -0.18$	

**Table 3.** Phenotypic  $(r_p)$  and genetic  $(r_a)$  correlations between cone count (Cc) and stem volume over bark (*Vob*) and between reproductive allocation (*RA*) and stem volume over bark for Aleppo pine and Maritime pine at different sites and measurement times. All correlations were significant

2000; Cobb et al., 2002) but to our knowledge, little attention has been paid to interspecific variation in early sex allocation (Richardson, 1998). Sexual specialization patterns between genotypes in relation to environmental conditions should be checked in the future, at the light of the diverse life history traits closely related to fire regime (Tapias et al., 2004). An early female but not male reproduction has been suggested as a consequence of a higher reproductive success in post-fire situations where some adult trees have survived in the surroundings and can pollinate precocious protogynous individuals (Ne'eman et al., 2004). Our data suggest differential selection pressures between Maritime pine populations that could be related to a complex combination of population history, local environment and perturbation regimes.

Moreover, a high additive genetic control for female reproductive allocation between families was found in Maritime pine. In PRIA and PREB, the almost perfect genetic correlations between both sites confirm that although plasticity exists, genetic differences of plasticity between families are negligible. This strong genetic correlation emerges in spite of heavy damage caused by pine weevil Hylobius abietis L. in the PRIA trial during the two first years after planting. The pine weevil attack, which differentially affected families (Zas et al., 2005), caused deep alterations of resource allocation and pine growth patterns (Sampedro et al., 2009). Although it is known that herbivory may also affect reproduction traits in pine trees (Cobb et al., 2002; Mueller et al., 2005), it seems to be not the case here, as family variation in reproductive allocation remained fairly consistent irrespective of the incidence of the herbivore. This meaningful result links to a wide scientific discussion about the plasticity of fitness traits, that largely overpasses the objectives of this paper (Schlichting, 2002; Sultan, 2003).

#### **Evolutionary implications**

The moderate or high differentiation in reproductive allocation among populations  $(Q_{st})$  is in agreement with a high fitness value of female early reproductive allocation, as it had been postulated based on the species' life histores (Ne'man et al., 2004; Tapias et al., 2004; Climent et al., 2008). Moreover, the very high additive genetic variation between families across populations  $(CV_a)$  observed both in Aleppo and Maritime pine could be also interpreted as a sign of high fitness value (Merilä and Sheldon, 1999) despite current controversy in this issue (Glazier, 2002). Nontheless, if a high differentiation between populations can be thought as a fingerprint of different directional selection processes in each environment, a high additive genetic variation provides the fuel for future adaptation to fast changes in perturbation regimes associated to global change. A precocious high female fecundity is advantageous in fire-prone habitats, as seen dramatically in Spain in repeteadly-burned forest stands (Gil et al., 2009) but it can also be thought to increase overall fitness under other environmental constraints, as postulated in Arabidopsis (Rutter and Fenster, 2007).

But early female fecundity, as a part of a reproductive strategy is not cost free, and seems to have implications in other traits, as reflected by the negative genetic correlations between reproductive allocation and tree size, also reported previously in pines (Schmidtling, 1981). Our data point towards a clear antagonism between reproduction and growth in both pine species, consistent with previous results in Aleppo pine, showing that most abundant cone yields are produced by middle-sized, not bigger individuals (Climent *et al.*, 2008). However, it should be noted that genetic correlation between reproductive allocation and growth were estimated here upon two non-independent variables, Vob and RA = Cc / Vob. The inclusion of Vob in the denominator for RA estimation may imply a mathematical artifact leading to spurious correlation (Brett, 2004). Although the negative correlation between cone count and volume observed in all trials except AMEG (Table 3) is supporting that the trade-off between growth and early reproduction does exist, bootstrap or Montecarlo simulations should be used in the future to confirm the actual magnitude of this trade-off (Brett, 2004). Actually, costs of reproduction measured as trade offs between reproductive allocation and vegetative growth are a classic and prolific research field, although mainly focused in herbaceous species (Karlsson and Méndez, 2005). Evaluation of costs of reproduction in trees is more challenging than in herbaceous plants, and thus examples are scarce and almost absent in Mediterranean pines. Some examples exist in which no costs were found, even for masting species like Picea abies (Seifert and Müller-Starck, 2009) and Fagus crenata (Yasumura et al., 2006). To explain that, the existence of compensatory mecanisms (resource storage, enhanced resource acquisition), rather than actual absence of costs of reproduction has been proposed (Karlsson and Méndez, 2005).

Results shown here point to a higher than suspected diversification among Aleppo pine and Maritime pine populations in early reproductive allocation, providing an excellent example of evolution as a response to ecological conditions in two widespread species. The fact that Maritime pine can start its reproductive phase either as male or female, while Aleppo pine consistently starts as female (confirmed by authors' unpublished data) deserves further attention, integrating genetic and environmental control and the different costs of male and female reproduction.

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Appendix

Species	Site (abr.)	Trial	Altitude	М (°С)	М (°С)	P (mm)	Sp (mm)	Longitude	Latitude
Pinus halepensis	Megeces AMEG	P-p	779	12.1	-0.1	413	66	4°33'30" W	41°25'18" N
-	Montañana AMON	P-p	216	14.9	1.5	350	70	0°49'31" W	41°41'5" N
Pinus pinaster	A Merca PMER	P-p	454	12.8	2.5	1,018	92	7°56'17" W	42°14'42" N
*	Cavada PCAV	P-p	349	12.6	3.1	1,328	183	6°32'36" W	43°25'15" N
	Rianxo PRIA	p	90	14.6	5.6	1,866	165	8°46'49" W	42°41'7" N
	Rebordelo PREB	p	530	12.5	2.8	2,335	235	8°28'35" W	42°27'40" N

Trial: P-p, provenance-progenies, p, progenies. M: mean annual temperature. m: mean temperature of the coldest month. P: mean annual rainfall. Sp: summer rainfall.

**Table A2.** List and location of the Spanish populations comprised in the AMEG and AMON Aleppo pine provenance progeny trials. Asterisks indicate seed sources from Northern Plateau afforestations **Table A3.** List and location of the populations comprised in PMER and PCAV Maritime pine provenance progeny trials

Population	Latitude	Longitude		
Altura	39°47' N	0°36' W		
Benamaurel	37°42' N	2°44' W		
Benicasim	40°04' N	0°01'E		
Cabanellas	42°14' N	2°47' E		
Carratraca	36°50' N	4°50' W		
Cazorla	38°06' N	2°47' W		
Escorca	39°49' N	2°53' E		
Frigiliana	36°49' N	3°55' W		
Hijar	41°06' N	0°25' W		
Lentegi	36°49' N	3°41' W		
Luna	42°13' N	0°00' W		
Monroyo	40°47' N	0°01'E		
Palma de Mallorca	39°08' N	2°56' E		
Paterna	38°37' N	2°16' W		
Quesada	37°44' N	3°09' W		
Ricote	38°08' N	1°25' W		
Sant Salvador de Guardiola	41°40' N	1°45' E		
Santanyí	39°17' N	3°02' E		
Santiago de la Espada	38°13' N	2°28' W		
Tibi	38°31' N	0°38' W		
Tivissa	41°03' N	0°00'E		
Tuéjar	39°49' N	1°09' W		
Valbuena de Duero*	41°39' N	4°17' W		
Valdeconcha	40°26' N	2°52' W		
Valtablado del Río	40°44' N	2°23' W		
Vega de Valdetronco*	41°35' N	5°04' W		
Velez Blanco	37°47' N	2°00' W		
Villa de Ves	39°10' N	1°14' W		
Villajoyosa	38°29' N	0°18' W		
Villanueva de Huerva	41°21' N	1°03' W		
Villavieja de Tordesillas*	41°36' N	4°55' W		
Zuera	41°55' N	0°55'W		

Population	Latitude	Longitude	Country
Alto de la Llama	43°17' N	6°29' W	Spain
Arenas de San Pedro	40°11' N	5°06' W	Spain
Armayán	43°18' N	6°27'W	Spain
Bayubas de Abajo	41°31' N	2°57'W	Spain
Cadavedo	43°32' N	6°25'W	Spain
Carbonero	41°10' N	4°16' W	Spain
Castropol	43°30' N	6°58' W	Spain
Cenicientos	40°16' N	4°29' W	Spain
Coca	41°15' N	4°29' W	Spain
Cuellar	41°22' N	4°29' W	Spain
Lamuño	43°33' N	6°13'W	Spain
Leiria	39°47' N	8°57' W	Portugal
Mimizan	44°08' N	1°18' E	France
Oria	37°31' N	2°21' E	Spain
Pineta	41°57' N	9°02'W	France
Pinia	42°01' N	9°27'W	France
Pleucadec	47°46' N	2°20' W	France
Puerto de vega	43°32' N	6°37' W	Spain
Rodoiros	43°25' N	6°32'W	Spain
San Cipriano de Ribarteme	42°07' N	8°21' W	Spain
San Leonardo	41°50' N	3°3'W	Spain
Sergude	42°49' N	8°27'W	Spain
Sierra de Barcia	43°31' N	6°29' W	Spain
Tamrabta	33°36' N	5°01'W	Morocco
Valdemaqueda	40°30' N	4°05' W	Spain

# APPENDIX II

Genetic differentiation for size at first reproduction through male versus female functions in the widespread Mediterranean tree *Pinus pinaster*.
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# Genetic differentiation for size at first reproduction through male versus female functions in the widespread Mediterranean tree *Pinus pinaster*

L. Santos-del-Blanco<sup>1,2,\*</sup>, J. Climent<sup>1,2</sup>, S. C. González-Martínez<sup>1,2</sup> and J. R. Pannell<sup>3</sup>

<sup>1</sup>INIA-CIFOR, Ctra A Coruña km 7.5, E-28040, Madrid, Spain, <sup>2</sup>Sustainable Forest Management Research Institute, University of Valladolid, Avda. Madrid 44, E-34071, Palencia, Spain and <sup>3</sup>Department of Ecology and Evolution, Biophore Building, University of Lausanne, CH-1015, Lausanne, Switzerland \*For correspondence. E-mail santos.luis@inia.es

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• *Background and Aims* The study of local adaptation in plant reproductive traits has received substantial attention in short-lived species, but studies conducted on forest trees are scarce. This lack of research on long-lived species represents an important gap in our knowledge, because inferences about selection on the reproduction and life history of short-lived species cannot necessarily be extrapolated to trees. This study considers whether the size for first reproduction is locally adapted across a broad geographical range of the Mediterranean conifer species *Pinus pinaster*. In particular, the study investigates whether this monoecious species varies genetically among populations in terms of whether individuals start to reproduce through their male function, their female function or both sexual functions simultaneously. Whether differences among populations could be attributed to local adaptation across a climatic gradient is then considered.

• *Methods* Male and female reproduction and growth were measured during early stages of sexual maturity of a *P. pinaster* common garden comprising 23 populations sampled across the species range. Generalized linear mixed models were used to assess genetic variability of early reproductive life-history traits. Environmental correlations with reproductive life-history traits were tested after controlling for neutral genetic structure provided by 12 nuclear simple sequence repeat markers.

• *Key Results* Trees tended to reproduce first through their male function, at a size (height) that varied little among source populations. The transition to female reproduction was slower, showed higher levels of variability and was negatively correlated with vegetative growth traits. Several female reproductive traits were correlated with a gradient of growth conditions, even after accounting for neutral genetic structure, with populations from more unfavourable sites tending to commence female reproduction at a lower individual size.

• *Conclusions* The study represents the first report of genetic variability among populations for differences in the threshold size for first reproduction between male and female sexual functions in a tree species. The relatively uniform size at which individuals begin reproducing through their male function probably represents the fact that pollen dispersal is also relatively invariant among sites. However, the genetic variability in the timing of female reproduction probably reflects environment-dependent costs of cone production. The results also suggest that early sex allocation in this species might evolve under constraints that do not apply to other conifers.

Key words: *Pinus pinaster*, conifers, sex-dependent threshold size for first reproduction, size-dependent sex allocation, clinal variation, neutral genetic structure.

#### INTRODUCTION

It is widely appreciated that plants are enormously variable in key life-history traits such as seed number, germination rate, growth rates and time to flowering. To the extent that life-history variation has an additive genetic component, these observations are puzzling, because natural selection is expected to deplete genetic variation for traits that affect individual fitness (reviewed by Barton and Keightley, 2002). Much of the variation in life-history traits within populations is attributable to phenotypic plasticity (Sultan, 2000), but it is also well established that plant populations contain large amounts of additive genetic variation for life-history traits, including seed size and number (Mazer, 1987), plant relative growth rate (Sánchez-Gómez *et al.*, 2010), time to flowering (Montague *et al.*, 2008), and flower size and number (Worley and Barrett, 2000). Understanding the nature and

maintenance of this variation remains an important challenge for plant evolutionary biologists. One idea is that life-history variation is rendered effectively neutral as a result of antagonistic pleiotropy, i.e. fitness trade-offs between different lifehistory traits (Walsh and Blows, 2009), or by fluctuating selection from among generations (Bonser and Aarssen, 2009; Childs *et al.*, 2010). Another idea is that much of the variation we observe is maintained in a balance between purifying selection, which depletes it, and its replenishment by mutation or immigration (Barton and Keightley, 2002).

The importance of immigration as a process that introduces new genetic variation for life-history traits into a population will ultimately depend on the maintenance of this variation among populations by adaptive responses to natural selection locally, i.e. on the adaptive divergence among populations for the life-history traits concerned. Measuring population genetic divergence for life-history traits is thus an important

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task, yet discriminating between variation due to phenotypic plasticity and that due to genetic divergence requires the measurement of traits expressed by individuals from different populations under comparable environmental conditions (i.e. tested in common gardens or reciprocal transplant experiments). In standard common-garden experiments, genotypes sampled from a range of different populations are grown together in a common environment. Unlike reciprocal transplant experiments, common gardens are unable to test the hypothesis of local adaptation for all populations, but they allow an assessment of genetic divergence among more populations that could be accommodated in fully reciprocal transplant experiments. Common garden experiments thus offer a promising avenue towards understanding the maintenance of life-history variation in geographically widespread species.

The age and/or size of first reproduction, i.e. the onset of reproduction, and the number of offspring produced represent particularly important life-history traits. Slight changes in size at maturity and fecundity can have dramatic effects on lifetime reproductive output in perennial species, because early reproduction can be costly in terms of future reproductive potential (Kozłowski, 1992; Roff, 2000). Theory predicts that selection will cause plants with a large life expectancy and a strongly positive relationship between size and fecundity to delay their onset of flowering (Roff, 1992). It is widely accepted that plants tend to use size rather than age as a developmental cue for reproduction (de Jong and Klinkhamer, 2005; but see Metcalf et al., 2003). However, size and age are clearly linked, not least because the age schedule of survival and the disturbance regime, which affects life expectancy, can have an important impact on size at reproduction (Klinkhamer and de Jong, 1987; Burd et al., 2006; Weiner et al., 2009).

Plant size at first reproduction has received relatively little attention in long-lived polycarpic trees (Thomas, 1996; Dodd and Silvertown, 2000; Niklas and Enquist, 2003; Wright *et al.*, 2005), although it is well studied in monocarps (Wesselingh *et al.*, 1997; Rees *et al.*, 1999; Callahan and Pigliucci, 2002; Kuss *et al.*, 2008; Kagaya *et al.*, 2009) and perennial herbs (Méndez and Karlsson, 2004; Brys *et al.*, 2011). These studies have not only documented substantial variability in size at first reproduction among populations, but they have also shown high within-population variability. Moreover, they confirmed theoretical predictions on the role of environmental effects that act through changes in disturbance regimes and growth conditions (see Méndez and Karlsson, 2004).

Although it may sometimes be useful to refer to size at first reproduction as a single trait, individuals in sexual populations transmit their genes through both male and female functions, and the size threshold for each may differ. Thus, in the case of dioecious species, males often begin flowering earlier than females (Delph, 1999). Similarly, in monoecious or hermaphroditic species, male function often precedes female function (Freeman *et al.*, 1981). In some species, this decoupling of male and female functions in relation to size, i.e. a sexdependent threshold size for first reproduction (TSFR), can result in the presence of individuals functioning as pure males, as females or as hermaphrodites expressing both sexes simultaneously during their early stages of sexual maturity (de Jong and Klinkhamer, 2005; Zhang, 2006). In the extreme, populations can express sequential hermaphroditism, as in the classical example of the jack-in-the-pulpit, where plants are male when small and either female or hermaphroditic one to several reproductive seasons later (Kinoshita, 1987). Such transitions in the functional gender of individuals as they grow have been interpreted in terms of adaptive responses to natural selection when the fitness gains through male and female functions depend on size (Cadet *et al.*, 2004; de Jong and Klinkhamer, 2005; Zhang, 2006).

Size effects on fitness expectations through male vs. female functions can be direct or indirect (Klinkhamer et al., 1997). In wind-pollinated species, pollen dispersal may be enhanced directly by growth in height, so that tall individuals preferentially express their male function (Pickup and Barrett, 2012). In animal-pollinated species, this 'direct effect' of height on male fitness is less likely. Here, larger plants are more likely to enhance their female function if the female fitness gain curve flattens off less quickly with investment than the male curve, as seems likely; these are the so-called 'budget effects' of size on sex allocation (Klinkhamer et al., 1997). Because a plant's resource base, and therefore its growth rate and ultimate size, will inevitably depend on habitat quality, we might then expect selection through these indirect budget effects to vary among populations of a species exposed to environments that differ across the species' geographical range. For example, the relative costs of male versus female functions may be different under different environments, such that female function may be borne optimally by large individuals in some, but not all, populations (see Klinkhamer et al., 1997). If so, differences in sex-specific TSFR may evolve in accordance with the differing trade-offs between reproduction through each sexual function and growth. Such geographical variation in this important life-history trait would allow a more subtle understanding of the action of natural selection on sex allocation that is possible through broader comparisons among species, for example with different pollination modes. There has, however, been very little research on among-population variation in TSFR through male versus female functions (hereafter male or female TSFR) despite the abundant recent literature dealing with sex allocation in plants.

Forest trees offer a valuable opportunity to test hypotheses concerning variation in size-dependent sex expression and sex allocation (SDS). Not only do they often have large distribution ranges (Petit and Hampe, 2006), but the several-fold differences in size between small and large plants within populations has probably favoured the selection of SDS patterns (Burd and Allen, 1988). For instance, variability in the gender of monoecious individuals in conifers does seem to be associated with variability in size related to environmental stress (Cobb *et al.*, 2002; Kang, 2007). Interestingly, this is in some ways similar to the well-established observation of an increased tendency towards dioecy with environmental stress in species that vary in the degree to which their sexes are separate (Ashman, 2006; and see Discussion).

In this study, we use a large common garden to assess patterns of genetic variation in early reproductive life-history traits among populations of the widespread Mediterranean species *Pinus pinaster* (maritime pine). *P. pinaster* is a

monoecious, wind-pollinated, wind-dispersed conifer with a remarkably wide ecological niche (Tapias et al., 2004). Previous work, based on data from common-garden experiments, has described among-population variation in size at reproduction for this species (Santos-del-Blanco et al., 2010). Here, we investigate whether P. pinaster shows differences in the threshold size at first reproduction through male versus female function among populations from a broad geographical range across a strong climatic gradient, from Atlantic (i.e. mild winters, high annual precipitation and low altitude) to dry continental environments. We sought evidence for a genetic trade-off between allocation to growth and early reproduction, and we tested the prediction that the median threshold size at first reproduction through each of the two sexual functions should correlate with the environment of the source populations. Finally, by comparing the amongpopulation genetic structure in terms of life-history traits with that displayed by neutral genetic markers, we tested the hypothesis that observed clinal variation in the former has resulted from responses to natural selection as opposed to drift or historical effects (Alleaume-Benharira et al., 2006; Grivet et al., 2011).

#### MATERIALS AND METHODS

#### Common garden

We established our common garden at Parderrubias, Ourense Province, north-west Spain (Fig. 1A). This common garden belongs to a broader series of experiments aimed at describing variability in adaptive traits in *Pinus pinaster*, and was chosen for the present study as it was placed in a transitional area from Atlantic to Mediterranean climate, as indicated by Gower's distance (Fig. 1B). The common garden comprised 194 openpollinated families from a total of 23 natural populations distributed over the greater part of the species' natural range, including Atlantic Iberian Peninsula and France, Mediterranean Spain, Corsica and Morocco (Fig. 1, Supplementary Data Table S1). Two additional non-natural populations present in the common garden were excluded from the present study.

Half-sib families in natural populations were sampled from trees that were spaced at least 100 m apart. In total, 2767 2-year-old seedlings were established from seed following standard nursery procedures, and were planted in the common garden in 2005 in a resolvable alpha design, with four replicates of 71 incomplete blocks and four contiguous plants from the same family per block. This design was originally chosen because it allows efficient control of spatial heterogeneity, which is a common problem in large common gardens of forest trees. Additional seedlings from a local provenance were used as a border as well as to fill in the gaps of missing plants. Individuals were planted at a spacing of  $2 \text{ m} \times 3 \text{ m}$ . The soil in the common garden was a humic cambisol. Despite generally uniform climatic conditions across the common garden, there was inevitably small-scale environmental heterogeneity due to a soil depth gradient, and this was probably responsible for some of the (random) variation in plant size. Of the 2767 trees initially planted, 2240 were still alive when we last measured them in 2010.

#### Measurement of traits and environmental variables

We measured reproductive and growth-related traits in 2007, 2009 and 2010, a period that covered the transition from a point where most individuals were still vegetative to one when most had become reproductive. We also calculated a number of derived size-dependent fitness components (Méndez and Karlsson, 2004), as set out in Table 1. Correlation among size-dependent fitness components was tested using Pearson's correlation coefficient. Population means of size-dependent fitness components were also included in association tests with environmental data (see below). Following previous research (Climent et al., 2008; Grivet et al., 2011), three spatial and six climatic variables were considered (Table 1). Climatic data for Iberian populations were obtained from a functional phytoclimatic model based on raw data from meteorological stations (Gonzalo-Jiménez, 2010). Climatic data for non-Iberian populations were obtained from WorldClim-Global Climate Data at 5' resolution (Hijmans et al., 2005).

#### Molecular analysis

Needles from between six and 30 individuals per population (mean = 16) were collected and dried in silica gel for subsequent DNA extraction. Twelve nuclear microsatellites (simple sequence repeats, SSRs) were genotyped: ITPH4516 (Mariette et al., 2001); RPtest1, ctg275, ctg4363, NZPR1078 and NZPR544 (Chagné et al., 2004); A6F03 (Guevara et al., 2005); pEST2669 (Steinitz et al., 2011); and gPp14, epi3, epi5 and epi6 (F. Sebastiani and G. G. Vendramin, Istituto di Genetica Vegetale, CNR, Florence, pers. comm., June 2011; Supplementary Data Table S2). Forward primers were 5' endlabelled with fluorochromes (HEX, FAM, VIC or PET) and amplified using the Qiagen Multiplex PCR Kit (Qiagen, Venlo, the Netherlands) following the manufacturer's instructions. Amplified allele fragments were separated using an ABI 3730 genetic analyser (Applied Biosystems, Carlsbad, CA, USA) and fragment sizes were determined with reference to the GeneScan<sup>TM</sup> –500 LIZ<sup>®</sup> Size Standard (Applied Biosystems) using GeneMapper software version 4.0 (Applied Biosystems).

#### Data analysis

Genetic variation among populations for flowering probability. Based on the common-garden data, individual male and female reproductive status was coded as a binary variable at each sampling time for each individual (4, 6 and 7 years old). Given the 2-year developmental cycle of female strobili, female reproduction was recorded as present for any tree bearing 1st- or 2nd-year female cones.

The probability of reproduction was modelled by a generalized linear mixed model (GLMM) with a binomial family distribution and logit link function fitted by Laplace approximation, as implemented by the lme4 package (Bates *et al.*, 2011) on the R platform (R Development Core Team, 2012). Total height (i.e. size), provenance and their interaction were used as fixed terms, and family was included as a random factor. Block structure was not included in these models, as preliminary analyses showed no influence of spatial heterogeneity



FIG. 1. Distribution map of *Pinus pinaster* source populations (circles) and common garden (triangle). Shaded areas indicate the species' natural distribution range; see key in (B) for population abbreviations. (B) Gower's (absolute) environmental distance between each population to the common garden location. Values closer to 0 indicate greater similarity between the population of origin and the common garden site with respect to the environmental variables used for the calculation of the distance. Symbols represent genetic groups, as indicated in the key.

on the allometry of reproduction (i.e. the block effect or spatial auto-correlation disappeared when tree size variation was included in the models). This first model was used to test the existence of genetic variation among populations for the relationship between size (estimated by height) and reproduction, both by using a likelihood ratio test (LRT) for models with and without the interaction term, and in terms of overall Akaike information criterion.

Ontogenetic and climatic effects on reproduction were necessarily confounded in our experiment, as all trees were the same age. Because we did not include age/year information in the model, our results should thus be interpreted as reflecting the overall behaviour of trees during their transition to maturity, but also to some extent influenced by among-year climatic variation.

We fitted a GLMM for both male and female reproduction using Markov chain Monte Carlo (MCMC) methods, with uninformative priors and fixing residual variance at 1 (MCMCglmm R package; Hadfield, 2011). This approach was preferred over restricted maximum-likelihood methods (Hadfield, 2010), which did not achieve satisfactory convergence for some populations when random effects were included, and also because MCMC methods allowed a more accurate estimation of errors in estimates of the median threshold size for reproduction. The Markov chain was run for 5500 000 iterations, sampling each 5000 after a burn-in period of 500,000 iterations. These settings proved suitable. as successive values of the chain were uncorrelated. Convergence of the chain was also checked graphically. From these models, posterior mode values for the parameters  $\mu$  (intercept) and  $\alpha$  (slope associated with height) were extracted as size-dependent fitness components commonly used in these studies (Méndez and Karlsson, 2004; Brys et al., 2011).

Age-dependent size distribution of reproductive stages. We classified each individual tree as female, male, cosexual or juvenile depending on the sex expressed for its first reproductive event; juveniles were those individuals that had still not started reproducing in 2010. We also recorded the height of individuals at which the first reproductive event took place. We tested whether plant size for each class of reproductive status differed significantly overall and among populations by using the mixed model  $H_{ijk} = \mu + R_i * P_j + F_k + e_{ijk}$ , where *H* was plant height,  $\mu$  was the general mean, *R* was the reproductive class (female, male, cosexual or juvenile), *P* was the population (fixed factor), *F* was the family (random factor) and *e* was the random error term. The GLMM was implemented in R. Finally, we recorded the changes in sex choice for all individuals across the last two years (Fig. 2).

Genetic population structure and environmental correlations. Maritime pine is characterized by strong population genetic structure at the range-wide scale (e.g. Bucci et al., 2007). When testing for environmental correlations at wide spatial scales, geographical genetic structure in maritime pine results in large numbers of false positives (estimated at approx. 84 % by Grivet et al., 2011). Thus, here we used linear models to test for genetic associations between early reproductive life-history traits and environmental data that controlled for neutral population structure (i.e. associations between plant traits and environmental data that could be due to demographical history) using molecular markers (nuclear SSRs) as covariates. Neutral population structure was assessed using the Bayesian clustering method implemented in STRUCTURE 2.2 (Pritchard et al., 2000). We ran an admixture model with correlated allele frequencies between clusters. Ten runs were performed for each number of clusters from K = 1 to K = 10, with a burn-in length of 50 000 and a

TABLE 1. Common gai	rden measured traits	, derived size-depender	t fitness	components	and nativ	e location	environmental	data of	
		Pinus pina	ster trees	1					

Variable		Description
Measured traits	ŝ	
Н	Height (cm)	Total height of each tree in 2007, 2009 and 2010
F. Rep	Female reproduction	Presence or absence of seed cones in each tree, yearly from 2007
M.Rep	Male reproduction	Presence or absence of pollen cones in each tree in 2009 and 2010
Derived size-de	ependent fitness components	
μ		Intercept of reproduction-size logistic regression at the population leve
α		Slope of reproduction-size logistic regression at the population level
TSFR	Median threshold size for first reproduction (cm)	Size at which the probability for a tree within a population to have reached sexual maturity was 50 %, equal to $\mu/\alpha$
SRI	Smallest reproductive individual (cm)	Size of the smallest reproductive individual in the population
LVI	Largest vegetative individual (cm)	Size of the largest vegetative individual in the population
RAN		Range between LVI and SRI
%VEG	Percentage of vegetative trees (cm)	Percentage of non-reproductive trees larger than the population TSFR
%REP	Percentage of reproductive trees (cm)	Percentage of reproductive trees in the population
Environmental	data	
Alt	Altitude (m)	
Long	Longitude (m)	
Lat	Latitude (m)	
AMT	Annual mean temperature (°C)	
MTWM	Mean temperature of the warmest month (°C)	
MTCM	Mean temperature of the coldest month (°C)	
CI	Continentality index (°C)	Difference between mean temperature of the warmest month and mean temperature of the coldest month
AP	Annual precipitation (mm)	-
PDM	Precipitation during the warmest quarter (mm)	Precipitation in June, July and August

run length of 500 000 iterations. The optimal number of population clusters, K, was calculated following Evanno et al. (2005) and the guidelines provided by STRUCTURE developers in the manual. Population means for individual assignment probability (Fig. 3) to each of the K clusters (i.e. the Q matrix, Yu et al., 2006) were used to control for neutral population structure in environmental associations (see similar appoaches in Eckert et al., 2010; Grivet et al., 2011). Additionally, to allow a more synthetic interpretation, we carried out a principal component analysis (PCA) with varimax rotation to reduce the number of environmental variables. To decide the number of principal components to retain, we ran a parallel analysis with 1000 iterations (Hayton et al., 2004) and selected only those principal components with eigenvalues for observed data larger than those obtained in simulations. The analysis was implemented by the psych package (Revelle, 2011) on the R platform (R Development Core Team, 2012). PCA loadings of retained components were used in correlation analysis with plant traits. Finally, we used a likelihood ratio test to compare a model including only neutral population genetic structure with one including both neutral population genetic structure and PCA loadings or single environmental variables to explain phenotypic differences among populations.

#### RESULTS

#### Phenotypic variation in early sex expression

Trees began to reproduce sexually from 5 years of age, i.e. in 2008, and most were reproductive by year 7 (in 2010). The majority of these early individuals expressed only their male

function, with similar lower numbers expressing either only female or both sexual functions (Table 2). In 2009, about half of the plants were juvenile, some remained juvenile in 2010, many began to express their male function and a minority expressed their female function, either purely or together with their male function. Individuals bearing only female or male cones in 2009 tended to produce only male cones or both male and female cones in 2010, so that individuals producing only female cones over two consecutive years were rare. Finally, all trees with female and male cones in 2009 continued producing male cones, but some abandoned their female function (Fig. 2).

Overall, juveniles were the smallest class of individual  $(206.0 \pm 23.9 \text{ cm})$ , followed by individuals expressing their female function  $(233.2 \pm 23.6 \text{ cm})$ , then those expressing their male function  $(321.1 \pm 23.3 \text{ cm})$  and finally those expressing both functions  $(372.9 \pm 25.2 \text{ cm})$ . However, this association between gender expression and plant size also depended significantly on the population of origin (population × gender interaction, P < 0.0001).

#### Associations between reproduction and size

Median TSFR varied significantly among populations for both male (P < 0.0001) and female functions (P < 0.0001). At the population level, the size effect was highly significant for all populations except the Pineta population (Corsica), which was dropped in subsequent analyses. Across all other populations, intercepts were similar for male and female logistic regressions, while slopes where approximately double for male compared with female reproduction, reflecting the faster transition from juvenile to male reproductive stage. Median male TSFR was relatively uniform among populations and lower than female TSFR, which varied substantially among populations (Fig. 4).

Average height of the largest vegetative individual (LVI) was sensibly larger for plants considering female function, whereas heights were only slightly smaller for female function compared with male function regarding the smallest reproductive individual (SRI). Accordingly, size ranges were larger for female function (Table 3). Size-dependent fitness components at the population level are given in Supplementary Data Tables S3 and S4.

## Correlations and trade-offs between size-dependent fitness components

There was low variability for male function, and correlations between early male reproductive traits with other traits were non-significant (data not shown). In contrast, significant correlations were found between female fitness components, height



FIG. 2. Transitions between reproductive developmental stages across *Pinus pinaster* populations grown in a common garden in north-west Spain. The *x*-axis represents morph classes at age 6, and within each class bars represent the number of individuals behaving as juveniles (J), females (F), males (M) or cosexuals (C) at age 7.

and the percentage of individuals reaching sexual maturity through female function (Table 4). Height was positively correlated with female TSFR (i.e. delayed female reproduction) and negatively correlated with the number of individuals that became reproductive through female function, thus indicating an overall negative correlation between growth in height and female function at the population level.

#### Environmental correlations

Parallel analysis associated with PCA for environmental traits revealed that only one principal component (PC1),

TABLE 2. Percentages of Pinus pinaster trees remaining non-reproductive (J), or having reached sexual maturity as females (F), males (M) or cosexuals (C) until year 7

Population	GG	% J	% F	% M	% C	n
Mimi	AF	15	13	43	30	96
Pleu	AF	12	36	20	31	108
Alto	AIP	9	12	68	11	65
Arma	AIP	14	8	58	20	71
Cada	AIP	14	3	76	7	87
Cast	AIP	15	15	50	20	74
Lamu	AIP	15	12	55	18	67
Leir	AIP	15	12	61	12	132
Puer	AIP	16	3	66	15	61
Sanc	AIP	10	31	33	26	39
Segu	AIP	7	13	55	24	165
Sier	AIP	11	7	72	11	46
Pine	CO	59	11	30	0	27
Pini	CO	11	6	70	13	54
Oria	MO	20	27	40	13	143
Tamr	MO	43	32	20	5	125
Aren	MS	12	16	48	24	126
Bayu	MS	20	35	31	13	143
Carb	MS	15	35	27	23	48
Ceni	MS	5	16	47	31	55
Coca	MS	32	25	41	1	68
Cuel	MS	14	44	30	12	153
Rodo	MS	17	16	46	21	76
SanL	MS	18	22	40	20	137
Vald	MS	14	18	54	15	74
Average or to	tal	16	19	48	17	2240

*N*, number of trees per population. GG, genetic group; see Fig. 1 for GG codes and Supplementary Data Table S1 for population codes.



F1G. 3. Barplot of individual assignment probability to each of the optimal K = 6 clusters representing five geographical genetic groups in *Pinus pinaster* as produced by STRUCTURE 2.2 software. Each individual is represented as a line segment which is vertically partitioned into *K*-coloured components, representing the individual's estimated proportions of ancestry in the *K* clusters. Population abbreviations are given in Supplementary Data Table S1.



FIG. 4. Median threshold size for first reproduction through male and female functions (as indicated) in *Pinus pinaster* populations grown in the common garden in north-west Spain. Bars represent the posterior mode of Bayesian estimates, with lower and upper 95 % credible intervals. Population abbreviations are given in Supplementary Data Table S1.

explaining 64 % of the variance, should be retained. Overall, this principal component reflected favourable growth conditions. This factor was positively related to longitude, mean temperature of the coldest month (MTCM), precipitation during the warmest quarter (PDM) and annual precipitation (AP), and negatively to altitude, mean temperature of the warmest month (MTWM) and continentality index (CI). Latitude and annual mean temperature (AMT) had loadings below 0.7.

Male TSFR was not correlated with PC1 or any single spatial or climatic variable (Table 5 and Supplementary Data Table S5), but we found significant positive associations between PC1 and female TSFR, height and the percentage of individuals expressing their male function, and negative associations between PC1 and the percentage of individuals expressing their female function. Among single climatic parameters, models for MTCM and CI showed the highest significance levels (see Supplementary Data Table S5). Trees with larger female TFSR tended to come from sites with more favourable growing conditions, as those with milder winters and a reduced degree of continentality (Fig. 5).

Nuclear SSR summary information is provided in Supplementary Data Table S2. A STRUCTURE analysis using Bayesian clustering based on SSRs showed the existence of five clearly differentiated genetic groups in P. pinaster corresponding roughly to Atlantic France, Atlantic Iberian Peninsula, Corsica, Mediterranean Spain (except the Oria population in southern Spain) and Morocco plus Oria (Fig. 3 and Supplementary Data Table S6). Atlantic and Mediterranean Iberian populations displayed varying degrees of admixture, which has been suggested to be the result of historical gene flow among Mediterranean and Atlantic maritime pine glacial refugia (de Lucas et al., 2009). Once neutral population genetic structure was integrated in the models, the overall correlations of early reproductive life-history traits with favourable growth conditions (as resumed in PC1) remained significant. However, several correlations for single environmental parameters became non-significant, including one for CI (see Supplementary Data Table S5). This indicated that either some true environmental associations were confounded

TABLE 3. Mean values for size-dependent fitness components for male and female reproduction, averaged across 22 natural Pinus pinaster populations; abbreviations are as defined in Table 1

	Male	Female		
μ	$-8.07 \pm 1.79$	$-5.75 \pm 0.90$		
α	$0.033 \pm 0.007$	$0.018 \pm 0.005$		
TSFR	$241.7 \pm 19.7$	$339.1 \pm 81.9$		
LVI	322 + 50	393 + 39		
SRI	113 + 17	105 + 22		
RAN	209 + 52	254 + 100		
%VEG	4.0 + 2.4	1.7 + 2.3		

with neutral genetic structure imposed by demographic history (and thus were not reliable), or that they were false-positives.

#### DISCUSSION

Our results provide evidence for clear genetic differences in key life-history traits among populations of *P. pinaster* sampled across a wide environmental range. Because much of this genetic variation was associated with a strong environmental gradient among the sites we sampled, some of which remained significant after neutral genetic structure was accounted for, the observed life-history trait clines are likely to be the result of local adaptation rather than simply the outcome of divergence through drift or historical accident. This is among the first reports of reproductive life-history trait variation in a widespread forest species (Climent *et al.*, 2008; Santos-del-Blanco *et al.*, 2010) but also, as far as we know, the first report of adaptive genetic differentiation for both male and female threshold sizes for reproduction in a (monoecious) plant species.

The *P. pinaster* individuals in our study tended to start reproducing preferentially as males, and they tended to maintain their male function fairly constantly through time. Accordingly, male median TSFR was lower than female TSFR. This finding is somewhat surprising, given that most

TABLE 4. Among-population	correlations for	early female	reproductive	traits in	Pinus	pinaster;	abbreviations	are	as	defined in	п
			Table 1.								

	fμ	fα	fTSFR	fLVI	fSRI	fRAN	%fVEG	%J	%F
Н	-0.305	-0.360	0.691***	0.654***	0.744***	0.269	-0.363	-0.634**	-0.747***
fμ		-0.523*	-0.148	-0.505*	-0.553 **	-0.222	0.117	-0.036	0.315
fα			-0.730***	-0.161	-0.094	-0.128	0.521*	0.152	0.517*
fTSFR				0.624**	0.544**	0.372	-0.694 ***	-0.231	-0.865***
fLVI					0.557**	0.815	-0.337	-0.149	-0.579 **
fSRI						-0.026	-0.324	-0.420	-0.689***
fRAN							-0.180	0.113	-0.217
%fVEG								0.284	0.768***
%J									0.322

Significant at: \*\*\**P* < 0.001, \*\**P* < 0.01, \**P* < 0.05.

conifer species tend to start reproducing as females (Williams, 2009). In perennial plants and, most notably, in trees, the existence of threshold sizes for reproduction arises from the need to invest all available resources in a large vegetative body during early developmental stages to maximize further reproduction (reviewed by Thomas, 2011). The existence of greater female costs of reproduction has been related to a generally lower threshold size for male reproduction in dioecious and monoecious plants and to failures in setting fruit in species with perfect flowers (Iwasa, 1991; de Jong and Klinkhamer, 2005). This pattern, largely observed in woody angiosperms, differs from that typically observed in conifers, which display earlier expression of female function. P. pinaster would thus seem to have evolved a pattern of early reproduction and sex allocation more similar to woody angiosperms than other conifers.

Although the trend for early male reproduction in *P. pinaster* is interesting and unusual, the more striking pattern we observed is that male TSFR is much more canalized, both within and among populations, than female TSFR: individuals in our common garden started producing male cones at around 2.5 m in height, whereas female function commenced in individuals over a very wide range of heights (Fig. 4). Why should male TSFR be so uniform, and why should the female TSFR vary so widely?

An increase in maleness with size in wind-pollinated plants is a well-established prediction from SDS theory (de Jong and Klinkhamer, 2005), and forest trees have been identified as suitable organisms to test it (Cruden and Lyon, 1985). The prediction rests on the presumed advantage to plants that release their pollen from greater heights, regardless of the seed dispersal mode (Friedman and Barrett, 2009). However, in conicalshaped conifers, like many pine species, male cones are not located in the treetop but rather on lower secondary branches, and it is the female cones that are borne by upper vigorous branches. This segregation is more evident in adult trees (Shmida et al., 2000), but we have observed it also in our young maritime pine specimens. This architectural pattern has been proposed either as a strategy to avoid selfing, as male and female flowering are synchronous in this species (Miguel-Pérez et al., 2002), and/or as a consequence of the larger size of female cones which could only be borne on stronger upper vertical branches (Ne'eman et al., 2011). Either way, the hypothesis that increased height favours male

TABLE 5. Environmental associations of Pinus pinasterreproductive life-history traits (first column) with environmentalvariables represented by population scorings on first componentof PCA analysis

Trait	Corrected		Uncorrected	
	Slope	Р	Slope	Р
fTSFR	$109.10 \pm 26.97$	0.001	$56.34 \pm 13.30$	0.000
mTSFR	$8.36 \pm 8.90$	n.s.	$-1.89 \pm 4.39$	n.s.
Н	$32.21 \pm 11.80$	0.016	$24.30 \pm 4.16$	0.000
%M	$15.28 \pm 6.15$	0.025	$10.43 \pm 2.97$	0.002
%F	$-12.34 \pm 4.85$	0.023	$-7.74 \pm 2.04$	0.001

Corrected values indicate slopes, standard errors and *P*-values for the association after including neutral genetic structure corrections (i.e. likelihood ratio test between a full model and a reduced model with just neutral genetic structure); uncorrected values indicate association parameters without correcting for neutral genetic structure (i.e. just environmental data and traits data in the model). n.s., not significant.



FIG. 5. Variation of *Pinus pinaster* median female threshold size for first reproduction along an environmental harshness gradient, as estimated by population scorings on PC1. Symbols represent genetic groups, as indicated in the key.

function would not offer a convincing explanation for sizedependent sex allocation in conifers (Fox, 1993). Moreover, the size advantage hypothesis only applies in the presence of clear differences in size among individuals within a population (Friedman and Barrett, 2009), which is not common in Mediterranean conifer forests. The established theory for sizedependent sex allocation thus would not seem to apply to conifers in a straightforward manner.

Given that the position of male and female cones is conserved across conical-shaped conifers and thus appears to be phylogenetically constrained, for whatever reason, a more plausible explanation for canalization of male TSFR is simply that pollen needs to be released above a threshold height to be lifted by horizontal winds and updrafts; if so, it would not be surprising if the same physical constraint applied to all individuals, irrespective of their provenance. The high value for the slope of the male flowering probability curve with size ( $\alpha = 0.033$ ) indicates low genetic variation for this trait and a common sharp threshold for male TSFR within and across populations. We therefore hypothesize that the low among-population variation in the threshold sizes for male reproduction found in our study points to the existence of similar male fitness gain curves and might be the result of rather uniform conditions for pollen release and transport across populations. To our knowledge, this possibility has not yet been considered in the SDS literature.

Female TSFR not only showed high variability within populations, but also clear evidence for high amongpopulation differentiation; this variation, in turn, was significantly correlated with most early reproductive life-history traits (Table 4). For example, we found a positive correlation among populations between the average height of individuals and female TSFR, and a negative correlation between the average height in the population and the number of individuals first reproducing as females (as opposed to early males). These patterns emerge as negative correlations of population means for reproductive and growth traits measured in a common garden environment, so our results should be considered at the genetic, not phenotypic or physiological, level (Reznick, 1985; de Jong and Klinkhamer, 2005). Thus, the observed patterns probably reflect genetic trade-offs between growth and female reproduction that have been selected under different environments.

A particularly interesting finding is that much of the observed among-population variation was associated with climatic differences among sites. At least some of this variation might underlie possible differences in the intensity of withinstand competition (e.g. competition might be more important at more mesic sites; Grime, 1977; Grivet *et al.*, 2011). Thus, in favourable (yet strongly competitive) environments, early investment in the more costly female function would increase the risk to individuals of being suppressed by their neighbours, so a delay of reproduction might be advantageous (Thomas, 2011). By contrast, in unfavourable (but less competitive) environments, selection would tend to favour stress tolerance at the expense of growth or competitive ability, so that a low female TSFR would be expected (Roff, 1992).

In accordance with this reasoning, we found lower female TSFR and a higher proportion of individuals first expressing their female function in populations from sites offering less favourable conditions for growth, a pattern that remained significant after accounting for neutral genetic structure. It would thus seem plausible that, on the one hand, populations have become differentiated for female TSFR in response to selection under possible differences in the competitive environment and disturbance regime among sites, and, on the other hand, a canalized male TSFR has evolved under the possible site-independent physical constraints facing pollen dispersal in conifers.

The idea that selection will have caused populations of P. pinaster to diverge for reproductive traits across its range is largely in keeping with observed clinal variation in morphological traits for a number of widespread species (Davis et al., 2005); in some cases, this clinal variation has evolved over short time periods, for example flowering-time variation in Verbascum thapsus (Ansari and Daehler, 2010) and Lythrum salicaria (Montague et al., 2008; Colautti et al., 2010). Similarly, artificial selection experiments on size at reproduction in Cynoglossum officinale also elicited a fast and direct response (Wesselingh and de Jong, 1995), and the cultivation of Eucalyptus trees in India resulted in the rapid evolution of fecundity traits (Varghese et al., 2009). These studies argue for the existence of large genetic variation for size at reproduction, allowing fast among-population differentiation due to strong selection.

If the above explanation is able to account for amongpopulation patterns in TSFR, how might we explain the differences in within-population variability in this important trait between male and female functions? A differential plastic response in female TSFR of populations grown out of their native habitat could account for variability registered in the common garden. This variability might not be present in native habitats, where a more uniform female TSFR could exist. However, we found no evidence of correlation between environmental distance and range of median threshold sizes in each population ( $t_{21} = 0.761, P > 0.45$ ). Another possibility is that the greater variance in TSFR in female function within populations is attributable to the evolution of a bet-hedging strategy, either through stochastic or plastic expression of the same underlying genotypes, or through the frequencydependent maintenance of genetic variation for TSFR under environmental stochasticity (Rees et al., 2004; Metcalf et al., 2008; Weiner et al., 2009; Childs et al., 2010). The role of environmental stochasticity on reproductive strategies has been described mainly for monocarps, which can suffer high rates of mortality before reaching maturity. Woody species usually have more stable demographic patterns than monocarps, probably buffered by ongoing trade-offs between investment in reproduction and growth or maintenance. Nevertheless, Mediterranean pine species suffer unpredictable fire return intervals as well as episodic droughts, both of which are likely to play a major role in reproductive strategies (Pausas et al., 2008). A bet-hedging strategy in a long-lived woody species prone to environmental stochasticity might be expected to evolve for the more expensive reproduction function, i.e. for cone production, as observed. Interestingly, previous studies on Pinus spp. report lower heritabilities for male compared with female cone production, for example in P. pinaster (Merlo and Fernández López, 2004) and P. sylvestris (Savolainen et al., 1993), suggesting that a similar explanation Page 10 of 12

might apply to several species evolving under a rather broad range of environmental challenges.

#### SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjourals.org and consist of the following. Table S1. List and location of the *Pinus pinaster* populations comprising the present study. Table S2. Summary information of nuclear SSR markers used to estimate range-wide population genetic structure in *Pinus pinaster*. Table S3. Size-dependent fitness components for male reproduction in 23 natural *Pinus pinaster* populations. Table S4. Size-dependent fitness components for female reproduction in 23 natural *Pinus pinaster* populations. Table S5. Associations of environmental variables with reproductive life-history traits in *Pinus pinaster*. Table S6. Number of trees per population included in the molecular marker analysis and average individual assignment probability for each of the optimal K = 6 clusters representing five geographical genetic groups in *Pinus pinaster*.

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# SUPPLEMENTARY DATA

TABLE S1. List and location of the *Pinus pinaster* populations comprised in the present study. GG= genetic group (as identified by STRUCTURE; see main text); MTCM = mean temperature of the coldest month (°C); MTWM = mean temperature of the warmest month (°C); AMT= annual mean temperature (°C); CI = continentality index (°C); PWM = precipitation during the warmest quarter (mm); AP = annual mean precipitation (mm).

Population and abbreviation		Latitude	Longitude	GG	Altitude	MTCM	MTWM	AMT	CI	PWM	AP
Mimizan	Mimi	44°80'N	1°18'W	AF	37	3.2	24.8	13.3	21.6	232	1235
Pleucadec	Pleu	47°46'N	2°20'W	AF	80	2.5	21.9	11.2	19.4	154	804
Alto de la Llama	Alto	43°17'N	6°29'W	AIP	503	2.6	23.4	11.7	20.8	149	1137
Armayán	Arma	43°18'N	6°27'W	AIP	498	2.0	24.0	11.8	22.0	152	1112
Cadavedo	Cada	43°32'N	6°25'W	AIP	210	5.0	22.0	13.2	17.0	204	1316
Castropol	Cast	43°30'N	6°58'W	AIP	391	4.5	22.0	12.6	17.5	184	1179
Lamuño	Lamu	43°33'N	6°13'W	AIP	125	5.3	22.7	13.4	17.4	192	1282
Leiria	Leir	39°47'N	8°57'W	AIP	20	7.4	24.4	15.4	17.0	44	811
Puerto de Vega	Puer	43°32'N	6°37'W	AIP	121	4.9	22.6	13.4	17.7	194	1283
San Cipriano de Ribarteme	San	42°70'N	8°21'W	AIP	300	2.7	26.0	12.3	23.3	121	1600
Sierra de Barcia	Sier	43°31'N	6°29'W	AIP	240	4.7	22.4	13.0	17.7	192	1339
Pineta	Pine	41°58'N	9°20'E	CO	750	6.7	26.0	15.5	19.3	42	583
Pinia	Pini	42°10'N	9°27'E	CO	10	6.3	26.9	15.6	20.6	49	580
Oria	Oria	37°31'N	2°21'W	MO	1223	0.4	30.7	13.1	30.3	29	357
Tamrabta	Tamr	33°36'N	5°10'W	MO	1758	-4.6	30.4	10.7	35.0.	49	745
Arenas de San Pedro	Aren	40°11'N	5°60'W	MS	733	1.2	33.4	14.2	32.2	73	1318
Bayubas de Abajo	Bayu	41°31'N	2°52'W	MS	998	-1.4	29.6	10.6	30.9	96	553
Carbonero	Carb	41°10'N	4°16'W	MS	845	-0.7	31.3	12.3	32.0	72	435
Cenicientos	Ceni	40°16'N	4°29'W	MS	1100	1.3	28.8	12.4	27.5	60	794
Coca	Coca	41°15'N	4°29'W	MS	800	-0.6	31.2	12.3	31.8	77	454
Cuellar	Cuel	41°22'N	4°29'W	MS	830	-0.6	30.9	12.2	31.6	72	468
San Leonardo	SanL	41°50'N	3°30'W	MS	1096	-2.7	27.9	9.3	30.6	120	753
Valdemaqueda	Vald	40°30'N	4°18'W	MS	890	0.5	29.2	12.1	28.7	70	681

TABLE S2. Summary information of nuSSR markers used to estimate range-wide population genetic structure in *Pinus pinaster*. Name = locus name; Motif = SSR repeated motif; Label = fluorescent label; Min, Max = minimun and maximun allele sizes per locus; N = number of alleles per locus.

Name	Motif	Label	Forward sequence	Reverse sequence	Min	Max	N
A6F03	(AC)17	VIC	CCTGAAAATCGACGGATCG	ATGGTATTTTGCGGGTTGC	258	272	8
rptest1	(ATC)7	NED	AGGATGCCTATGATATGCGC	AACCATAACAAAAGCGGTCG	131	171	18
Ctg4363	(AT)10	VIC	TAATAATTCAAGCCACCCCG	AGCAGGCTAATAACAACACGC	108	124	8
NZPR1078	(AC)10	PET	TGGTGATCAAGCCTTTTTCC	GTTGATGAGTGATGGCATGG	239	259	10
epi3	(TC)15	NED	AGCAACATTTCCCTGGACAC	GGAATAATTGCAGTTGCAGTAGC	202	216	7
gPp14	(ATT)9	VIC	TATTGACGGTGTCTCTTCCT	GACTTTGACCTAAAGCATGG	215	237	8
pEST2669	(TA)19	NED	ATTGCTTCTGAAAGGGCATC	TCCCTTGGCACCATGTTAAT	202	229	6
epi5	(TA)9	PET	GGCGCGAACTACTTCATCTG	CAATGCTGACAAACCCAGAA	166	202	16
NZPR544	(CA)5(AC)12(TA)5	FAM	GCGATGTGCAACCCTTGATA	TGCTATTCCGTCAAAAACCC	340	359	8
ctg 275	(AT)16	FAM	ACGGAGATATATTGCTGGCG	AAAGAATAACGTGAAAACAAACCC	259	269	6
epi6	(AT)9	FAM	CCCACCATGACAAGGTTGAT	CGCTGGGCTTGAACATCTA	161	185	11
ITPH4516	(CT)27	PET	TGATGCAAACAAGTTCCATG	AGCACTCGCTAAACTATGAAGG	223	238	6

TABLE S3. Size- dependent fitness components for male reproduction in 23 natural *Pinus pinaster* populations. GG = genetic Group;  $m\mu$  = intercept of logistic regressions;  $m\alpha$  = slope of logistic regressions; mTSFR = median threshold size for male reproduction; loCI,upCI = lower and upper 95% limits of Bayesian credible intervals for mTSFR; LVI = largest vegetative individual respect to male reproduction; SRI = smallest individual bearing male cones; mRAN = transition range (mLVI–mSRI); mVEG = number of vegetative individuals. na = not available.

Рор	GG	$m\mu \pm s.e.$	$m\alpha \pm s.e.$	mTSFR	loCI	upCI	mLVI	mSRI	mRAN	%mVEG
Mimi	AF	$-10.72 \pm 5.91 \times 10^{-2}$	$4.15 \times 10^{-2} \pm 2 \times 10^{-4}$	258.2	220.1	291.1	378	146	232	4.2
Pleu	AF	$-8.40 \pm 3.43 \times 10^{-2}$	$3.00 \times 10^{-2} \pm 1 \times 10^{-4}$	283.6	264.4	294.8	327	120	207	2.8
Alto	AIP	$-7.19 \pm 4.39 \times 10^{-2}$	$2.77 \times 10^{-2} \pm 2 \times 10^{-4}$	234.2	217.0	258.7	300	134	166	1.5
Arma	AIP	$-9.78 \pm 2.71  imes 10^{-2}$	$4.15 \times 10^{-2} \pm 2 \times 10^{-4}$	248.4	232.6	267.9	313	118	195	7.0
Cada	AIP	$-5.86 \pm 3.22 \times 10^{-2}$	$2.36 \times 10^{-2} \pm 1 \times 10^{-4}$	226.7	209.3	250.4	231	78	153	1.1
Cast	AIP	$-4.81 \pm 2.95 \times 10^{-2}$	$2.22 \times 10^{-2} \pm 1 \times 10^{-4}$	230.0	203.5	252.5	410	94	316	4.1
Lamu	AIP	$-8.88 \pm 5.13  imes 10^{-2}$	$3.56 \times 10^{-2} \pm 2 \times 10^{-4}$	232.1	209.4	248.3	257	113	144	1.5
Leir	AIP	$-8.68 \pm 3.72 \times 10^{-2}$	$3.61 \times 10^{-2} \pm 1 \times 10^{-4}$	260.4	248.0	271.8	376	100	276	0.8
Puer	AIP	$-9.30 \pm 6.27 \times 10^{-2}$	$4.18 \times 10^{-2} \pm 2 \times 10^{-4}$	247.2	217.5	271.4	280	126	154	4.9
San	AIP	$-11.61 \pm 7.56 \times 10^{-2}$	$5.21 \times 10^{-2} \pm 3 \times 10^{-4}$	227.1	209.2	244.0	267	137	130	2.6
Sier	AIP	$-4.96 \pm 4.14 \times 10^{-2}$	$2.16 \times 10^{-2} \pm 2 \times 10^{-4}$	233.4	217.1	248.8	282	119	163	4.3
Pine	CO	na	na	na			467	127	340	na
Pini	CO	$-6.23 \pm 4.67 \times 10^{-2}$	$2.51 \times 10^{-2} \pm 2 \times 10^{-4}$	238.2	215.4	263.9	347	111	236	1.9
Oria	MO	$-8.59 \pm 3.06 \times 10^{-2}$	$3.56 \times 10^{-2} \pm 1 \times 10^{-4}$	231.9	219.8	243.8	338	101	237	4.9
Tamr	MO	$-9.54 \pm 4.84  imes 10^{-2}$	$3.47 \times 10^{-2} \pm 2 \times 10^{-4}$	296.6	273.1	316.7	383	105	278	3.2
Aren	MS	$-7.49 \pm 3.55  imes 10^{-2}$	$3.14 \times 10^{-2} \pm 1 \times 10^{-4}$	240.2	226.0	252.7	392	112	280	6.3
Bayu	MS	$-8.12 \pm 3.04 \times 10^{-2}$	$3.21 \times 10^{-2} \pm 1 \times 10^{-4}$	239.1	229.7	255.3	361	108	253	4.2
Carb	MS	$-8.38 \pm 6.01 \times 10^{-2}$	$3.64 \times 10^{-2} \pm 3 \times 10^{-4}$	228.8	213.0	249.8	290	117	173	8.3
Ceni	MS	$-6.03 \pm 4.93 \times 10^{-2}$	$2.86 \times 10^{-2} \pm 2 \times 10^{-4}$	208.6	172.1	246.4	353	132	221	1.8
Coca	MS	$-9.13 \pm 4.27 \times 10^{-2}$	$3.35 \times 10^{-2} \pm 2 \times 10^{-4}$	250.8	237.2	269.7	296	118	178	10.3
Cuel	MS	$-7.07 \pm 3.17 \times 10^{-2}$	$3.01 \times 10^{-2} \pm 1 \times 10^{-4}$	246.8	228.6	264.4	345	106	239	3.3
SanL	MS	$-9.77 \pm 4.18 \times 10^{-2}$	$4.00 \times 10^{-2} \pm 2 \times 10^{-4}$	229.7	207.7	255.7	283	80	203	4.4
Vald	MS	$-6.92 \pm 3.93 \times 10^{-2}$	$2.93 \times 10^{-2} \pm 2 \times 10^{-4}$	225.6	200.9	242.3	272	111	161	4.1

TABLE S4. Size-dependent fitness components for female reproduction in 23 natural *Pinus pinaster* populations. GG = genetic Group;  $f\mu$  = intercept of logistic regressions;  $f\alpha$  = slope of logistic regressions; fTSFR = median threshold size for female reproduction; loCI,upCI = lower and upper 95% limits of Bayesian credible intervals for fTSFR; fLVI = largest vegetative individual respect to female reproduction; fSRI = smallest individual bearing female cones; fRAN = transition range (fLVI–fSRI), fVEG = number of vegetative individuals. na not available.

Рор	GG	$f\mu \pm s.e.$	$f\alpha \pm s.$	e.	fTSFR	loCI	upCI	fLVI	fSRI	fRAN	%fVEG
Mimi	AF	$-6.79 \pm 2.23 \times 10^{-2}$	$2.00 \times 10^{-2} \pm$	$1 \times 10^{-4}$	316.8	301.2	346.9	404	129	249	2.1
Pleu	AF	$-5.75 \pm 1.71 \times 10^{-2}$	$2.15 \times 10^{-2} \pm$	$1 \times 10^{-4}$	258.0	236.3	280.9	422	115	212	7.4
Alto	AIP	$-5.47 \pm 2.22 \times 10^{-2}$	$1.16 \times 10^{-2} \pm$	$1 \times 10^{-4}$	426.6	368.0	522.8	403	126	174	0
Arma	AIP	$-5.32 \pm 2.32 \times 10^{-2}$	$1.51 \times 10^{-2} \pm$	$1 \times 10^{-4}$	381.7	341.0	451.0	391	92	221	0
Cada	AIP	$-6.42 \pm 3.05 \times 10^{-2}$	$1.43 \times 10^{-2} \pm$	$1 \times 10^{-4}$	458.4	399.0	531.9	470	138	93	0
Cast	AIP	$-5.53 \pm 2.59 \times 10^{-2}$		$1 \times 10^{-4}$	363.6	299.4	418.7	410	84	326	1.4
Lamu	AIP	$-4.97 \pm 1.76 \times 10^{-2}$		$1 \times 10^{-4}$	385.9	336.2	426.8	430	100	157	0
Leir	AIP	$-5.26 \pm 1.96 \times 10^{-2}$		$1 \times 10^{-4}$	490.4	399.4	576.8	448	100	276	0
Puer	AIP	$-6.62 \pm 3.02 \times 10^{-2}$	$1.39 \times 10^{-2} \pm$	$1 \times 10^{-4}$	450.6	391.5	524.0	429	149	131	0
San	AIP	$-4.31 \pm 2.27 \times 10^{-2}$		$1 \times 10^{-4}$	262.7	228.1	303.4	353	78	189	2.6
Sier	AIP	$-5.16 \pm 3.42 \times 10^{-2}$	$0.96 \times 10^{-2}$ ±	$1 \times 10^{-4}$	480.2	381.3	578.9	364	118	164	0
Pine	CO	na	na		na			467	263	204	na
Pini	CO	$-8.44 \pm 6.33 \times 10^{-2}$		$2 \times 10^{-4}$	368.8	294.0	448.9	458	147	200	0
Oria	MO	$-5.08 \pm 1.48 \times 10^{-2}$		$1 \times 10^{-4}$	284.7	256.3	310.2	366	85	253	2.1
Tamr	MO	$-6.40 \pm 1.73 \times 10^{-2}$		$1 \times 10^{-4}$	276.8	259.0	300.3	383	78	305	6.4
Aren	MS	$-5.77 \pm 1.99 \times 10^{-2}$		1 1 0	325.9	293.1	371.9	400	112	280	1.6
Bayu	MS	$-4.46 \pm 1.25 \times 10^{-2}$		$1 \times 10^{-4}$	246.9	233.0	269.6	366	94	267	3.5
Carb	MS	$-5.26 \pm 2.49 \times 10^{-2}$		$1 \times 10^{-4}$	233.2	211.2	271.3	314	99	191	6.3
Ceni	MS	$-5.43 \pm 2.53 \times 10^{-2}$		$1 \times 10^{-4}$	287.2	242.7	333.5	364	111	242	1.8
Coca	MS	$-5.41 \pm 3.47 \times 10^{-2}$		$1 \times 10^{-4}$	306.4	241.0	501.0	367	82	214	0
Cuel	MS	$-6.37 \pm 2.08 \times 10^{-2}$		$1 \times 10^{-4}$	229.5	203.3	281.8	383	72	273	3.3
SanL	MS	$-5.71 \pm 2.07 \times 10^{-2}$		$1 \times 10^{-4}$	285.2	236.7	335.7	349	103	180	0
Vald	MS	$-6.54 \pm 3.04 \times 10^{-2}$	$1.84 \times 10^{-2}$ ±	$1 \times 10^{-4}$	339.7	260.6	423.3	377	99	173	0

TABLE S5. Associations of environmental variables (first column) with reproductive life-history traits (second column) in *Pinus piaster*. Corrected values indicate slopes, standard errors and *P*-values for the association after including neutral genetic structure corrections (i.e. likelihood ratio test between a full model and a reduced model with just neutral genetic structure); uncorrected values indicate association parameters without correcting for neutral genetic structure (i.e. just environmental data and traits data in the model).

		Co	orrected		U	ncorrected	
	Trait	Slope		<i>P</i> -value	Slop	be	<i>P</i> -value
Alt	fTSFR	$-2.02 \times 10^{-1}$ ±	$7.02 \times 10^{-2}$	0.011	$-9.87 \times 10^{-2}$ ±	3.13×10 <sup>-2</sup>	0.005
	mTSFR	$-3.37 \times 10^{-2}$ ±	$1.86 \times 10^{-2}$	0.090	$1.39{ imes}10^{-3}$ ±	$9.22 \times 10^{-3}$	0.881
	Η	$-4.06 \times 10^{-2}$ ±	$3.06 \times 10^{-2}$	0.205	$-4.68 \times 10^{-2}$ ±	$9.77 \times 10^{-3}$	0.000
	%M	$-2.21 \times 10^{-2}$ ±	$1.54 \times 10^{-2}$	0.171	$-1.90 \times 10^{-2}$ ±	$6.65 \times 10^{-3}$	0.010
	% F	$1.57{ imes}10^{-2}$ ±	$1.24 \times 10^{-2}$	0.223	$1.33 \times 10^{-2}$ ±	$4.75 \times 10^{-3}$	0.011
Lat	fTSFR	$-8.22 \times 10^{-5}$ ±	$1.97 \times 10^{-4}$	0.682	$-7.02 \times 10^{-5}$ ±	$5.81 \times 10^{-5}$	0.241
	mTSFR	$-5.23 \times 10^{-5}$ ±	$4.45 \times 10^{-5}$	0.258	$2.09 \times 10^{-6}$ ±	$1.45 \times 10^{-5}$	0.887
	Н	$1.84 \times 10^{-5}$ ±	$7.31 \times 10^{-5}$	0.805	$-1.14 \times 10^{-6}$ ±	$2.25 \times 10^{-5}$	0.960
	%M	$2.01 \times 10^{-5}$ ±	$3.67 \times 10^{-5}$	0.593	$-6.81 \times 10^{-7}$ ±	$1.24 \times 10^{-5}$	0.957
	% F	$-1.95 \times 10^{-5}$ ±	$2.90 \times 10^{-5}$	0.512	$1.96 \times 10^{-7}$ ±	8.80×10 <sup>-6</sup>	0.982
Long	fTSFR	$2.08 \times 10^{-5}$ ±	$1.41 \times 10^{-4}$	0.885	$6.58 \times 10^{-5}$ ±	$5.81 \times 10^{-5}$	0.271
	mTSFR	$5.50 \times 10^{-5}$ ±	$3.01 \times 10^{-5}$	0.087	$-8.97 \times 10^{-5}$ ±	$1.43 \times 10^{-5}$	0.537
	Н	$1.84 \times 10^{-5}$ ±	$5.22 \times 10^{-5}$	0.729	$6.02 \times 10^{-5}$ ±	$1.79 \times 10^{-5}$	0.003
	%M	$-1.24 \times 10^{-5}$ ±	$2.63 \times 10^{-5}$	0.645	$1.61 \times 10^{-5}$ ±	$1.18 \times 10^{-5}$	0.189
	% F	$1.16 \times 10^{-5}$ ±	$2.09 \times 10^{-5}$	0.585	$-1.08 \times 10^{-5}$ ±	$8.42 \times 10^{-6}$	0.214
AMT	fTSFR	$30.58$ $\pm$	16.22	0.079	$29.46 \pm$	10.76	0.013
	mTSFR	-4.59 ±	4.07	0.277	$-1.63 \pm$	3.01	0.595
	Н	9.24 ±	6.24	0.159	10.26 ±	4.12	0.022
	%M	6.86 ±	2.88	0.031	6.46 ±	2.16	0.007
	% F	-4.87 ±	2.37	0.058	-4.31 ±	1.57	0.012
MTCM	fTSFR	29.43 ±	7.48	0.001	20.01 ±	3.99	0.000
	mTSFR	-0.91 ±	2.49	0.719	$-1.20 \pm$	1.42	0.407
	Н	8.36 ±	3.31	0.023	7.67 ±	1.44	0.000
	%M	4.83 ±	1.56	0.007	3.99 ±	0.85	0.000
	% F	-3.35 ±	1.33	0.024	-2.72 ±	0.63	0.000
MTWM	fTSFR	$-18.34 \pm$	6.06	0.009	$-14.18 \pm$	3.66	0.001
	mTSFR	-2.19 ±	1.72	0.221	-0.19 ±	1.16	0.872
	H	$-5.14 \pm$	2.52	0.059	$-5.61 \pm$	1.30	0.000
	%M	-2.12 ±	1.33	0.132	$-2.32 \pm$	0.85	0.013
ar	<u>% F</u>	<u>1.52 ±</u>	1.08	0.177	<u>1.78 ±</u>	0.59	0.006
CI	fTSFR	$-14.08 \pm$	3.39	0.001	-9.71 ±	1.91	0.000
	mTSFR	$-0.72 \pm 2.07$	1.15	0.542	$0.21 \pm 2.70$	0.69	0.762
	H		1.53	0.020		0.67	0.000
	%M	$-1.94 \pm$	0.79	0.026	$-1.75 \pm$	0.45	0.001
4.D	% F	$\frac{1.37 \pm}{10^{-2}}$	0.65	0.052	$\frac{1.27 \pm 1.27}{1.12 \pm 10^{-1}}$	0.31	0.001
AP	fTSFR	$4.22 \times 10^{-2} \pm$	$5.85 \times 10^{-2}$	0.482	$1.13 \times 10^{-1} \pm$	$4.28 \times 10^{-2}$	0.015
	mTSFR	$2.47 \times 10^{-3} \pm$	$1.40 \times 10^{-2}$ $2.06 \times 10^{-2}$	0.862	$\begin{array}{rrr} -6.57{\times}10^{-3} & \pm \\ 4.69{\times}10^{-2} & \pm \end{array}$	$1.19 \times 10^{-2}$ 1.52 \times 10^{-2}	0.586
	H	$3.09 \times 10^{-2} \pm 7.67 \times 10^{-3}$		0.153		$1.53 \times 10^{-2}$ $9.22 \times 10^{-3}$	0.006
	%M % E	$7.67 \times 10^{-3} \pm$	$1.10 \times 10^{-2}$	0.495	$2.00 \times 10^{-2} \pm$		0.043
	% F	$-1.23 \times 10^{-2} \pm$	$8.28 \times 10^{-3}$	0.157	$-1.77 \times 10^{-2} \pm$	$6.09 \times 10^{-3}$	0.009
PDM	fTSFR mTSEP	$1.14 \pm 1.14$	$5.19 \times 10^{-1}$	0.044	$5.54 \times 10^{-1} \pm$	$2.66 \times 10^{-1}$ 7.05 × 10 <sup>-2</sup>	0.050
	mTSFR	$1.47 \times 10^{-1} \pm$	$1.35 \times 10^{-1}$	0.294	$-1.11 \times 10^{-2} \pm$	$7.05 \times 10^{-2}$ $9.27 \times 10^{-2}$	0.877
	H V M	$2.39 \times 10^{-1} \pm$	$2.12 \times 10^{-1}$	0.277	$2.61 \times 10^{-1} \pm$		0.011
	%M % E	$1.81 \times 10^{-1} \pm$	$1.02 \times 10^{-1}$ $7.49 \times 10^{-2}$	0.096	$9.66 \times 10^{-2} \pm$	$5.64 \times 10^{-2}$ $3.81 \times 10^{-2}$	0.102
	% F	$-1.85 \times 10^{-1}$ ±	7.49×10	0.026	$-8.79 \times 10^{-2}$ ±	3.81×10	0.032

Population	Ν	C1	C2	C3	C4	C5	C6	GG
Mimi	19	0.74	0.01	0.02	0.01	0.06	0.15	AF
Pleu	21	0.67	0.02	0.03	0.01	0.12	0.16	AF
Alto	9	0.14	0.01	0.59	0.02	0.09	0.14	AIP
Arma	9	0.08	0.01	0.70	0.02	0.14	0.05	AIP
Cada	10	0.09	0.00	0.82	0.01	0.04	0.04	AIP
Cast	10	0.12	0.01	0.80	0.01	0.04	0.03	AIP
Lamu	9	0.10	0.01	0.73	0.01	0.10	0.06	AIP
Leir	24	0.19	0.01	0.24	0.01	0.42	0.13	AIP
Puer	8	0.05	0.01	0.76	0.04	0.10	0.05	AIP
San	12	0.15	0.01	0.51	0.02	0.25	0.06	AIP
Segu	21	0.12	0.01	0.50	0.01	0.23	0.13	AIP
Sier	10	0.21	0.01	0.41	0.02	0.13	0.21	AIP
Pine	10	0.05	0.03	0.03	0.80	0.05	0.04	CO
Pini	14	0.01	0.02	0.02	0.92	0.01	0.01	CO
Oria	29	0.10	0.54	0.07	0.03	0.16	0.11	MO
Tamr	24	0.01	0.95	0.01	0.01	0.01	0.01	MO
Aren	27	0.14	0.02	0.09	0.03	0.34	0.38	MS
Bayu	27	0.22	0.01	0.07	0.03	0.14	0.53	MS
Carb	6	0.28	0.04	0.11	0.06	0.20	0.31	MS
Ceni	9	0.30	0.01	0.13	0.02	0.19	0.35	MS
Coca	19	0.26	0.08	0.13	0.06	0.26	0.22	MS
Cuel	28	0.34	0.02	0.07	0.01	0.20	0.36	MS
Rodo	8	0.16	0.02	0.03	0.02	0.42	0.35	MS
SanL	20	0.14	0.06	0.07	0.06	0.11	0.56	MS
Vald	16	0.11	0.01	0.05	0.03	0.36	0.44	MS
Total	399							1110

TABLE S6. Number of trees per population included in the molecular marker analysis and average of individual assignment probability for each of the optimal K=6 clusters (C1 through C6) representing five geographical genetic groups (GG) in *Pinus pinaster*.

# APPENDIX III

Plasticity in reproduction and growth among 52 range-wide populations of
a Mediterranean conifer: adaptive responses to environmental stress.
Santos-del-Blanco L, Bonser SP, Valladares F, Chambel MR, Climent J.
2013. Journal of Evolutionary Biology. 26: 1912-1924.

## Plasticity in reproduction and growth among 52 range-wide populations of a Mediterranean conifer: adaptive responses to environmental stress

L. SANTOS-DEL-BLANCO\*<sup>†</sup>, S. P. BONSER<sup>‡</sup>, F. VALLADARES<sup>§</sup>, M. R. CHAMBEL<sup>\*</sup> & J. CLIMENT<sup>\*</sup><sup>†</sup>

\*Department of Forest Ecology and Genetics, INIA-CIFOR, Madrid, Spain

†Sustainable Forest Management Research Institute, INIA-University of Valladolid, Palencia, Spain

‡Evolution ∂ Ecology Research Centre, School of Biological, Earth and Environmental Sciences, The University of New South Wales, Sydney, NSW, Australia §Laboratorio Internacional de Cambio Global LINC-Global, Museo Nacional de Ciencias Naturales, MNCN-CSIC, Madrid, Spain

#### Keywords:

ecotypic trends; genotype-by-environment interaction; reproductive allometry; threshold size for reproduction; trade-offs.

#### Abstract

A plastic response towards enhanced reproduction is expected in stressful environments, but it is assumed to trade off against vegetative growth and efficiency in the use of available resources deployed in reproduction [reproductive efficiency (RE)]. Evidence supporting this expectation is scarce for plants, particularly for long-lived species. Forest trees such as Mediterranean pines provide ideal models to study the adaptive value of allocation to reproduction vs. vegetative growth given their among-population differentiation for adaptive traits and their remarkable capacity to cope with dry and low-fertility environments. We studied 52 range-wide Pinus halepensis populations planted into two environmentally contrasting sites during their initial reproductive stage. We investigated the effect of site, population and their interaction on vegetative growth, threshold size for female reproduction, reproductive-vegetative size relationships and RE. We quantified correlations among traits and environmental variables to identify allocation trade-offs and ecotypic trends. Genetic variation for plasticity was high for vegetative growth, whereas it was nonsignificant for reproduction. Sizecorrected reproduction was enhanced in the more stressful site supporting the expectation for adverse conditions to elicit plastic responses in reproductive allometry. However, RE was unrelated with early reproductive investment. Our results followed theoretical predictions and support that phenotypic plasticity for reproduction is adaptive under stressful environments. Considering expectations of increased drought in the Mediterranean, we hypothesize that phenotypic plasticity together with natural selection on reproductive traits will play a relevant role in the future adaptation of forest tree species.

#### Introduction

The timing of the onset of reproduction and the number of offspring produced by an individual are two fundamental life-history traits closely linked to fitness in an environment (Stearns, 1992; Braendle *et al.*, 2011). According to life-history theory, individuals that start reproducing earlier in life tend to be favoured under harsh environments, due to reduced life expectancy (Roff, 1992). The initiation of reproduction in plants is often related to size rather than age (De Jong & Klinkhamer, 2005). For example, individuals should build a large vegetative body and invest all available resources in reproduction just before death, that is, a bang-bang strategy (King & Roughgarden, 1982). But uncertainty about the moment of death, for example, due to disturbances will tend to favour reproduction at smaller sizes (and younger ages), a graded reproductive investment and bet-hedging strategies (Childs *et al.*, 2010). Thus, it is

*Correspondence:* Luis Santos-del-Blanco, Department of Ecology and Evolution, University of Lausanne, Biophore Building, CH1015 Lausanne, Switzerland. Tel.: +41 21 622 42 47; fax: +41 21 692 41 65; e-mail: luis.santosdelblanco@unil.ch

expected that plants, particularly long-lived perennials, will delay reproduction in favourable environments until they reach an optimal size for reproduction both by means of genetic change and phenotypic plasticity provided that selective forces act at local and broad scales (Kozłowski, 1992; Roff, 1992).

Experiments on herbaceous plants demonstrate that varying environmental factors - namely resource availability and competition - induce plasticity in reproductive strategies (Sultan, 2000; Weiner et al., 2009b; Anderson et al., 2011; Nicholls, 2011). In addition, plant populations are often genetically differentiated along environmental clines for size at reproduction and reproductive allometry, that is, the relationship between reproductive output and vegetative size (Lacey, 1988; Alexander et al., 2009; Guo et al., 2012). However, phenotypic plasticity of reproduction is driven to an important extent by size effects, as a strong positive relationship between vegetative and reproductive size is typically found and vegetative traits commonly respond plastically to environmental conditions. In comparison, phenotypic plasticity of the relationship between vegetative and reproductive size has been claimed to have a minor contribution to reproductive output, but this is still debated (Weiner et al., 2009a,b).

Long generation time in long-lived perennials implies that the same genotypes cope with year-to-year changing environmental conditions. On the other hand, populations of annuals or short-lived perennials can undergo genetic changes in shorter periods (Franks & Weis, 2008). Therefore, plasticity might be of greater importance as an adaptive strategy in trees and woody plants compared with short-lived plant species (Willson, 1983) such that long-lived species might exhibit plasticity in both vegetative (Chambel et al., 2005) and reproductive traits like size at reproduction and reproductive investment. The few studies published on long-lived species highlight strong selection on the threshold size at first reproduction and the allometry of reproduction, leading to genetic differentiation at large spatial scales (Thomas, 1996; Matziris, 1997; Niklas & Enquist, 2003; Climent et al., 2008; Santos-del-Blanco et al., 2010) and promoting phenotypic plasticity in life histories at local scales (Fang et al., 2006). Despite consistent predictions of plasticity in the threshold size of reproduction, little is known about the costs of plasticity in terms of final reproductive output relative to vegetative size (Roff, 2000). Reproductive efficiency (RE) can be defined as the slope of the reproductive-vegetative size developmental trajectory that connects threshold size for reproduction with reproduction at a given developmental stage or at the onset of senescence (Bonser & Aarssen, 2009). It is expected that early reproduction will imply lower RE, modifying reproductive allometries and, in turn, reproductive variability within and among populations. However, this has rarely been tested even in shortlived semelparous species (but see Bonser et al., 2010).

The Mediterranean pine Pinus halepensis Mill. (Aleppo pine) is a suitable model species for testing hypotheses on the evolution of reproductive strategies in long-lived perennials. It is precocious, bearing female cones from as early as 3 to 6 years of age, and commits heavily and regularly to reproduction, most notably female reproduction (Ne'eman et al., 2004). Pinus halepensis is widespread over a large circum-Mediterranean distribution area, and low population differentiation in neutral markers has been reported in the Iberian Peninsula due to recent range expansion (Soto et al., 2010). Pinus halepensis shows a wide ecological breadth among populations and is putatively adapted to a large range of abiotic stressors and perturbations, particularly fire and drought (Ne'eman et al., 2004), although intense drought episodes might be detrimental to reproduction (Girard et al., 2011). However, information regarding among-population variation in phenotypic traits in this species remains scarce.

Previous works described significant ecotypic differentiation for size at maturity in P. halepensis (Climent et al., 2008; Santos-del-Blanco et al., 2010). In this study, we focus on phenotypic plasticity and among-populations genetic variation in plasticity for reproductive allometry in range-wide P. halepensis populations assessed in a common garden experiment replicated in two contrasted sites (low and high environmental stress). Our objectives are to (i) assess the existence of phenotypic plasticity for size at maturity and the reproductive-vegetative size (R-V) relationship in range-wide populations subject to contrasting field conditions; and (ii) to compare genotype × environment patterns for vegetative and reproductive traits and correlations between both sets of traits representing trade-offs that might describe adaptive strategies. First, we expect that similar environmental cues defining favourable or unfavourable growth conditions will act in the same direction considering genetic differentiation and plasticity (Anderson et al., 2012; Chevin et al., 2012). Based on life-history theory, this would imply that the more stressful the environment (both at the origin of populations and at the trial site), the greater amount of resources would be devoted to reproduction. Specifically, we expect that environmental stress will induce reproduction at smaller sizes, associated with higher slopes of the R-V relationship. Finally, we also expect reproductive strategies to be governed by trade-offs between precocity and lifetime fitness, so that individuals that reproduce late benefit from a higher lifetime reproductive investment relative to their size.

#### **Materials and methods**

#### Study species and common gardens

A multisite *P. halepensis* common garden experiment was set up in 1997 replicated at six different sites in

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eastern and central Spain. The trial includes 52 native populations from continental Spain, Balearic Islands (Spain), France, Italy, Greece and Tunisia, as well as four non-native populations (see Climent *et al.*, 2008 for details) (Fig. 1, Table S1), thus covering most of the species' range. Only data relative to native populations were used in the present study. The minimum requirements for assessing plasticity in our experiment were, first, data measured at same age and identical protocols between sites and, second, contrasted enough environments. Only two of the six sites fulfilled both requirements.

Summary data of environmental conditions at both trial sites obtained from a functional model (Gonzalo-Jiménez, 2010) are shown in Table 1. Valdeolmos trial site (hereafter 'low-stress site') has sandy loam deep soil, whereas Rincón de Ademuz trial site (hereafter 'high-stress site') has shallow and rocky soil. In addition, mean annual rainfall is ca. 25% higher in the low-stress site, and winters are slightly warmer compared with the high-stress site. As a result of combined effects of poorer soil, lower rainfall and slightly colder winters, the high-stress site is much more limiting for P. halepensis vegetative growth compared with the lowstress site. This constant environmental difference between sites should not be confounded with withinsite year-to-year meteorological variation that has been previously described in this species (Girard et al., 2011).

Population seedlots were obtained by bulking openpollinated seeds from a subsample of 20 to 30 trees spaced at least 100 m apart in each population. At both sites, 832 one-year-old seedlings from native populations were planted in 1997 in a row-column design on the intersections of a  $2.5 \times 2.5$  m grid, with four replicates and four contiguous plants per population and replicate (16 trees per population). One replicate in the low-stress site was lost due to rabbit herbivory and was not included in this study (624 trees remaining). Due to other causes of mortality, final sample size for this study was 589 in the low-stress site and 633 in the high-stress site.

#### Measurement of traits and environmental variables

We measured height for each tree at ages 7, 11 and 13 years for both sites (2003, 2007 and 2009, respectively). Diameter at breast height was measured at both sites at ages 11 and 13 years and used to infer biomass from allometric models (Montero *et al.*, 2005) (Table 2).

The onset of female and male reproductive functions in *P. halepensis* is decoupled, with trees generally starting reproduction as females (protogyny) and male reproduction being delayed for up to several years (Shmida *et al.*, 2000). Thus, we focused on the study of *P. halepensis* early investment in female function to estimate threshold sizes for first reproduction and reproductive investment.

Female cones in *P. halepensis* remain attached to the branches even after dehiscence (normally also delayed several years, Ne'eman *et al.*, 2004). Differences in size and colour allow discrimination of several cohorts within tree crowns (Ne'eman *et al.*, 2011) and therefore enable retrospective record of female reproduction. Up to three successive cohorts of female cones were counted at ages 7 and 13 years (2003 and 2009) therefore dating back to the very first reproductive events up to the generalization of reproduction at both sites.

We defined the cumulative reproductive investment (CRI) as the sum of all counted female cones produced



**Fig. 1** Distribution map of *Pinus halepensis* source populations (circles) and common garden (stars). Green areas indicate the species' natural distribution range.

Table 1 Climatic descriptors for two Pinus halepensis common
garden study sites, derived from functional climatic models
(Gonzalo-Jiménez, 2010).

Abr.	Site	Valdeolmos Low-stress site	Rincón de Ademuz High-stress site
Long	Longitude	3°26'44"W	1°14′14″W
Lat	Latitude	40°38'42"N	40°06'38"N
Alt	Altitude (m)	731	844
SP	Spring precipitation (mm)	129	99
PDQ	Precipitation of the warmest quarter (mm)	62	94
PDM	Precipitation of the driest month (mm)	13	23
Ρ	Annual precipitation (mm)	475	364
AMT	Annual mean temperature (°C)	12.9	12.3
MTWM	Mean temperature of the warmest month (°C)	29.9	27.6
MTCM	Mean temperature of the coldest month (°C)	0.7	0.2

by an individual until last measurement at age 13 (Table 2).

We collected data for six meteorological and three spatial variables describing the environmental conditions found in the natural populations (Table 1, Table S1). Meteorological data for Iberian populations were obtained from a functional model (Gonzalo-Jiménez, 2010), and data for other populations (i.e. Balearic Islands, France, Tunisia, Italy and Greece) were obtained from WorldClim-Global Climate Data at 5' resolution (Hijmans *et al.*, 2005). To test hypotheses of local adaptation, we also calculated the Gower's ecological distance for each population at both trial sites (Rutter & Fenster, 2007). This adimensional index informs about the environmental distance between the native environment of each population and the environment where they were grown in the common garden. The analysis was limited to Iberian and Balearic populations due to the unbalanced number of eastern Mediterranean populations in the experiment.

#### Data analysis

All reported models and tests were implemented in *R* (R Development Core Team., 2012) using packages lme4 (Bates *et al.*, 2011) and MCMCglmm (Hadfield, 2010).

#### Survival and vegetative size

Mixed linear models for size (height and biomass) at age 13 and generalized linear mixed models (logit link, binomial error) for survival and proportion of reproductive individuals at age 13 were fitted. In all models, site effect was treated as fixed. Population, site-by-population interaction and replicate within site were treated as random. A common interpretation of model parameters is as follows: significant differences among populations indicate intraspecific genetic variability; significant differences between sites reflect phenotypic plasticity, and significant site  $\times$  population interaction indicates genetic variation for plasticity among populations (Schlichting, 1986). However, deviations from that framework need also to be considered. For example, environmental factors can significantly affect seeds

**Table 2** Common garden measured and derived vegetative and reproductive traits of *Pinus halepensis* trees. Measurement age inparenthesis.

Variable		Description
Measured traits		
Н	Height (cm)	Total height of each tree (7, 11, 13)
DBH	Diameter at breast height (cm)	Diameter of the tree trunk at 1.30 m (11,13)
CC	Cone number (n)	Number of seed cones in each tree per cohort (7, 13)*
Surv	Survival	Status of each tree: dead (0) or alive (1) (7, 11, 13)
Derived traits		
Vegetative		
Biomass	Biomass (kg)	Biomass inferred from DBH (Montero et al., 2005)
Reproductive		
TSFR	Median threshold size for first reproduction (cm)	Size at which the probability for a tree within a population to have reached sexual maturity was 50%
CRI	Cumulative reproductive investment (n)	Sum of all seed cones produced by a tree
BA	Reproductive allocation ( <i>n</i> per kg)	Number of seed cones divided into total tree biomass
R–V intercept	heproductive anosation (7 per kg)	Intercept of R–V GLMM Poisson model
R-V slope		Slope of R–V GLMM Poisson model
RE	Reproductive efficiency	CRI/(height at last measurement – TSFR)

\*7 corresponds to ages 5 and below, 6 and 7 years; 13 corresponds to 11, 12 and 13 years.

during development, causing epigenetic changes in gene expression (Johnsen *et al.*, 2005). Also, a significant site  $\times$  population interaction can indicate local adaptation if populations have a better performance in the site most similar to the conditions of their site of origin (Vergeer & Kunin, 2013). To test the significance of site, population and site  $\times$  population terms, we performed likelihood ratio tests (LRTs) comparing full models containing all terms with those lacking the relevant term to be tested. Variance components and adjusted means for size at final measurement of each population were derived from analogous models fitted for each trial site.

#### Size at first reproduction

A generalized linear mixed model (logit link, binomial error) was fitted for cumulative female reproduction (either present or absent) data at ages 7 and 13. We included height as covariate and height × site interaction as fixed term. Then, independent models were fitted for each population and site by Markov Chain Monte Carlo (MCMC) methods (see Santos-del-Blanco et al., 2012 for further details), and median threshold size for first reproduction (TSFR) was defined as the size at which the probability for a tree to have reached sexual maturity was 50% (Wesselingh et al., 1997) and computed by dividing slope by intercept estimates. We also calculated the size of the smallest reproductive individual (SRI) at each population and used this information to classify nonreproducing trees into juvenile (smaller than SRI) or vegetative (larger than SRI) (Mendez & Karlsson, 2004).

*Fecundity and reproductive–vegetative size relationships* Generalized linear mixed models (log link, Poisson error) were fitted to the CRI. Juveniles were removed from the data set prior to analysis. The models also included an individual-level random effect to model additive overdispersion (Elston et al., 2001).

Reproduction in plants is typically size dependent (Niklas & Enquist, 2003; Weiner, 2004). We accounted for size-dependent differences in reproductive allocation by calculating reproductive allocation per population first as the mean value across individuals (CRI/biomass) and second as the expected reproductive value based on fitted reproductive–vegetative size (R–V) models and then divided into average size. Thus, each approach represents the mean reproductive allocation per population and reproductive allocation of an average-sized individual in a population, respectively. Similar values for both indexes would indicate that the estimation of reproductive allocation is robust, although issues remain about spurious correlations with size.

Reproductive–vegetative size models describe the relationship between reproductive and vegetative allocation using two parameters – an intercept and a slope (Weiner *et al.*, 2009a; Guo *et al.*, 2012). However,

depending on the range of sizes used to fit the models, those two parameters might not be independent in a set of populations due to collinearity (Pinheiro & Bates, 2000). Thus, to summarize reproductive output and compare R–V relationships while accounting for tree vegetative size, we fitted generalized linear mixed models (log link, Poisson error) with independent intercept and slopes to CRI data according to:

$$\eta_i = \ln(\mu_i) = \mathbf{x}'\boldsymbol{\beta} + \mathbf{z}'\mathbf{b} + \mathbf{e}; \mathbf{b} \sim N(0, \sigma^2); \mathbf{e} \sim \operatorname{Pois}(\lambda)$$

where  $\eta_i$  is the linear predictor linked to the expected value of natural logarithm of CRI [ln  $(\mu_i)$ ]. x' represents the design matrix containing the values for the fixed size effects.  $\beta$  is a vector containing the fixed intercept and slope associated with size, to be estimated. z' is the design matrix for the random populations effects. b is the vector of random coefficients that follow a normal distribution. e is the vector containing the errors that follow a Poisson distribution. Two models were fitted per site, the first one with b containing random effects for intercepts and the second containing random effects for the slopes. AIC values from both models at each site were very close, indicating that either random intercept or random slope models had similar explanatory power. Population-adjusted intercepts and slopes were derived from MCMC models fitted at each site and used as fecundity indicators; this allowed us to compare general estimates from both sites. Random intercepts associated with population reflect constant deviations across sizes from the general model, that is, a constant higher or lower commitment to reproduction across sizes. Random slopes associated with population represent deviations on reproductive output proportional to vegetative size, that is, enhanced or decreased commitment to reproduction along vegetative size. Our analysis of size at first reproduction and reproductive output divided in two steps (binomial and Poisson submodels) was thus similar to a hurdle model (Brophy et al., 2007; Haymes & Fox, 2012).

#### *Reproductive efficiency*

We estimated RE as the slope of the size-reproduction developmental trajectory, linking vegetative size at first reproduction and vegetative and reproductive output at final development (age 13) (Bonser & Aarssen, 2009). RE was estimated at the population level for both sites. We tested whether there were significant correlations between the threshold size for reproduction and RE at each site and whether RE was affected by the environment, comparing RE values between both sites with a paired *t*-test.

#### Local adaptation patterns

Pearson's correlation tests at each trial site were used to test the relationship between Gower's distance and fitness. We used CRI and female TSFR, as the variables most closely related to fitness but also explored the correlation between Gower's distance and vegetative growth traits (Leimu & Fischer, 2008). We also tested whether increased environmental distances were correlated with changes in trait means.

#### Plant trait correlations and ecotypic trends

We calculated Pearson's correlations among plant traits at the population level and among those traits and environmental conditions found in the natural populations. Correlations among plant traits can be interpreted as genetic correlations modified by common environmental effects. Correlations were conducted at each trial site separately to check whether trait–trait correlations and ecotypic trends of variation were site dependent. We also obtained the site-to-site correlations for phenotypic traits, as a double-check of site-by-population interaction (Pigliucci, 2001).

#### Results

#### Vegetative traits

We found that plants in the high-stress site had lower biomass and height compared with those in the lowstress site, thus confirming that overall environmental differences between both sites had an effect on vegetative growth (Table 3, Fig. 2). In addition, tree survival was significantly lower in the high-stress site compared with the low-stress site ( $\chi_1^2 = 76.2$ , *P* < 0.001) (Table 3). All populations attained larger sizes in the low-stress than in the high-stress site. However, there was no evidence for population effect alone, but differences between populations were site specific, and a significant site-by-population interaction was found for all vegetative traits (Table 4, Figs 2 and 3), consistent with genetic variation in plasticity for vegetative traits among populations and, possibly, local adaptation. Among-population variance was larger for biomass in

**Table 3** Mean values ( $\pm$  standard errors or credible intervals in brackets) for *Pinus halepensis* vegetative and reproductive traits at two experimental sites with contrasting environmental conditions. Values at each site are averaged across 52 natural populations; abbreviations and units are as defined in Table 2.

	Low-stress site	High-stress site
Н	339.6 ± 3.4	274.9 ± 3.0
Biomass	$12.16 \pm 0.67$	$4.10 \pm 0.24$
Survival	$0.65 \pm 0.03$	$0.61 \pm 0.01$
CRI	$17.1 \pm 1.1$	$16.3\pm0.8$
RA	$5.61 \pm 1.11$	$20.09 \pm 3.99$
TSFR	$204.5 \pm 6.9$	$147.3 \pm 6.1$
Intercept	1.60 (1.36–1.77)	2.05 (1.87-2.20)
Slope	0.033 (0.030–0.042)	0.070 (0.056–0.090)

CRI, cumulative reproductive investment; TSFR, threshold size for first reproduction; RA, reproductive allocation.

the low-stress site [45.0 (31.6–69.2)] compared with the high-stress site [4.7 (3.3–7.2)], but no significant differences were found for height [829 (582–1275) lowstress site; 797 (560–1225) high-stress site]. Population means for vegetative traits at the low- and high-stress sites can be accessed in Tables S2 and S3, respectively.

#### **Reproduction and threshold sizes**

Mean size of reproductive individuals was greater than that of nonreproductive ones at both sites and both years (low-stress site: biomass  $\chi_1^2 = 21.0$ , height  $\chi_1^2 = 34.7$ ; high-stress site: biomass  $\chi_1^2 = 85.8$ , height  $\chi_1^2 = 242.1$ , all tests P < 0.001). At the early measurement date (age 7 years), the proportion of reproductive individuals was slightly greater in the low-stress site than in the high-stress site ( $\chi_1^2 = 58.6$ , P < 0.001). However, at 13 years of age, 96% of trees were reproductive in the stressed environment, whereas only 84% were in the more favourable environment ( $\chi_1^2 = 34.7$ , P < 0.001). As a result, at final measurement, the number of vegetative individuals was higher in the low-stress than in high-stress site (34 vs. 5).

We found a significant effect of both site and population on the threshold size for reproduction, as shown by the significant site and population terms (Table 4). Thus, threshold size for reproduction is both highly plastic and variable among populations (Figs 2 and 3). By contrast, the site-by-population interaction term was not significant, indicating that there was no significant genetic variation for plasticity in the threshold size for reproduction among populations. The probability of reproducing at a given size was significantly smaller in the low-stress site than in high-stress site, evidenced by a reduced slope of the model (data not shown).

We were able to fit independent threshold models for all but three populations in the low-stress site and all populations but one in the high-stress site. For all but two populations, the point estimate of the threshold size for reproduction was higher in the low-stress site than in the high-stress one (Figs 2 and 3).

For CRI, site and population effects were significant, but not, although marginally, population-by-site interaction (Table 4). When tree biomass was included as a covariate in the R-V model, site (indicating a different R-V relationship in both sites), and population terms were significant, but not site × population interaction (Table 4). When height was used as a covariate, similar results were obtained although site was not significant (Table 4). The subsequent GLMM models fitted by MCMC aimed at estimating fecundity at the population level while controlling for size effects revealed an enhanced reproductive allocation in the high-stress site with respect to the low-stress one, defined by a larger intercept and slope (Table 3). Here, a positive intercept must not be regarded as biologically implausible, as it represents a population, not an individual developmental trajectory. Mean reproductive allocation per



**Fig. 2** Site—site graphs for the interpretation of phenotypic plasticity in several vegetative and reproductive traits measured in two *Pinus halepensis* provenance common gardens subject to contrasting environmental conditions (low and high environmental stress). Points represent mean values per source population. Units are as defined in Table 2.

population and expected reproductive allocation of an average-sized individual per population yielded similar results (data not shown), so we used the first index to describe reproductive allocation (RA) as it was derived primarily from the data. Consistently with fecundity descriptors, RA was larger in the high-stress than in the low-stress site. Population means for reproductive traits at the low- and high-stress sites can be accessed in Tables S2 and S3, respectively.

#### Plant trait correlations and ecotypic trends

Differences in CRI were related to the Gower's environmental distance at both trial sites. That is, there was a significant positive relationship between the environmental similarity between each population with respect to the common garden and the number of cones it produced as revealed by Pearson's correlation tests (Table 5). Female threshold size for reproduction was also significantly negatively related to Gower's distance in the low-stress site but only marginally in the high-stress site. By contrast, for vegetative traits, even when correlations were only marginally significant, they showed opposite patterns at each trial site. Correlations at the high-stress site were negative and at the low-stress site were positive (Table 5). The threshold size for reproduction was the only variable significantly correlated with a change in environmental distance. Closer distances were related to larger thresholds for reproduction. Correlation with CRI was negative but nonsignificant (Table 5). **Table 4** Results of general and generalized linear mixed models for *Pinus halepensis* vegetative and reproductive traits measured in two experimental sites with contrasting environmental conditions. Full models were fitted including all terms. Site, population and population-by-site models were fitted excluding the relevant terms to test plasticity, genetic variation and genetic variation for plasticity. log-likelihood (logLik) is given for each model. Chi-square statistic (Chisq), degrees of freedom (d.f.) and *P*-value are given for likelihood ratio tests between the full model and reduced models.

	logLik	Chisq	d.f.	P-value
Vegetative traits				
Biomass				
Full model	-3449			
Site	-3452	5.6	1	0.018*
Population	-3450	1.7	1	0.190
Site × population	-3466	33.7	1	<0.001***
Н				
Full model	-5153			
Site	-5156	6.135	1	0.013*
Population	-5153	0.908	1	0.341
Population × site	-5166	25.6	1	<0.001***
Reproductive traits				
CRI:				
Full model	-1646			
Site	-1649	5.8	1	0.016*
Population	-1658	23.3	1	<0.001***
Site × population	-1648	3.8	1	0.051†
CRI~h‡				
Full model	-1544			
Site	-1544	0.025	1	0.875
Population	-1564	40.59	2	<0.001***
Population × site	-1545	3.2	2	0.200
CRI~biomass‡				
Full model	-1575			
Site	-1581	13.3	1	<0.001***
Population	-1592	34.97	2	<0.001***
Site × population	-1576	2.7	2	0.257
TSFR§				
Full model	-857			
Site	-876	37.6	1	<0.001***
Population	-871	27.1	2	<0.001***
Site × population	-859	3.0	2	0.222

CRI, cumulative reproductive investment; TSFR, threshold size for first reproduction.

‡Poisson model.

§Binomial model.

\*P < 0.05, \*\*\*P < 0.01; †P < 0.1.

Site–site correlations of population-adjusted means were not significant for vegetative traits, but strong positive correlations were found for reproductive traits (Table S4). This corroborates the high site-by-population interaction for vegetative traits vs. nonsignificant site-by-population interaction for reproductive traits seen by previous analyses (Table 4).

Within sites, correlations among traits representing potential trade-offs between growth and reproduction (e.g. height and reproductive allocation) showed, in general, stronger correlations in the high-stress site than in the low-stress one (Table S4). We found significant positive correlations between height and TSFR in the high-stress site, but not in the low-stress one. All other correlations between vegetative and reproductive traits were nonsignificant. We also found extensive correlations within vegetative traits and reproductive traits (Table S4).

We found no significant difference in RE between sites ( $t_{47} = 1.442$ , P = 0.156), nor for growth above the site-specific median threshold size for reproduction ( $t_{47} = -1.037$ , P = 0.305). RE was negatively correlated with TSFR in the low-stress site but not in the high-stress site (Table S4).

Correlations between environmental factors and plant traits, reflecting ecotypic trends of variation, were higher in reproductive traits compared with vegetative traits. In turn, they were higher in the low-stress site, but in both sites, the sign of the correlation was the same. Traits indicative of more precocious or abundant reproduction were related to higher altitude, and warmer summers and colder winters (therefore higher continentality index). However, no significant correlations were found between rainfall and any of the phenotypic traits measured at either site (Table S4).

#### Discussion

Our experiment showed that at the more stressful site, P. halepensis trees started reproducing at smaller sizes and completed female reproductive maturity earlier both in size and in time – than at the least stressful site, therefore following theoretical expectations (Roff, 1992). By definition, threshold size for reproduction accounts for differences in size, so a plastic response in this trait should be considered as a true plastic response and not driven solely by plasticity in size (Sugiyama & Bazzaz, 1998; Weiner, 2004). Hypotheses regarding plasticity of threshold size for reproduction have been addressed in plants only in few cases (Bonser & Aarssen, 2009; Kagaya et al., 2009; Bonser et al., 2010), in contrast with predictions of the high relevance of this type of plasticity (Burd et al., 2006). However, evidences pointing at this phenomenon are common through the literature (Nagy & Proctor, 1997; Fang et al., 2006). We relied on two natural environments to test our hypothesis, which also allowed us to study local adaptation patterns. Nonetheless, a more precise control of environmental stress could be achieved by artificially inducing drought or watering or by setting the experiment at contrasting soil depths and/or nutrient levels, for example, leading to more general conclusions.

The adaptive value of reproduction at larger sizes in favourable conditions relies on a positive relationship between fecundity and size at reproduction, so that

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Fig. 3 Reaction norms for vegetative and reproductive traits measured in two *Pinus halepensis* provenance common gardens subject to contrasting environmental conditions (low and high environmental stress). Lines connect mean values per source population at each common garden. Units are as defined in Table 2.

Table 5         Pearson's correlations and <i>t</i> -tests between <i>Pinus halepensis</i>
adjusted population mean values for plant traits and Gower's
environmental distance at either high- or low-environmental-
stress experimental sites. 'Between sites' refers to the correlation of
differences in Gower's distance between sites and differences in
mean values for plant traits.

Trait	Site	r	t	d.f.	P-value
Vegetative tra	iits				
Biomass	High stress	-0.31	-2.07	40	0.045*
	Low stress	0.18	1.17	40	0.250
	Between sites	0.10	0.65	40	0.521
Н	High stress	-0.29	-1.90	40	0.065†
	Low stress	0.27	1.76	40	0.086†
	Between sites	-0.05	-0.35	40	0.730
Reproductive	traits				
CRI	High stress	-0.35	-2.38	40	0.022*
	Low stress	-0.52	-3.85	40	0.000***
	Between sites	-0.25	-1.65	40	0.108
TSFR	High stress	0.27	1.80	40	0.080†
	Low stress	0.49	3.42	37	0.002**
	Between sites	0.45	3.07	37	0.004**

*r*, Pearson's correlation, *t*, *t* statistic, d.f., degrees of freedom; CRI, cumulative reproductive investment; TSFR, threshold size for first reproduction.

Significant values \*\*\*< 0.001, \*\*0.01, \*0.05, †0.1.

attaining a larger size implies an increased lifetime reproductive output (Metcalf *et al.*, 2003). Although this relationship is clear in semelparous species, as they only have one reproductive event in their life (bangbang strategy), in iteroparous species like trees the relationship is not straightforward because individuals allocate significant amounts of resources to maintenance each season throughout their lives (De Jong & Klinkhamer, 2005). For example, two trees with a similar adult size might differ in reproductive output due to differential investment in maintenance along their lives, whereas in annual plants, no differences in reproductive output are expected for similar-sized individuals (Weiner *et al.*, 2009a).

In plants, favourable environments for growth generally favour lower reproductive investment relative to size (Matyas & Varga, 2000; Ortiz *et al.*, 2011; Haymes & Fox, 2012). As these conditions are typically associated with increased competition (Grime, 1977), delayed reproduction in these environments could be driven both by a positive relationship between size at reproduction and lifetime reproductive investment, but also by a persistent allocation to growth and maintenance in crowded stands (Zhang, 2006).

At the population level, a low threshold size for reproduction was correlated with steeper slopes for the R-V relationship at both trial sites. Thus, genetic and

environmental factors promoting allocation to reproduction acted consistently along the development of the trees in our experiment. Trade-offs between reproduction and growth are predicted to be more relevant in limiting conditions (Karlsson & Mendez, 2005). Accordingly, in the more stressful site, we found a negative correlation between vegetative traits and reproductive precocity (hence positive with TSFR), and correlation coefficients between reproductive allocation and size among populations were higher than in the less stressful site. However, contrary to our expectations, we found no differences for RE between sites; that is, delayed onset of reproduction in the low-stress site was not rewarded with a proportionally higher reproductive output. We did not extensively test for adaptive plasticity by inducing plants to express an inappropriate phenotype in a given environment (see Sultan, 2000). Nonetheless, our findings regarding enhanced reproduction at the more stressful site highlight the adaptive value of plasticity for reproduction (Anderson et al., 2012) as they support theoretical expectations (Pigliucci, 2001). Indeed, if reproduction at the high-stress site had followed the same allometric trend as in the low-stress site, the risk of becoming locally extinct after a severe disturbance would be dramatically higher. However, at the same time, our results raise uncertainty about what the benefits of delayed reproduction are in environments favourable to vegetative growth. Also, as we did not consider male reproduction in our analysis as the trees were still young, we cannot rule out a possible trade-off between female and male reproduction, so that female reproduction is reduced in favourable environments, but male reproduction could be enhanced. To gain better insight into these uncertainties, data covering a longer time period for both sexual functions would be needed.

In our experiment, tree size was a weak predictor of reproductive output in both trial sites. Although our trees were young and the relationship may strengthen with time (Weiner et al., 2009a), a loose, although significant, relationship between size and reproduction is common in perennial species, notably in trees (Climent et al., 2008; Haymes & Fox, 2012). Population  $\times$  site interaction was important for vegetative traits, but we found no evidence for larger plants corresponding to shorter environmental distances (Vergeer & Kunin, 2013). Actually, for the low-stress site, the trend was opposite, being larger plants from ecologically distant populations. Instead, reproductive output did show a negative relationship with the environmental distance of the original populations to each trial site. As reproductive output is closely linked to fitness, this suggests that populations have adapted to local climate conditions, and climate is important in controlling the expression of reproduction (Leimu & Fischer, 2008). Thus, we advise against using tree size as a proxy for

fitness and encourage the use of reproductive output in tree evolutionary ecology studies.

Several additional factors might interact with raw reproductive output to define individual fitness (Braendle et al., 2011). Within populations, some individuals remained nonreproductive well above their population TSFR, a phenomenon also described in biennials (Wesselingh & Klinkhamer, 1996), and the highest reproductive output was typically achieved by mediumsized individuals in consistency with other experiments in this species (Climent et al., 2008). This pattern was more evident in the low-stress site, where a higher number of trees remained vegetative. A likely explanation for this observation is a diversifying bet-hedging strategy (Simons, 2007), with individuals reproducing according to a genetically determined allocation curve and others situated below that curve (Weiner, 2009a). If a disturbance occurred at either trial site, population and individual would be the most important factors determining the number of available seeds for the next generation. This would imply that if the primary reason for delaying reproduction was a larger future reward through increased size and greater potential future reproduction, many individuals would be making a nonprofitable investment. However, an enhanced allocation to growth would also increase fitness through an increased likelihood of survival (Zhang, 2006). The relative importance of these nonexclusive explanations deserves more attention to better understand adaptive responses in trees.

Contrary to expectations, reproductive output for the whole set of populations was very similar between the two contrasting environments. Plasticity for cumulative cone production was much lower (up to twofold) than that for biomass (up to 10-fold) (Fig. 2). Reproductive output emerged from a combination of plastic responses in growth (larger in the less stressful site) and allometry (higher reproduction for a given size in the more stressful site). An ecotypic trend of enhanced reproduction towards higher altitudes and more extreme temperatures, already described in Climent *et al.* (2008), was not related to population differentiation in plasticity. Interestingly, we found plasticity for both reproductive allometry and vegetative traits, but only genotype-by-environment interaction for vegetative traits.

Phenotypic plasticity is expected to arise in environments that change in a predictable fashion (Van Kleunen & Fischer, 2005). Within species, higher plasticity is generally expected in populations subject to greater interannual variance in precipitation and extreme temperatures and also those living in more patchy environments (Sultan & Spencer, 2002; Baythavong, 2011). In addition, traits might differ in their sensitivity to the environment, or may be constrained resulting in some being more plastic than others (Matesanz *et al.*, 2010), as is the case in our experiment, where not only phenotypic plasticity but also its variation among

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populations was higher for growth than for reproduction, consistent with findings in different plant genera (reviewed in Weiner et al., 2009a). In our experiment, the lack of population differentiation for plasticity for reproductive allometry could be due to nonexclusive causes such as (i) a strong stabilizing selection for plasticity of reproductive allometry among populations, (ii) a canalization or total dependence of reproductive traits on vegetative traits, like internal cues, or (iii) the perception of environmental heterogeneity, selecting for plasticity, differing between reproductive and vegetative traits. For example, vegetative traits may be more dependent on fine-grain variability of soil depth or nutrient availability, but reproductive traits depend more on factors acting at a larger scale like climate and severe perturbations. In the high-stress site, variation in responses for vegetative traits was constrained, whereas in the low-stress site, among-population differences were neatly expressed, revealing cryptic genetic variation (Schlichting, 2008). Reproductive traits, however, displayed similar levels of variation at both trial sites, so in the environment with most limiting conditions, variation for reproductive traits was more relevant than that for vegetative traits.

Considering expectations of increased drought in the Mediterranean due to climate change (Lindner *et al.*, 2010) and assuming a high heritability of reproductive allometry (Santos-del-Blanco *et al.*, 2010 and in prep; Wesselingh & De Jong, 1995), we hypothesize that phenotypic plasticity coupled with subsequent natural selection on this trait (Anderson *et al.*, 2012; Chevin *et al.*, 2012) will play a relevant role in future adaptation of forest species.

#### Acknowledgments

The authors declare no conflict of interest. The data used in this research are part of the Spanish Network of Genetic Trials (GENFORED). Data are publicly available upon request through www.genfored.es. We thank E. Ballesteros, F. del Caño, D. Barba and E. Alamillo for fieldwork support. Luis Samperdo (CSIC), Jill T. Anderson (U. South Carolina) and two anonymous reviewers provided insightful comments that improved the manuscript. This study was funded by the Spanish Ministry of Economy and Competitiveness through the project Mitigenfor (RTA 2011-00016-00-00). Regina Chambel was granted by the Spanish Ministry of Economy and Competitiveness for technical support to GENFORED network.

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#### **Supporting information**

Additional Supporting Information may be found in the online version of this article:

**Table S1** List and location of the *Pinus halepensis* populations comprised in the present study and planted at two trial sites with contrasting environmental conditions (high and low environmental stress).

**Table S2** Vegetative and reproductive traits in 52 natural *Pinus halepensis* populations grown in a common garden placed at Valdeolmos (Madrid, Spain), referred as low stress site.

**Table S3** Vegetative and reproductive traits in 52 natural *Pinus halepensis* populations grown in a common garden placed at Rincón de Ademuz (Valencia, Spain), referred as high stress site.

**Table S4** Pearson correlation coefficients for the correlation at the population level between climatic variables and *Pinus halepensis* traits in each site, the low stress site above diagonal and the high stress site below the diagonal.

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1	Supp	lementary	material
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1	Supplementary material
2 3	Plasticity in reproduction and growth among 52 range-wide populations of a
4	Mediterranean conifer: adaptive responses to environmental stress
5	Authors: Santos-del-Blanco, Luis <sup>1,2</sup> *; Bonser, Stephen P. <sup>3</sup> ; Valladares, Fernando <sup>4</sup> ; Chambel,
6	Maria Regina <sup>1</sup> , Climent, José <sup>1,2</sup> .
7	<sup>1</sup> INIA-CIFOR, Dpt. Forest Ecology and Genetics. Ctra A Coruña km 7.5, E-28040, Madrid,
8	Spain
9	<sup>2</sup> Sustainable Forest Management Research Institute, INIA-University of Valladolid,
10	http://sostenible.palencia.uva.es
11	<sup>3</sup> Evolution & Ecology Research Centre, School of Biological, Earth and Environmental
12	Sciences, The University of New South Wales, NSW 2052, Sydney, Australia
13	<sup>4</sup> Laboratorio Internacional de Cambio Global LINC-Global, Museo Nacional de Ciencias
14	Naturales, MNCN-CSIC, E-28006, Madrid, Spain
15	Running title:
16	Plasticity in plant reproductive traits
17	*corresponding author:
18	e-mail address: santos.luis@inia.es
19	Tel +34 91 3476862,
20	Fax +34 91 3476767.
21	INIA-CIFOR, Dpt. Forest Ecology and Genetics.
22	Ctra A Coruña km 7.5,
23	E-28040, Madrid, Spain
24 25	

26 TABLE S1. List and location of the *Pinus halepensis* populations comprised in the present

study and planted at two trial sites with contrasting environmental conditions (high and low

28 environmental stress).

Code	Population	Country	Altitude	Latitude	Longitude	AP	PDQ	AMT	MTWM		CI
11	Cabanellas	Spain	258	977212	4693568	858	208	14.2	28.3	1.3	27.0
21	Tivissa	Spain	336	815989	4552144	566	75	15.3	30.6	3.2	27.4
31	Sant Salvador de Guardiola	Spain	318	896567	4624516	569	123	13.9	30.0	0.9	29.1
61	Zuera	Spain	576	672350	4642850	488	95	12.9	31.6	-0.1	31.7
82	Valdeconcha	Spain	837	510850	4477650	579	80	12.7	32.2	-1.2	33.4
83	Alcantud	Spain	1057	558120	4490600	819	100	10.5	30.9	-2.2	33.1
84	Colmenar de Oreja	Spain	692	471558	4437895	450	55	13.7	33.0	0.0	33.0
91	Cirat	Spain	445	715916	4436492	472	109	15.4	31.1	2.6	28.5
92	Tuéjar	Spain	729	657550	4409300	385	91	15.0	33.2	1.0	32.2
93	Enguidanos	Spain	990	615980	4388800	572	85	12.5	32.2	-0.7	32.9
101	Tibi	Spain	976	704994	4266071	503	58	13.0	28.7	1.5	27.
102	Altura	Spain	662	704300	4407300	472	71	14.5	29.2	2.9	26.
103	Villa de Ves	Spain	864	651250	4338000	489	82	13.5	30.6	2.0	28.
104	Jarafuel	Spain	563	671850	4336700	518	71	14.7	30.5	3.5	27.
105	Bicorp	Spain	587	685200	4330500	545	76	15.1	30.5	3.2	27.
111	Benicasim	Spain	468	758019	4440766	699	90	14.2	26.3	4.4	21.
112	Gilet	Spain	152	727450	4394300	562	69	16.3	28.9	5.3	23.
131	Villajoyosa	Spain	126	735148	4264302	431	51	16.8	29.4	5.6	23.
141	Ricote	Spain	688	637500	4222800	321	44	15.8	32.6	3.5	29.
142	Monovar	Spain	820	678402	4250548	387	51	14.7	30.7	2.7	28.
143	Monovar	Spain	601	681100	4251500	335	47	15.8	31.2	3.7	27.
144	Paterna	Spain	1028	562950	4275800	552	67	13.2	32.2	-0.3	32.
145	Abaran	Spain	657	651900	4237900	343	45	15.6	32.5	2.4	30.
151	Quentar	Spain	1226	463550	4120300	657	29	13.0	31.6	-0.2	31.
152	Benamaurel	Spain	908	523020	4172800	394	44	14.3	32.9	1.1	31.
152	Velez Blanco	Spain	785	586700	4183100	357	37	14.2	32.6	1.1	31
155	Santiago de la Espada	Spain	842	546600	4231150	608	66	13.6	32.9	0.3	32.
154	Lorca	-	831	592647	4178631	329	42	15.7	32.9	2.7	29.
150		Spain Spain	765	628924	4191849	433	42 47	14.9	29.8	3.9	25.
157	Alhama de Murcia	Spain	703	498321	4191849	433 510	33	14.9	29.8 34.1	2.1	32
	Quesada	Spain								4.3	26
171	Lentegi	Spain	363	438677	4075439	593	16 25	16.0	30.7		
172	Carratraca	Spain	635	336500	4079000	774	35	14.3	30.4	2.9	27.
173	Frigiliana	Spain	595	417900	4075100	696	26	15.2	28.9	4.3	24.
182	Palma de Mallorca	Spain	32	1013545	4350235	554	38	16.8	28.7	6.9	21
183	Santanyí	Spain	19	1021743	4365904	568	40	16.8	28.7	6.9	21
184	Alcúdia	Spain	185	1027856	4431910	697	59	16.0	28.0	6.2	21
185	Calviá	Spain	243	971329	4396425	485	50	16.3	28.6	6.2	22
186	Alcotx	Spain	85	1112427	4449307	616	47	16.5	28.1	7.1	21
187	Atalix	Spain	67	1103104	4442252	606	45	17.0	28.5	7.5	21
191	Cala d'hort	Spain	329	868343	4312901	503	46	16.6	28.4	6.0	22
192	Ses Salines	Spain	10	881834	4308362	445	37	17.4	29.0	6.9	22.
193	Ses Salandres	Spain	65	874946	4331016	507	50	16.3	28.2	5.8	22.
211	Istaia-eyboia	Greece	53	2818134	4637783	506	35	17.4	32.5	5.3	27.
212	Amfilohia	Greece	429	2618397	4595421	966	66	14.8	31.1	2.1	29
213	Tatoi-attica	Greece	253	2824738	4603335	619	49	14.1	29.7	2.1	27.
214	Kassandra	Greece	402	2803882	4800501	518	63	14.1	29.2	1.5	27
221	Gemenos	France	391	1201790	4843798	712	102	12.4	25.9	1.0	24.
231	Litorale Tarantino	Italy	204	2206311	4696011	550	76	15.2	29.5	3.9	25
232	Gargano Monte Pucci	Italy	382	2073826	4816115	553	97	13.4	26.0	3.4	22
233	Gargano Marzini	Italy		2075777	4774891	472	76	16.1	30.7	4.5	26
241	Thala	Tunisia	948	1558095	3999027	463	63	14.9	34.6	1.0	33
242	Tabarka	Tunisia	287	1583691	4108559	553	36	17.8	36.1	4.5	31.
	AP = annual mean preci										

annual mean temperature (°C); MTWM = mean temperature of the warmest month (°C); MTCM = mean temperature of the coldest month (°C); CI = continentality index (°C).

32	TABLE S2. Vegetative and reproductive tra	ts in 52 natural Pini	is halepensis populations
33	grown in a common garden placed at Valdeoln	ios (Madrid, Spain), re	eferred as low stress site.
		$log(\mathbf{P}\mathbf{A})$	Slope*

II III a C	Jonnio	i garuen	placed at Va			a, span), tetene	<u>u as 10w sucss suc</u> .
	Н	Biomass	TSFR	log(ARI)	log(RA)	Intercept*	Slope*
Code	(cm)	(kg)	(cm)	(n cones)	(cones/kg	(cones)	log(cones/kg
	. ,		(em)	. ,	biomass)	. ,	biomass)
11	371±19	16.1±3.8		$2.9\pm0.2$	$-0.36\pm0.56$	6.19 (3.05-10.74)	0.043 (0.013-0.076)
21	323±21	$9.4 \pm 4.4$	221 (182-288)	$2.1\pm0.2$	$-0.04\pm0.64$	3.74 (2.08-8.25)	0.033 (-0.012-0.086)
31	342±20	$11.5 \pm 4.1$	202 (165-269)	2.0±0.2	-0.34±0.6	3.43 (1.68-6.36)	0.017 (-0.01-0.066)
61	325±20	$8.6 \pm 4.1$	149 (118-168)	3.2±0.2	$1.09\pm0.6$	10.61 (6.98-23.45)	0.058 (0.001-0.112)
82	357±19	$10.6 \pm 3.8$	220 (174-257)	2.3±0.2	-0.11±0.56	5.42 (2.75-9.08)	0.036 (-0.004-0.08)
83	348±20	$10.8 \pm 4.1$	174 (138-200)	2.4±0.2	$-0.02\pm0.6$	5.46 (2.93-10.37)	0.053 (-0.002-0.097)
84	$304\pm20$	$6.4\pm4.1$	116 (92-139)	3.5±0.2	$2.47 \pm 0.6$	17.39 (9.47-31.05)	0.078 (0.009-0.147)
91	338±22	9.2±4.5	172 (147-206)	3.6±0.2	$1.03 \pm 0.66$	13.89 (6.3-25.96)	0.068 (0.026-0.144)
92	294±20	$5.4 \pm 4.2$	202 (144-240)	1.5±0.3	$0.02 \pm 0.62$	2.54 (1.18-4.73)	0.015 (-0.035-0.102)
93	366±20	$14.0\pm4.2$		3.0±0.2	$0.19 \pm 0.62$	7.31 (3.24-11.83)	0.049 (0.009-0.07)
101	318±19	9.3±4.0	211 (193-248)	$2.5\pm0.2$	$0.65 \pm 0.58$	6.13 (3.64-11.47)	0.048 (0.000-0.087)
102	337±20	$13.4 \pm 4.1$	234 (182-276)	$2.2\pm0.2$	$-0.17\pm0.6$	4.28 (2.48-8.58)	0.025 (-0.007-0.062)
103	315±19	8.7±3.8	190 (131-263)	$2.7\pm0.2$	$0.49 \pm 0.56$	4.99 (2.99-9.75)	0.055 (0.011-0.096)
104	333±19	9.1±3.8	172 (169-176)	$2.6\pm0.2$	$0.65 \pm 0.56$	6.07 (3.1-10.48)	0.038 (-0.003-0.079)
105	336±20	$11.5 \pm 4.2$	205 (163-246)	$2.8\pm0.2$	$0.24 \pm 0.62$	6.55 (3.17-12.08)	0.037 (0.005-0.081)
111	325±20	7.3±4.2	212 (169-268)	1.7±0.2	$-0.45\pm0.62$	3.23 (1.44-5.77)	0.029 (-0.031-0.082)
112	365±20	$32.9 \pm 4.2$	312 (238-502)	$2.0\pm0.2$	$-1.34\pm0.62$	0.63 (0.28-1.71)	0.002 (-0.017-0.011)
131	331±20	8.1±4.2	236 (184-286)	2.1±0.2	$-0.34\pm0.62$	3.13 (1.47-6.94)	0.04 (-0.026-0.089)
141	308±20	$6.6 \pm 4.2$	215 (165-271)	2.3±0.2	$-0.12\pm0.62$	4.95 (2.23-9.16)	0.035 (-0.019-0.11)
142	345±19	$11.6 \pm 3.8$	209 (84-334)	2.1±0.2	$-0.43\pm0.56$	3.27 (1.68-5.77)	0.02 (-0.029-0.049)
143	311±22	8.1±4.5	236 (184-272)	$1.9\pm0.2$	$-0.02\pm0.66$	3.05 (1.49-7.41)	0.023 (-0.045-0.07)
144	379±21	$23.4 \pm 4.4$	180 (178-199)	3.2±0.2	$0.05 \pm 0.64$	7.26 (2.84-12.57)	0.028 (0.004-0.052)
145	383±19	$15.2 \pm 4.0$	150 (105-192)	3.0±0.2	$0.03\pm0.58$	6.67 (3.76-11.99)	0.046 (0.013-0.077)
151	301±24	$7.8\pm5.0$		3.5±0.2	$1.86\pm0.73$	13.15 (7.45-38.19)	0.06 (-0.004-0.135)
152	318±19	$8.6 \pm 4.0$	137 (94-160)	3.1±0.2	$1.35\pm0.58$	9.08 (5.92-19.59)	0.059 (0.011-0.095)
153	319±20	$7.7 \pm 4.1$	219 (177-259)	$2.2\pm0.2$	$0.23\pm0.6$	4.84 (2.39-8.67)	0.026 (-0.02-0.099)
154	343±19	$14.3 \pm 4.0$	173 (150-198)	$2.9\pm0.2$	$0.42\pm0.58$	6.33 (3.12-10.81)	0.038 (0.015-0.076)
156	328±19	$7.6 \pm 4.0$	177 (159-202)	$2.4\pm0.2$	$0.58 \pm 0.58$	5.63 (2.84-10.11)	0.051 (-0.022-0.09)
157	$349\pm20$	$16.2 \pm 4.1$	152 (129-181)	3.0±0.2	$0.34 \pm 0.6$	6.65 (3.04-10.91)	0.032 (0.011-0.063)
158	353±19	$13.2 \pm 4.0$	131 (123-142)	3.5±0.2	$0.83 \pm 0.58$	13.14 (7.04-22.96)	0.075 (0.038-0.117)
171	364±20	$14.4 \pm 4.1$	217 (184-263)	$2.5\pm0.2$	$-0.22\pm0.6$	3.96 (2.11-8.83)	0.022 (-0.004-0.066)
172	$326 \pm 20$	$8.7 \pm 4.1$	132 (106-159)	3.4±0.2	1.17±0.6	13.83 (6.78-21.65)	0.074 (0.023-0.129)
173	327±20	$8.0 \pm 4.1$	209 (169-241)	$2.4\pm0.2$	$0.08 \pm 0.6$	5.07 (3.1-10.42)	0.047 (-0.001-0.105)
182	339±20	13.3±4.1	214 (145-275)	1.8±0.2	-0.85±0.6	2.72 (1.3-5.02)	0.004 (-0.026-0.047)
183	346±25	$10.9 \pm 5.2$	272 (217-331)	2.0±0.3	$-0.58\pm0.77$	3.23 (1.09-8.58)	0.021 (-0.035-0.082)
184	350±20	9.3±4.2	251 (168-333)	$1.8\pm0.2$	$-0.47\pm0.62$	2.8 (1.59-6.2)	-0.014 (-0.053-0.062)
185	341±20	$16.9 \pm 4.2$	209 (206-218)	2.1±0.2	$-0.51\pm0.62$	2.22 (1.17-5.22)	0.023 (-0.006-0.045)
186	315±22	$8.6 \pm 4.5$	210 (145-296)	1.6±0.3	$-0.14 \pm 0.66$	2.68 (1.52-6.78)	0.033 (-0.046-0.082)
187	361±20	$15 \pm 4.2$	165 (82-217)	$2.8\pm0.2$	$0.14 \pm 0.62$	5.54 (2.58-10.55)	0.034 (0.000-0.066)
191	346±21	$10.5 \pm 4.4$	264 (207-318)	1.7±0.3	$-0.75\pm0.64$	2.74 (1.33-5.82)	0.011 (-0.045-0.061)
192	329±22	$10.8 \pm 4.5$	248 (194-307)	$1.6\pm0.3$	$-0.82 \pm 0.66$	2.18 (0.99-5.01)	0.018 (-0.020-0.060)
193	354±26	$12.8 \pm 5.6$	243 (201-321)	1.9±0.3	$-0.5\pm0.83$	3.01 (0.93-9)	0.023 (-0.035-0.083)
211	359±23	$15.2 \pm 4.7$	290 (242-362)	$2.5\pm0.2$	-0.76±0.69	2.77 (1.44-8.06)	0.038 (-0.010-0.070)
212	394±21	31.1±4.4	334 (248-496)	1.3±0.3	$-1.88\pm0.64$	1.03 (0.4-2.1)	-0.014 (-0.034-0.01)
213	378±19	$26.0 \pm 3.8$	169 (95-238)	$2.5\pm0.2$	-0.53±0.56	3.62 (1.67-6.03)	0.015 (-0.006-0.03)
214	$328\pm20$	$10.8 \pm 4.2$	254 (218-310)	1.7±0.2	-0.6±0.62	2.22 (1.09-4.76)	0.013 (-0.048-0.051)
221	379±20	$14.5 \pm 4.1$	269 (161-397)	$1.8\pm0.2$	-0.91±0.6	2.38 (1.11-5.23)	0.008 (-0.028-0.042)
231	$332\pm22$	13.1±4.5	166 (119-207)	4.0±0.2	1.31±0.66	16.76 (8.62-36.74)	0.074 (0.034-0.108)
232	366±21	$15.0{\pm}4.4$	235 (182-308)	3.1±0.2	$0.21 \pm 0.64$	5.88 (3.17-12.77)	0.047 (0.013-0.079)
233	325±21	$10.7 \pm 4.4$	189 (136-224)	3.5±0.2	$0.36\pm0.64$	6.51 (2.89-12.98)	0.076 (0.024-0.118)
241	323±21	$7.0\pm4.4$	134 (73-176)	4.5±0.1	$2.62\pm0.64$	42.63 (17.92-79.1)	0.078 (0.026-0.186)
242	$314\pm24$	$7.0\pm5.0$	140 (118-156)	4.6±0.2	2.5±0.73	44.23 (15.91-91.17)	0.074 (0.017-0.173)
* Inter	rcept and s	slope were	derived from inc	lependent n	nodels		

\* Intercept and slope were derived from independent models

35 TABLE S3. Vegetative and reproductive traits in 52 natural *Pinus halepensis* populations

- 36 grown in a common garden placed at Rincón de Ademuz (Valencia, Spain), referred as high
- 37 stress site.

	Н	Biomass	TSFR	log(ARI)	log(RA)	Intercept*	Slope*
Code	(cm)	(kg)	(cm)	(n cones)	(cones/kg	(cones)	log(cones/kg
			(ciii)	(ii colles)	biomass)	. ,	biomass)
11	$281 \pm 18$	$4.2 \pm 1.4$	165 (156-174)	2.4±0.2	$1.57 \pm 0.38$	5.73 (3.41-9.09)	0.056 (0.015-0.131)
21	319±18	6.5±1.4	176 (156-231)	2.4±0.2	$1.24\pm0.38$	6.74 (3.82-9.51)	0.047 (-0.006-0.092)
31	288±19	4±1.5	149 (110-184)	$2.5\pm0.2$	1.81±0.4	8.57 (4.68-12.4)	0.087 (0.01-0.159)
61	274±17	$4.5 \pm 1.4$	118 (105-152)	3.0±0.1	$2.05 \pm 0.36$	9.03 (5.9-14.33)	0.113 (0.045-0.16)
82	$260 \pm 18$	$2.8 \pm 1.5$	122 (93-157)	$2.2\pm0.2$	2.12±0.39	5.67 (3.14-8.57)	0.08 (-0.003-0.156)
83	257±19	2.3±1.5	110 (92-165)	$2.6\pm0.2$	$2.15\pm0.4$	8.8 (5.49-14.2)	0.088 (0.005-0.179)
84	250±18	4.1±1.5	89 (69-118)	3.1±0.1	$2.99 \pm 0.39$	9.03 (6.11-15.62)	0.049 (0.018-0.108)
91	260±21	2.7±1.7	127 (92-136)	2.4±0.2	$2.15\pm0.45$	6.38 (3.72-11.46)	0.085 (-0.013-0.168)
92	$272 \pm 18$	$3.2 \pm 1.4$	170 (146-172)	$2.5\pm0.2$	$1.72\pm0.38$	6.67 (4.48-10.74)	0.087 (0.022-0.176)
93	284±19	4.6±1.5	225 (197-251)	2.3±0.2	$1.47\pm0.4$	5.69 (3.25-9.04)	0.031 (-0.043-0.085)
101	296±18	5±1.5	161 (155-167)	$2.7\pm0.2$	$1.66 \pm 0.39$	8.71 (4.8-11.91)	0.071 (0.021-0.133)
102	$280 \pm 18$	4±1.5	139 (117-170)	2.6±0.2	$2.07\pm0.39$	7.02 (4.95-12.3)	0.068 (0.014-0.146)
103	$269 \pm 20$	3.8±1.6	121 (80-167)	2.7±0.2	2.13±0.43	7.86 (4.88-14.07)	0.107 (0.029-0.165)
104	288±22	3.7±1.8	121 (76-155)	2.9±0.2	$1.9\pm0.48$	10.15 (5.46-17.53)	0.084 (0.017-0.188)
105	286±17	4.9±1.3	135 (102-168)	3.1±0.1	1.93±0.35	11.06 (7.3-15.66)	0.076 (0.026-0.124)
111	293±20	4.5±1.6	134 (122-155)	2.3±0.2	$1.29 \pm 0.43$	6.34 (3.38-9.57)	0.065 (0.001-0.136)
112	243±19	2.6±1.5	162 (122-164)	2.1±0.2	2.42±0.4	6.29 (3.58-9.45)	0.083 (-0.017-0.142)
131	248±22	$2.4{\pm}1.8$	147 (143-190)	2.3±0.2	$2.15\pm0.48$	7.14 (3.14-11.15)	0.072 (-0.011-0.168)
141	307±20	6.8±1.6	120 (83-166)	3.1±0.1	$1.65 \pm 0.42$	10.88 (5.46-15.11)	0.099 (0.046-0.149)
142	284±17	3.7±1.3	169 (127-205)	2.6±0.1	1.79±0.35	6.78 (4.77-10.35)	0.069 (0.015-0.147)
143	259±18	3.4±1.4	132 (108-162)	2.9±0.1	$2.14\pm0.38$	8.96 (6.04-14.55)	0.062 (-0.017-0.118)
144	270±17	3.4±1.3	128 (111-166)	2.2±0.2	$1.5\pm0.35$	5.24 (3.3-7.88)	0.089 (0.012-0.137)
145	268±16	3.7±1.3	131 (115-157)	2.7±0.1	$2.04\pm0.35$	8.75 (5.41-12.72)	0.081 (0.026-0.145)
151	299±22	$5.4 \pm 1.8$	119 (68-176)	3.9±0.1	$2.09\pm0.48$	15.32 (8.3-26.98)	0.1 (0.057-0.193)
152	258±20	2.9±1.6	114 (89-148)	2.8±0.2	2.3±0.43	8.97 (5.66-17.11)	0.091 (0.015-0.175)
153	261±18	$4.7 \pm 1.4$	160 (113-203)	1.9±0.2	1.31±0.38	3.23 (1.89-5.54)	0.037 (-0.008-0.08)
154	258±18	2.8±1.5	106 (70-135)	2.9±0.1	2.5±0.39	11.67 (6.89-17.87)	0.108 (0.033-0.194)
156	254±17	3.5±1.4	100 (85-124)	2.6±0.2	2.37±0.37	7.25 (5.02-12.25)	0.077 (0.012-0.145)
157	264±20	3.7±1.6	120 (100-143)	2.7±0.2	$2.26\pm0.42$	8.12 (4.44-13.14)	0.089 (0.02-0.156)
158	255±19	2.7±1.5	96 (83-121)	2.8±0.2	2.17±0.4	7.34 (4.89-12.63)	0.094 (0.015-0.181)
171	273±19	3.3±1.5	145 (120-176)	2.4±0.2	1.91±0.4	5.97 (3.8-10.78)	0.071 (0.017-0.157)
172	256±20	3.7±1.6	113 (96-143)	3.5±0.1	$2.72\pm0.42$	13.19 (9.83-26.95)	0.098 (0.045-0.186)
173	278±17	$4.1 \pm 1.4$	147 (128-174)	2.7±0.2	1.81±0.37	7.94 (4.73-11.6)	0.087 (0.022-0.151)
182	247±18	$2.4{\pm}1.5$	166 (146-174)	2.2±0.2	2.18±0.39	5.55 (4.1-10.11)	0.064 (-0.004-0.157)
183	259±22	$2.9 \pm 1.8$	176 (143-224)	20±0.2	1.57±0.48	4.04 (2.56-9.3)	0.067 (-0.01-0.16)
184	267±25	3.6±1.9	187 (93-216)	2.2±0.3	$1.83 \pm 0.54$	5 (2.55-12.37)	0.076 (-0.008-0.164)
185	291±20	4.7±1.6	148 (129-249)	2.7±0.2	1.47±0.43	6.46 (4.06-11.78)	0.094 (0.006-0.144)
186	244±18	2.2±1.5	118 (79-154)	2.4±0.2	2.74±0.39	8.65 (5.05-12.74)	0.093 (0.005-0.179)
187	237±22	2.3±1.8	105 (98-116)	3.0±0.2	2.91±0.48	9.9 (5.24-17.1)	0.089 (0.008-0.191)
191	314±22	6.2±1.8	226 (176-286)	2.1±0.2	1.15±0.48	4.57 (2.44-8.6)	0.024 (-0.036-0.09)
192	288±23	4.3±1.8	256 (195-330)	1.9±0.3	1.26±0.51	4.32 (2.15-8.57)	0.059 (-0.028-0.138)
193	292±18	3.7±1.5	218 (178-225)	1.9±0.2	$1.26\pm0.39$	4.51 (2.73-7.24)	0.058 (-0.039-0.115)
211	323±22	9.8±1.8	209 (203-224)	3.5±0.2	1.7±0.48	7.5 (5.24-16.55)	0.059 (0.023-0.099)
212	336±22	11.1±1.8	210 (177-222)	2.8±0.2	$1.04\pm0.48$	4.75 (2.28-7.93)	0.042 (-0.009-0.067)
212	300±23	6±1.8	139 (104-193)	3.0±0.2	1.84±0.51	7.84 (4.81-16.24)	0.074 (0.009-0.129)
213	292±25	3.9±1.9	283 (170-293)	2.1±0.3	1.57±0.54	6.27 (2.84-11.72)	0.068 (-0.015-0.149)
214	292±25 283±17	4.3±1.4	167 (142-202)	2.1±0.3 2.1±0.2	1.42±0.34	4.82 (3.10-7.57)	0.056 (-0.014-0.116)
231	278±17	4.2±1.3	120 (88-147)	3.4±0.1	2.59±0.35	15.87 (11.03-24.33)	0.118 (0.061-0.191)
232	256±18	2.6±1.4	116(86-143)	3.0±0.1	2.94±0.38	10.68(6.93-16.9)	0.100(0.038-0.200
232	304±19	5.8±1.5	135(111-190)	3.5±0.1	1.90±0.4	12.93(8.22-21.21)	0.113(0.066-0.174)
233 241	$304\pm19$ 249±21	$2.6\pm1.7$	63(33-87)	4.2±0.1	3.69±0.45	27.69(16.75-51.32)	0.134(0.032-0.228)
241	$243\pm26$	2.8±2.1	05(55 07)	4.2±0.1 3.7±0.2	3.68±0.59	14.11(5.61-28.13)	0.084(0.002-0.188)
			derived from in			1.11(3.01 20.13)	5.00 ((0.002 0.100)
inter	cept and	stope were		acpendent I	noucis		

38 39 TABLE S4 Pearson correlation coefficients for the correlation at the population level between climatic variables and *Pinus halepensis* traits in each site, the low stress site above diagonal and the high stress site below the diagonal. N = 52 populations. Values for climatic variables are the same for both trial sites as they refer to the environmental conditions found in the area where populations where sampled. Pale blue site-to-site correlations lie on the diagonal. Pale red, ecotypic trends with growth traits. Mid – red, ecotypic trends with reproductive traits. Red, trade-offs between reproduction and growth. Green, correlations with reproductive efficiency. Significant (p-value <0.05) correlations are indicated in boldface.

45

								Biom	Н	TSFR	$ARI_b$	logRA	ARI	slope	RE		
CI								-0.15	-0.22	-0.48	0.38	0.50	0.36	0.34	0.01		
									Alt	0.40	0.25	0.32	-0.17	-0.25	-0.02	-0.12	0.28
									Long	0.23	0.25	0.26	-0.11	-0.20	-0.03	-0.07	0.24
									Lat	0.15	0.27	0.09	0.01	-0.15	0.03	-0.04	-0.10
									AP	0.04	0.17	-0.01	0.04	-0.09	0.09	0.08	0.14
									PDQ	0.10	0.04	0.30	-0.30	-0.31	-0.27	-0.29	-0.04
									ATM	0.03	-0.13	-0.52	0.36	0.39	0.40	0.32	0.18
									TMWM	0.06	0.05	0.41	-0.41	-0.40	-0.38	-0.36	-0.12
	CI	Alt	Long	Lat	AP	PDQ	ATM	TMWM	TMCM	-0.02	-0.10	-0.53	0.45	0.46	0.45	0.40	0.17
Biom	0.08	0.15	0.13	-0.04	0.02	-0.08	-0.03	-0.08	0.03	0.28	0.81	-0.03	-0.12	-0.40	0.17	-0.21	-0.01
Н	0.03	0.20	0.23	0.06	0.14	-0.12	-0.21	-0.07	-0.07	0.87	0.14	0.12	-0.18	-0.55	0.07	-0.25	-0.12
TSFR	-0.38	0.28	0.19	-0.01	0.02	0.23	-0.31	0.23	-0.30	0.24	0.43	0.73	-0.74	-0.69	-0.72	-0.67	-0.33
ARI <sub>b</sub>	0.19	0.18	-0.05	0.08	-0.10	-0.07	0.29	-0.13	0.24	0.08	-0.04	-0.55	0.75	0.84	0.88	0.82	0.76
logRA	0.02	0.09	-0.11	0.01	-0.17	0.12	0.32	0.04	0.15	-0.54	-0.71	-0.68	0.63	0.71	0.67	0.71	0.60
ARI	0.27	0.13	-0.11	0.02	-0.14	-0.06	0.37	-0.15	0.30	0.20	0.02	-0.60	0.94	0.59	0.71	0.83	0.82
slope	0.12	0.11	0.03	0.03	-0.01	-0.07	0.17	-0.08	0.14	-0.22	-0.21	-0.59	0.66	0.49	0.60	0.50	0.74
RE	-0.04	0.52	0.27	-0.02	-0.09	-0.03	-0.02	-0.07	0.03	0.08	0.17	0.47	0.12	-0.03	0.06	0.02	0.20

CI, Continentality Index; Alt, Altitude; Long, Longitude; AP, Annual mean Precipitation; PDQ, Precipitation during the driest quarter; ATM, Annual mean Temperature; TMWM, Mean Temperature of the Warmest Month; MTCM, Mean Temperature of the Coldest Month. Biom, biomass; H height; TSFR, Threshold Size for First Reproduction (female); ARI<sub>b</sub> Accumulated Reproductive Investment after accounting for plant size (biomass); logRA logReproductive Allocation; ARI, Accumulated Reproductive Investment; slope, slope associated with biomass in the Reproductive-Vegetative size relationship (Poisson model); RE, Reproductive Efficiency.

# APPENDIX IV

Correlated effects on reproductive traits derived from domestication of a forest tree are relevant for future forest resilience. **Santos-del-Blanco L**, Alía R, González-Martínez S, Sampedro L, Lario F, Climent, J. 2013.

1	Correlated effects on reproductive traits derived from domestication of a forest tree are
2	relevant for future forest resilience
3	
4	Santos-del-Blanco, L. <sup>1,2,3</sup> , Alía, R. <sup>1,2</sup> , González-Martínez, S.C. <sup>1,2</sup> , Sampedro <sup>4</sup> , L., Lario,

5 F. <sup>5</sup> , Climent, J	J.	1,2
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- <sup>6</sup> <sup>1</sup> Department of Forest Ecology and Genetics. INIA-CIFOR, Ctra A Coruña km 7.5, E-
- 7 28040, Madrid, Spain.
- 8 <sup>2</sup> Sustainable Forest Management Research Institute, www.sostenible.palencia.uva.es
- <sup>3</sup>Department of Ecology and Evolution. University of Lausanne. Biophore Building.
- 10 CH-1015, Lausanne, Switzerland.
- <sup>4</sup> Misión Biológica de Galicia CSIC. Carballeira, 8, E-36143, Salcedo, Pontevedra,

12 Galicia, Spain.

- <sup>5</sup> Vivero de Maceda. Dirección Técnica. TRAGSA. Ctra.cesa Maceda-Valdrey km 2,
- 14 E-32700, Maceda, Ourense, Spain.
- 15
- 16 Corresponding autor: Santos-del-Blanco, L., Department of Ecology and Evolution.
- 17 Biophore Building. University of Lausanne. CH-1015. Lausanne, Switzerland Phone.
- 18 +41(0)216924247, e-mail address Luis.SantosdelBlanco@unil.ch
- 19
- 20 Running title: Correlated effects of domestication in trees
- 21
- 22 BIOLOGICAL SCIENCES: Agricultural sciences

23

## 24 Abstract

25 Trade-offs between growth and reproduction are well established across the tree of life. 26 According to allocation theory, artificial selection aimed at increasing timber yield in 27 forest trees is expected to indirectly alter life-history reproductive traits, diverting 28 resources from reproduction to growth. This hypothesis was tested in a long term 29 artificial selection experiment with *Pinus pinaster* (Maritime pine), a widespread forest 30 tree in its early stages of domestication. Growth and reproductive traits of progenies 31 from phenotypically selected individuals were compared with those of progenies from 32 control trees, not subjected to selection within the same population. Female and male 33 threshold sizes for reproduction were much higher -up to above one meter- and 34 reproductive investment relative to size for female and male functions were lower in the 35 selected group compared with the control group. A single event of artificial selection 36 proved sufficient to produce a significant change in traits fundamental for individual 37 fitness and for population persistence. Importantly, the direction of this change towards 38 delayed and reduced reproduction, opposes natural selective pressures derived from 39 climate change. Widespread plantation of genetically growth-improved forest trees 40 should therefore be done with caution, as they may constraint adaptive responses to 41 climate change in natural or naturalised future forests.

42

Key Words: fitness traits, genetic change, adaptation, artificial selection, domestication
syndrome.

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### 47 Introduction

48 Domestication of forest trees is still in its infancy compared to cultivated crops 49 (1). The most advanced breeding programs have been running for only a few 50 generations, and genetic breeding tools are still to be implemented (2). However, 51 domestication of many widespread tree species is growing, as it is the area of planted 52 forests in response to a rising demand of timber and cellulose (3). Managed forests 53 occupy large extensions providing goods and services with an ecological relevance 54 transcending the landscape scale. They constitute natural or naturalized ecosystems 55 where improved tree genetic materials often interact with natural un-improved 56 populations. As key pieces determining structure and function or extensive terrestrial 57 ecosystems (4, 5), even slight changes in the adaptive ability of individual trees can 58 have broad consequences. The incipient domestication of forest trees is probably 59 interfering with their ability to adapt to the environment, but how and how much is 60 currently unknown.

61 Common target traits in forest tree breeding are growth, timber yield, stem form 62 and physical and chemical wood properties (6, 7). Unintended correlated responses to selection, already predicted by Darwin (8), are rarely reported in the forestry literature 63 making it difficult to define a "domestication syndrome" for forest trees (9). Correlated 64 65 responses to selection or unintended effects of management are well defined for other 66 and fish-farming have provided several paradigmatic organisms. Fisheries examples(10). For instance, over-exploitation of fisheries has driven the evolution of 67 68 slow growing and precocious individuals (11), revealing fundamental genetic 69 correlations between growth and reproductive life-history traits. Life-history traits lay at 70 the base of adaptive processes and are closely linked to individual fitness, effective

population size and thus to population persistence. For example, micro-evolutionary
changes of plant size at reproduction or reproductive phenology in response to climate
change are well documented (12, 13), and a global trend towards earlier reproduction
has been described (14).

75 Interestingly, monoecious forest trees allow the separate study of female and 76 male sexual functions in the same individuals. Female reproduction is typically more 77 costly in plants (15) and a different effect of domestication on male and female 78 reproduction could reflect differential costs of reproduction (16). Therefore, separating 79 investment in female and male reproductive functions, particularly at the onset of 80 reproduction, is key to understand constrains to adaptive evolution for tree life-history 81 traits. Size at maturity and reproductive allocation are key fitness traits defining 82 contrasting life histories in all kind of organisms (17). As those traits imply shifting 83 resource allocation between vegetative growth and reproduction, genetic breeding for 84 allocation to growth is predicted to impact them.

85 Specifically, based on allocation theory, a decreased investment in reproduction 86 would be expected if selection is focused solely on growth. Here we provide evidence for this hypothesis, for the first time in a forest tree, with a long term artificial selection 87 88 experiment of *Pinus pinaster* (Maritime pine), a monoecious conifer widespread in 89 southwestern Europe in its early stages of domestication. Growth and reproductive traits 90 of the progenies from phenotypically selected individuals were compared with those 91 from control trees, not subjected to selection within the same population, growing 92 together in a common garden in North Spain. Indirect effects of selection for high 93 allocation to stem growth delayed the ontogenetic program for reproduction in maritime 94 pine. More specifically, we tested the hypothesis that progenies from selected trees 95 would reproduce at larger sizes and/or produce fewer cones at a given size. Besides,

4

96 considering the expected greater cost in terms of growth for female reproduction, we
97 hypothesised that female function of progenies selected for high stem growth should
98 show a greater indirect change, as compared to male function.

99

100 **Results** 

101 Direct and indirect effects of domestication Progenies of trees selected for timber 102 yield (hereafter 'selected group') were on average 11.7 cm higher at age 10 than those 103 sampled at random from the wild base population (hereafter 'control group') (Table 1). 104 As expected, phenotypic selection on mother trees lead also to better stem form scores 105 in the selected than in the control group (P < 0.001; Table 1). Trees from the selected 106 group were consistently taller than those in the control group for juvenile, vegetative, 107 female and cosexual groups (all P < 0.05; Fig. 1). However, height of male trees did not 108 significantly differ between groups and showed large variability within groups (Fig. 1, 109 Table S1).

110 Selection for timber yield established large differences in reproductive traits 111 between the selected and control group, leading to delayed reproduction both in age and 112 size terms in the former (Table 1). The proportion of reproductive trees was higher in the control group (66%) than in the selected one (51%) ( $X_{1}^{2}$  14.3 P < 0.001). In both 113 114 groups, trees started their reproductive phase most commonly as females, followed by 115 cosexuals and then males (Table S1). Among reproductive individuals, the selected 116 group had a lower proportion of cosexuals (36%) ( $X_{1}^{2}$  13.2 P < 0.001) and higher proportion of males (24%) ( $X_{1}^{2}$  14.3 P < 0.007). compared to the control group (54%) 117 118 and 13% respectively) (Table S1). Differences in reproductive traits between both 119 selected and control groups were highly significant (p<0.001, Table 1). The selected 120 group had significantly greater female and male threshold size for first reproduction

121 (TSFR), and showed a consistently lower allocation to reproduction at a given size 122 (reproductive – vegetative allocation, female and male R-V; Table 1). The maximum 123 change in TSFR was recorded for the female function, which increased in 106.9 cm in height in the selected population (Fig. 2). In the control group, female TSFR was 124 significantly lower than male TSFR (Fig. 2, Table S1), but both parameters did not 125 126 differ in the selected group as selection had a greater effect on female TSFR (Fig. 2, 127 Table S1). The control group showed the smallest reproductive individual and 128 significantly lower male TSFR than any of the 31 selected progenies. Regarding the 129 female TSFR, all but two selected families showed point estimates above the control, 130 but wider credible intervals made those differences non-significant for about half of the 131 selected families (Fig. S1).

132

## 133 Quantitative genetic parameters of growth and reproductive traits

134 Models were improved by including genetic relatedness for height ( $\Delta$  Deviance 135 Information Criterion, DIC 7.3). Additive genetic variance for height was very low 136 compared to total variance. Thus its heritability was centred at zero, although with wide 137 credible intervals (Table 2). Heritability for stem form was 0.13 and significantly 138 different from zero. Male and female reproductive functions expressed sharp differences 139 in genetic control between selected and control groups. While narrow sense 140 heritabilities for female TSFR and relative reproductive investment (R-V) were very 141 high (> 0.53), those for male function were indistinguishable from zero (Table 3).

In the selected group we found significant negative genetic correlations between female TSFR and female relative reproductive investment (r = -0.82; Table S2a) indicating that precocious trees were also more prolific at this stage. Male and female thresholds for first reproduction were positively correlated (r = 0.39) but not the relative
investment in male and female functions (Table S2a). Within the 31 selected families, we only found evidence of genetic trade-offs between height and reproduction for male function, but not for female function (Table S2a). This showed that taller families tended to have a delayed male size for first reproduction (r = 0.38; Table S2).

150

# 151 **Discussion**

In this study we show how a single event of artificial selection aiming at domestication of a forest tree interfered with adaptive potential of a widespread conifer, *Pinus pinaster*. Our study is, to our knowledge, the first to report on the correlated effects in reproductive life-history traits caused by selection for increased timber yield in a forest tree.

157 In our experiment, phenotypic selection in natural stands pursued an ideotype of 158 tall, thick straight trees with short horizontal branches and high apical dominance. There 159 was a positive phenotypic correlation between growth and absolute female and male 160 reproduction (Table S2). However, due to "unconscious selection" (8) of traits 161 genetically correlated with high stem allocation, female and male threshold sizes for 162 reproduction were both increased and male and female reproductive allocation 163 decreased in the selected group, revealing an underlying negative genetic correlation 164 between allocation to growth and reproduction in the base population (18). Differences 165 in ontogenetic development were also reflected in a higher proportion of reproductive 166 individuals in the control group compared to the selected group. Considering actual 167 growth rates, two to three years would be needed for the selected group to achieve a 168 similar reproductive status as the control group. Thus indirect selection effects on male 169 and female reproductive traits caused delayed reproduction both in age and size terms.

Altogether, direct and indirect effects of selection contributed to define a "domesticationsyndrome" for timber selection in trees (19).

172 A few artificial selection experiments in plants have focused on reproductive 173 life-history traits, such as the threshold size for reproduction (20, 21) and flowering 174 phenology (22, 23), revealing abundant additive genetic variation for these traits. 175 Narrow-sense heritabilities reported in those studies are moderate, and close to the 176 lower bound of our estimates for female reproduction in *Pinus pinaster*. Numerous 177 studies in forest trees, in particular in conifers, also report high broad-sense heritabilities 178 for cone and pollen production in clonal seed orchards, usually in the range of 0.25-0.80 179 (24).

180 Female reproduction has been usually found to be more costly than male 181 reproduction (15). In agreement with this idea, we found smaller average size for 182 protogynous than protrandrious trees in maritime pine, indicative of higher female 183 reproductive costs (Fig 1). Despite we were not able to detect negative genetic 184 correlations between reproduction and growth in the selected group, our results 185 evidence that strong negative genetic association between growth and early reproductive 186 effort existed in the native original population subjected to selection. At the genetic 187 level, greater costs of female reproduction could be thought of as stronger negative 188 correlations for growth (25), thus, a more intense indirect change of the threshold size 189 for female reproduction, compared to the male threshold, fitted our expectations. The 190 change due to a single selection event implied an increase close to half of the species 191 range of the threshold size for female reproduction and similar to the species male range 192 of the threshold size for male reproduction (1.0 m female, 0.8 m male). Our results 193 partially support greater genetic costs of female reproduction, as the change in female 194 function was more important in absolute terms (greater change in female function), but195 not in relative terms (greater change in male function).

# 196 Implications for adaptation of forest trees and forest management in a changing 197 environment

198 The observed correlated genetic effects of breeding for timber production on 199 reproductive life-history traits are closely linked to individual fitness, but may also 200 impact effective population size at a given age, and thus on population "in situ" 201 persistence and resilience against disturbance. Climate warming has led to dramatic 202 increases of severe droughts and more frequent forest fires in many areas such as the 203 Mediterranean basin (26). Under forecasts of increased disturbance frequency, and thus 204 reduced life expectancy, precocious trees could be naturally selected for (27), as 205 mounting evidence indicates for non-woody species (12–14, 27). Indeed, under climate 206 change scenarios, both survival in situ and potential for migration and extended gene 207 flow rely on enhanced early allocation to reproductive function(28).

208 Present-day population differentiation along climate clines can also help us to 209 infer responses to changing climate. In that sense, genetically-based phenotypic clines 210 are common in forest trees (4). Particularly, poorer growth conditions of origin have 211 been related to earlier female reproduction in several pine species, including *P. pinaster* 212 (27, 28, 29), suggesting that populations facing increased environmental stress would 213 benefit from enhanced early reproduction. However, correlated genetic effects observed 214 after a single artificial selection event for increased timber yield lie in the opposite 215 direction of the putatively adaptive trend observed in natural populations evolving in the 216 wild. In a context of changing environment, and taking into account the long life cycles 217 and extensive pollen dispersal, intensive breeding of native forest species could be

compromising the adaptation of managed forest and also of natural surrounding forestsinteracting genetically with them.

220 The impact of forest management on genetic resources of forest trees is a 221 declared matter of concern for conservation practices (29). Silvicultural activities such 222 as selective logging (30), clear cuttings (31) and seed tree regeneration (32) have not 223 been not found to strongly impact, within limits, forest stands in terms of neutral genetic 224 diversity. Also, assuming a positive correlation between neutral and adaptive diversity, 225 production of genetically improved material in seed orchards ideally seeks a 226 maximization of genetic diversity as assessed by neutral markers (33). However, 227 breeding, by definition implies genetic changes that are reflected in phenotypes. The 228 fitness consequences of deployment of material improved for increased allocation to 229 growth (this study) or selective harvesting in natural tree stands (30, 34) can be of great 230 importance, as has also been demonstrated in fisheries and fish-farming (10). Indeed, 231 growth traits should not be considered alone as adaptive traits of forest trees (35, 36). 232 Due to long life cycles, extensive distribution areas and key environmental relevance, 233 reproductive traits and tolerance to biotic and abiotic stressors must be also taken into 234 account to assess adaptation of forest trees in the long term. An excessive focus on 235 growth traits might imply a correlated negative response in reproductive investment, 236 with detrimental consequences on forest regeneration and long-term survival, in 237 particular under environmental stressful conditions.

In spite of impressive recent development, with first full-genome draft sequences underway, molecular genetic tools are still not accurate enough to describe adaptive variation in forest trees, particularly in species with huge genomes like pines. Screening for differences between domesticated and wild populations of forest trees might be a powerful tool to unveil the genetic basis of adaptation to both natural and human

- environments (19) and further improve domestication (2). Meanwhile, quantitativegenetic tools cannot be dismissed for policy guidance in the sustainable management of
- 245 forest genetic resources aimed at ensuring their evolutionary potential.

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#### 383 Methods

#### 384 Studied species and selection experiment

*Pinus pinaster* is a Western Mediterranean monoecious conifer, with a disjoint distribution in South-Western France, Iberian Peninsula (Portugal and Spain), Italy and North Africa (Morocco, Algeria and Tunisia). A complex phylogeographic history has been postulated for this species based on different molecular markers (37), and significant within population variation has been reported for growth, wood properties and stem form (7), defence (38), and also for reproductive life-history traits (39).

391 A long term selection experiment aimed at increasing timber yield was carried 392 out in the 'Montaña de Soria-Burgos' provenance area, Central-North Spain, during the 393 1990's. Thirty one plus trees were selected in natural stands according to their superior 394 phenotypes for growth and stem form, i.e. timber production. Altogether, less than 1 out 395 of 10,000 trees were selected as plus trees. Then, in 2001, their progeny was planted in 396 a common garden together with the progeny from an unselected control lot randomly 397 selected within the same population. The common garden was placed in the vicinity of 398 the surveyed stands (Latitude 41° 55' 15" N; Longitude 3° 11' 35" W; 1,153 m a.s.l.; Fig. 399 3). A detailed description of the selection protocol and common garden design can be 400 found in the Supplementary Material (ST1).

401 A total of 1,087 trees were included in this study, 833 belonging to progenies of 402 selected trees (selected group) and 254 to progenies of unselected trees (control group). 403 Mortality at ten years was very low (2.4%) and did not differ between selected and 404 control groups ( $X^{2}_{1} = 0.266$ , P = 0.606). Some trees showing damage caused by insects 405 or pathogens were excluded from the study, and thus the final sample size was 868 406 trees.

# 407 Field measurements and variable description

408 Two measurements were carried out in late May in 2009 and 2011, when the trees were 409 eight and ten years old, respectively. In both dates, we measured tree height below the 410 elongating bud (this is, previous winter height) and diameter at breast height (for those 411 trees > 1.3 m) and female cones were counted for the following cohorts: female strobili 412 (red-coloured and placed at the end of the current year flush), one-year old immature 413 conelets and two-year old and above mature cones (SFig. 2). Male (pollen) cone 414 abundance was categorically scored from 0 (absent) to 3 (very abundant). Stem form 415 and branching habit was assessed through an ordinal scoring from 1 to 6 in 2009 (40). A 416 higher score implied high apical dominance and straight stem with few branches, 417 inserted at high angles (closer to horizontal position) (41). Total biomass was estimated 418 from diameter at breast height using standard formulae in Montero et al. (2005).

419 Considering its reproductive status, each tree was classified as either 420 reproductive or non-reproductive. According to their first reproductive event, trees were 421 assigned to female (F, those that started their reproductive life as females, i.e. 422 protogynous), male (M, those that started their reproductive life as males, i.e. 423 protrandious) or cosexual (C, those that started their reproductive life with both female 424 and male strobili) groups. Non-reproductive trees were further classified as juveniles (J, 425 size below the family or group-specific smallest reproductive individual) or vegetative 426 (V, size above the family or group-specific smallest reproductive individual).

427 <u>Statistical analyses</u>

We used Bayesian approaches for fitting quantitative genetic models as they provide more flexibility and accuracy in estimating errors of model parameters and their combinations, e.g. heritability, than Maximum Likelihood procedures (43). We computed posterior mode and 95% credible intervals (95% CIs) for fixed effects, variance components, threshold sizes for reproduction, heritabilities and phenotypic

433 correlations across a variety of models. A detailed description of priors used can be
434 accessed in the Supplementary Material (ST2). Analyses were run in R, version 2.15.2
435 (R Development Core Team, Vienna), using the package MCMCglmm, version 2.01
436 (44).

Tree height at ten years was modelled as normal distributed traits with Gaussian errors including block as a random factor. Tree form was modelled as an ordinal trait with a generalised model with probit link and residual variance ( $V_R$ ) fixed to one by convention. Stem form model included block as a random factor. Fixing  $V_R$  to a particular value affects proportionally all variance components in a model, and thus heritability estimates are independent of the  $V_R$  value.

443 Threshold size for first reproduction (TSFR) was calculated as minus the result 444 of dividing intercept by slope estimates from a generalised binomial model with logit 445 link, where categorical reproduction (present or absent) was the response variable. 446 Height was included as a covariate (20).  $V_R$  was fixed to one in binomial models by 447 convention. Cumulative quantitative female reproduction, i.e. number of cones 448 produced throughout a tree's life, was modelled as a Poisson generalised model with log 449 link, using log biomass as a continuous predictor (Female R-V). This was equivalent to 450 standard log-log allometric Reproductive vs. Vegetative size (R-V) regressions but 451 benefited from the inclusion of zeroes in the response variable (non-reproductive 452 individuals), otherwise commonly discarded or transformed (45). Only data from nonjuvenile trees were used in R-V models in order to avoid zero inflation (46). Male 453 454 reproduction at ten years, an ordinal trait, was modelled analogously to stem form, but 455 including tree log biomass as a covariate (Male R-V). Inclusion of size (height or 456 biomass) as a covariate in TSFR and R-V models accounts for small scale

457 environmental variation in common gardens (47), and so, block effects were not

458 considered.

#### 459 **Direct and indirect effects of early domestication**

We first tested for differences between progenies from the selected (hereafter "selected group") and non-selected (hereafter "control group") mother trees. We analysed height, stem form, threshold for first reproduction (TSFR), and reproductive-vegetative (R-V) size relationships for female and male reproduction by fitting independent univariate models and using fixed effects (selected – control) 95% credible intervals (95% CIs) to evaluate the significance of their difference. Average sizes per reproductive group (juvenile, vegetative, female, male or cosexual) were also calculated. For TSFR and R-

467 V models we evaluated the significance of separate additive (intercept) and

468 multiplicative (slope) selection effects.

# 469 Genetic control and correlation among traits

470 To evaluate the quantitative genetic basis of variation in traits affected by early 471 domestication, we fitted 'animal models' to data from the selected group to estimate 472 additive genetic variance. A detailed description of model specification and estimation 473 of quantitative genetic parameters can be accessed in the Supplementary Material (ST3). 474 As 95% CIs for variance components cannot overlap 0, it is not possible to test the null 475 hypothesis of zero variance. Instead, significance of variance components was assessed 476 by means of Deviance Information Criteria (DIC), comparing DIC values of nested 477 models (48). However, this was done only for Linear Mixed Models (height) as 478 interpretations of DIC in Generalised Linear Mixed Models with latent variables may 479 not be clear. In these cases (binomial, Poisson and ordinal models), parameter posterior 480 distribution was used to illustrate the magnitude of variance components (49).

481 Genetic correlations were estimated as the Pearson's correlation between traits for
482 family means derived from mixed models where family was coded as random (50).
483 Phenotypic correlations were estimated by fitting bivariate models to data from the
484 selected and control groups.

TABLE 1. Effects of a single artificial selection event aimed at improving growth and
timber yield in a *Pinus pinaster* wild population on growth and reproductive function of
the following generation. Selected and unselected groups were grown in a common
garden close to the original population in Central-North Spain. Effect size and 95%
credible interval, CI, are shown. For models where a covariate was used (height or
biomass), we indicate both add., additive (intercept) and mult., multiplicative (slope)
effects of selection. Results are reported on the corresponding latent linear scale.

494

Trait	Effect size	95% CI		p-value	
Height		11.7	1.9	27.7	0.024
Stem form		0.375	0.218	0.618	<0.001
Female TSFR	add. mult.	-1.242 -0.003	-1.452 -0.006	-0.893 0.001	< <b>0.001</b> 0.198
Female R-V	add. mult.	-1.627 0.123	-1.918 -0.128	-1.197 0.301	< <b>0.001</b> 0.45
Male TSFR	add. mult.	-1.616 -0.002	-1.957 -0.007	-1.272 0.003	< <b>0.001</b> 0.422
Male R-V	add. mult.	-0.886 -0.073	-1.278 -0.274	-0.605 0.176	<b>&lt;0.001</b> 0.764

495

TSFR, Threshold size for first reproduction; R-V, relative Reproductive – Vegetative
 effort obtained from a Poisson (female) or ordinal (male) model with number of cones

498 (female) or qualitative pollen production (males) as the response variable and

499 log(biomass) as a covariate

- 501 TABLE 2. Posterior modes  $(h^2)$  and credible intervals (95% CI) of narrow sense
- 502 heritabilities for growth and reproductive traits recorded on *Pinus pinaster* trees. Values
- 503 for stem form, TSFR and RV are reported on the corresponding latent linear scale.
- 504

trait	h <sup>2</sup>	95% CI		
Н	0.00	0.00	0.19	
Stem form	0.13	0.03	0.36	
Female TSFR	0.53	0.35	0.91	
Female R-V	0.73	0.42	0.81	
Male TSFR	0.00	0.00	0.24	
Male R-V	0.00	0.00	0.43	

505

TSFR, Threshold size for first reproduction; R-V, relative Reproductive – Vegetative
 effort obtained from a Poisson (female) or ordinal (male) model with number of cones

(female) or qualitative pollen production (males) as the response variable and vegetativesize as a covariate.





520 FIGURE 2. Comparison of height density distribution and threshold size (height) for 521 first reproduction between *Pinus pinaster* progenies of trees either selected (S) or not 522 selected (C, control) for timber production after one generation. Bell-shaped lines 523 represent height probability distribution at age 10 yrs. Control group, orange lines; 524 selected group, blue lines. Vertical lines represent threshold sizes for reproduction. 525 Solid line, female function; broken line, male function. 95% Credible intervals for 526 threshold sizes are represented by horizontal lines. Arrows show changes in threshold 527 size for reproduction in male and female function due to selection. Control group, left; 528 selected group, right.



FIGURE 3. Location of the *Pinus pinaster* region of provenance Montaña de SoriaBurgos (light green), and the forest stand (dark green) where artificial selection for
growth and timber yield was carried out. Boxes 1-3 describe the breeding program from
phenotypic selection to common garden establishment. Star denotes the location of the
common garden.



#### 1 Correlated effects on reproductive life-history traits derived from domestication of

## 2 a forest tree are relevant for future forest resilience

3 Santos-del-Blanco, L., Alía, R., González-Martínez, S.C., Sampedro, L., Lario, F.,

4 Climent, J.

5

# 6 SUPPLEMENTARY INFORMATION

#### 7 SUPPLEMENTARY TEXT 1. Phenotypic selection in the wild and sampling

8 A breeding program for the Pinus pinaster region of provenance Montaña de Soria-9 Burgos in Northern Spain (1) was designed in the 1990's. The first stage involved 10 phenotypic selection of plus trees growing in natural stands. Best valued traits were 11 high stem volume growth, straight stem, high apical dominance and short branches 12 inserted at high angles (closer to horizontal position) (2). Trees were selected between 13 1992 and 1995 by subjective comparison with neighbouring individuals. In order to 14 avoid biasing the selection of best phenotypes towards favourable sites, a geographical 15 subdivision of the stands was made, such that best scoring individuals of all 16 subdivisions were chosen for progeny testing. All individuals were above 28 years old 17 (at breast height), between 20 and 28 m tall and between 35 and 60 cm of diameter over 18 bark at breast height. Some candidate plus trees had to be discarded as they did not 19 produce enough seed for progeny testing. Considering the area of sampled stands and 20 the average tree density, less than 1 out of 10,000 trees were selected as plus trees. The 21 history of forest management in the area and the age of all mother trees allowed us to 22 discard seed transfers from unknown origins.

Ripe cones of 31 selected plus trees were collected in autumn 1998. Also, a control seed lot was built the same year from cones collected randomly from 30 trees felled during forest management activities. This ensured that control seed lot was representative of the average phenotype of the study stand, avoiding a frequent bias of commercial seed lots collected from standing trees, therefore over-representing highly reproductive mothers (3). The control seed lot was built in the same study area where plus tress were selected. This ensured that comparisons were not biased by fine-scale local adaptation and that pooled data from these 31 plus tree progenies gathered a priori a similar genetic variability than the control seed lot.

All cones harvested from selected and control trees were processed together, including their opening in an oven at 60° C after being air-dried. Seeds were extracted manually, and kept dry at 4°C until sowing in 2000. Seedlings were produced in 250 cc containers with a mixture of fertilized peat and vermiculite following best nursery practices for the species and planted in the common garden in 2001. Plants were placed at the intersections of a 3 x 3 m grid following a resoluble alpha design with 28 complete blocks and single-tree plots.

The common garden for progeny testing was set in a former marginal arable land, within the area where phenotypic selection was carried out (Figure 1). The plot is placed at 1.153 m above sea level, with approximately 10% of slope facing eastwards. The mother rock is siliceous sandstone and conglomerates, the predominant soil type in the 'Montaña de Soria-Burgos' area, but soil impoverishment and erosion from inadequate previous farming practices were evident.

#### 46 SUPPLEMENTARY TEXT 2. Prior specification for statistical models

47 Default priors for fixed effects followed a normal distribution centred around cero and had a large  $(10^8)$  variance. Non-informative diffuse proper priors were used for variance 48 components. By default, we used inverse-Wishart priors with parameter V = 1 and a 49 50 degree of belief of 0.002. However, given the null or very low value of some variance 51 components, parameter expanded priors were also used in order to improve chain 52 mixing. For height mixed model, the working parameter prior was normally distributed, 53 with a mean of 0 and a variance of 1000, and the location effect prior was inverse-54 Wishart distributed, forming a scaled non-central F distribution, with a degree of belief 55 parameter and a limit variance of one. In the case of binomial (male and female 56 threshold size for reproduction) and ordinal models (stem form and male reproduction) 57 residual variance was fixed to 1, causing inflation in heritability estimates due to high 58 additive genetic variances driven by long tails in prior distribution. In order to keep 59 additive genetic variance within expected boundaries, we used a Chi-square distributed 60 prior with one degree of freedom for variance components (4). All MCMC analyses 61 were run yielding in every case a total of 1,000 data points for each analysis and 62 ensuring autocorrelations below 0.1.

# 64 SUPPLEMENTARY TEXT 3. Model specification and estimation of quantitative

#### 65 genetic parameters

Animal models use pairwise coefficients of relatedness (0.25 between half-sibs) among individuals to define a matrix that is proportional to the variance-covariance structure of additive genetic effects. In these models, individual (animal) effects estimate the variance in phenotypes that can be related to additive genetic variance ( $V_A$ ). For stem form and TSFR generalised models, an additional individual effect was added to account for overdisperssion, i.e. individual variance ( $V_I$ ) larger than that specified by the model link due to causes other than additive genetic effects.

73 The heritability of normally distributed growth variables (height and biomass) 74 was calculated by dividing  $V_A$  by total variance ( $V_A$  + residual variance,  $V_R$ ). Latent 75 scale heritabilities can be interpreted as the heritability of a continuous, directly 76 immeasurable variable that describes a variation in a discontinuous trait. For example, 77 increasing continuous values of liability for reproduction are translated into two 78 phenotypes (reproduction absent or present) separated by a threshold. Heritability of tree form and male reproduction on the underlying (probit) latent scale, was calculated 79 as:  $h_{\text{form latent}}^2$ :  $V_A / (V_A + V_I + V_R + V_L)$ , and  $h_{RVm \text{ latent}}^2$ :  $V_A / (V_A + V_R + V_L)$ , respectively, 80 81 with  $V_R = 1$  by convention and latent scale variance  $V_L = 1$  as the probit variance. 82 Heritability of the threshold size for female and male reproduction was also calculated on the underlying (logit) latent scale as:  $h^2_{TSFR | atent|}$ :  $V_A / (V_A + V_I + V_R + V_L)$ , with  $V_R = 1$ 83 by convention and  $V_L = \pi 2/3$  as the logit variance. Heritability on the underlying 84 85 (Poisson) latent scale for female reproductive investment after accounting for size was calculated as: 86

87  $h_{RVf \, latent}^2$ :  $V_A / \{V_A + V_R + log[1/exp(\beta_0) + 1]\}$ , where  $\beta_0$  is the predicted trait value given 88 certain fixed effects (5). We calculated  $h_{RVf \, latent}^2$  taking  $\beta_0$  as the average cone 89 production per tree.

90 Genetic correlations were estimated as the Pearson's correlation between traits 91 for family means derived from mixed models where family was coded as random (6, 7). 92 Genetic correlations were only calculated for those variables with significant genetic 93 variance or heritability. Phenotypic correlations were estimated by fitting bivariate 94 models to data from the selected and control groups. Models were fitted with two 95 variables at each time given that there were slight differences in number of observations 96 per variable due to the existence of juveniles regarding male or female function. The 97 variables included were height, biomass, stem form and male (qualitative scale) and 98 female (cone counts) reproduction. Height and biomass were considered Gaussian 99 variables while quantitative female reproduction was treated as Poisson. Originally, 100 stem form and male reproduction were recorded according to an ordinal distribution, but 101 here were considered also as count data (Poisson error) in order to enable the calculation 102 of phenotypic correlations (8). Genetic and phenotypic correlations thus do not 103 represent the same set of traits as phenotypic correlations for male and female 104 reproductive data were not corrected for size effects.

SUPPLEMENTARY FIGURE 1. Effect of a single event of artificial selection for timber yield within a *Pinus pinaster* population on the threshold size for first reproduction of the progeny of selected plus trees and control, unselected, base population. Smallest reproductive individual (black dots), and threshold size for female (grey squares) and male first reproduction (empty squares plus) per family and control group are shown. Family or group mean ± 95% credible intervals are shown. Control, control group; 26P-52S codes, half-sib family codes; SW Female and Male: female and male Species-Wide ranges for the threshold size for first reproduction (reported by Santos-del-Blanco L, Climent J, González-Martínez SC, Pannell JR (2012) Genetic differentiation for size at first reproduction through male versus female functions in the widespread Mediterranean tree *Pinus pinaster*. Ann Bot 110:1449–1460)



SUPPLEMENTARY FIGURE 2. Description of *Pinus pinaster* seed cone development. Top left, current year female strobilus. Mid-left, first year female cone. Mid-right, second year mature female cone. Top right, whorls of serotinous female cones older than two years attached to the main stem.



116 SUPPLEMENTARY TABLE 1. Posterior modes (mode) and credible intervals (95% CI) of height (cm) per reproductive class (juvenile,

117 vegetative, female, male or cosexual) and experimental group (selected, control) in *Pinus pinaster* trees. TSFR indicates the median threshold

118 size for first reproduction per sex. Cells marked with an asterisk (\*) indicate significant (P<0.05) differences between selected and control

119 groups.

120

				selected			conti	ol	
	Reproductive class	n	%	mode	95% CI	n %	6 mode	95% CI	sig.
Height	Juvenile	167		165.4	156.1 184.1	15	100.0	67.2 137.3	*
	Vegetative	155		268.6	251.7 280.2	51	199.7	182.5 221.2	*
	Female	134	40	285.1	264.7 294.1	42 33	3 237.6	213.4 254.5	*
	Male	81	24	316.8	306.6 340.2	17 13	305.3	272.5 331.9	
	Cosexual	119	36	346.3	334.3 365.2	70 54	308.9	292.0 325.5	*
	Total	656	51			195 60	ó		
TSFR	Female			322.6	293.0 362.4		215.7	187.1 241.7	*
	Male			323.9	310.6 338.2		246.5	230.5 267.7	*

121

122 n, number of trees per class. Total, total number of trees per group and age. %, referred to F, M and C represents the percentage of those classes

123 respect to reproductive trees. % referred to total represent the percentage of reproductive trees. Pines are monoecious species and individual trees

124 usually produce both female and male cones, in our study, Female and Male trees represent protogynous and protrandrious individuals.

125 Vegetative trees where those that could reproduce according to their size, but they did not.

126 SUPPLEMENTARY TABLE 2. Genetic and phenotypic correlations between vegetative and

127 reproductive traits measured in *Pinus pinaster* trees grown in a common garden in North Spain.

128 Significant correlations (P< 0.05) in bold face.

**a**) Half-sib family correlations (above diagonal) and corresponding P-values (below diagonal)

130 among traits in the selected group for timber production.

		Female		Ma	le	Vegetative	
		TSFR	R-V	TSFR	R-V	Height	Stem form
Female	TSFR		-0.82	0.39	-0.19	0.12	-0.38
Fer	R-V	< 0.001		-0.52	0.00	-0.14	0.25
ile	TSFR	0.03	0.00		0.02	0.38	-0.31
Male	R-V	0.31	0.99	0.90		0.06	0.02
Vegetative	Height	0.51	0.45	0.04	0.74		-0.36
Veg	Stem form	0.04	0.17	0.09	0.92	0.05	

131 TSFR, Threshold size for first reproduction; R-V, reproductive investment relative to size.

- 132
- 133 b) Phenotypic correlations among raw traits for the selected (above diagonal) and control,

134 unselected, (below diagonal) groups. 95% credible intervals in brackets.

- 135
- 136

	Female	Male	Η	Stem Form
Female		0.62	0.35	0.15
		(-0.06-0.85)	(0.27 - 0.46)	(-0.13-0.46)
Male	0.76		0.52	0.12
	(0.53-0.92)		(0.44 - 0.58)	(-0.24-0.33)
Height	0.28	0.45		0.52
	(0.10-0.36)	(0.45-0.66)		(0.31-0.54)
Stem Form	0.23	0.12	0.52	
	(-0.18-0.54)	(-0.26-0.49)	(0.44-0.58)	

138 Female, absolute female reproduction; Male, qualitative male reproduction (scale 0-3).

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  differentiation for size at first reproduction through male versus female functions in the
  widespread Mediterranean tree *Pinus pinaster*. Ann Bot 110:1449–1460.

# APPENDIX V

Somatic and genetic costs of reproduction in a conifer tree. Santos-del-Blanco L, Climent J. •

- 1 Somatic and genetic costs of reproduction in a conifer tree.
- 2 Luis SANTOS-DEL-BLANCO<sup>1,2,3\*</sup>, José CLIMENT<sup>2,3</sup>
- <sup>3</sup> <sup>1</sup> Department of Ecology and Evolution, University of Lausanne, Biophore Building, CH-
- 4 1015, Lausanne, Switzerland.
- <sup>5</sup> <sup>2</sup> Departamento de Ecología y Genética Forestal, CIFOR-INIA, Carretera de La Coruña km
- 6 7.5, 28040 Madrid, Spain
- 7 <sup>3</sup> Instituto Universitario de Gestión Forestal Sostenible www.sostenible.uva.es
- 8 \*Corresponding author:
- 9 Luis Santos-del-Blanco
- 10 Departamento de Sistemas y Recursos Forestales, CIFOR-INIA, Carretera de La Coruña km
- 11 7.5, 28040 Madrid, Spain
- 12 Telephone: +34 91 3473904
- 13 Fax: +34 91 3476767
- 14

#### 15 ABSTRACT

Costs of reproduction lie at the core of basic ecological and evolutionary theories, and their 16 existence is commonly invoked to explain adaptive processes. Despite their sheer importance, 17 empirical evidence for the existence and quantification of costs of reproduction in tree 18 19 species, comes mostly from correlational studies and more comprehensive approaches remain missing. Manipulative experiments are a preferred approach to study cost of reproduction, as 20 they allow controlling for otherwise inherent confounding factors like size or genetic 21 22 background. Here, we conducted a manipulative experiment in a Pinus halepensis common garden comprising Spanish populations, removing developing cones from a group of trees 23 and comparing growth and reproduction after treatment with a control group. Manipulated 24 trees grew slightly more than control trees just after treatment, but not significantly. Larger 25 differences were observed for the number of female cones initiated one year after treatment, 26 27 with an increase of 70% more cones in the manipulated group. Phenotypic and genetic correlations between vegetative size and female reproduction were also calculated, providing 28 29 no evidence for costs of reproduction.

#### 30 INTRODUCTION

31 Life history theory assumes that costs, or trade-offs, exist between functions linked to fitness, namely survival, growth and reproduction (Roff, 1992). The study of the costs of 32 33 reproduction in terms of growth, survival or subsequent reproduction has a long tradition in evolution and ecology, but the approaches used by both disciplines usually differ and focus 34 on different conceptions of costs of reproduction (Reznick, 1992). When considering 35 36 evolutionary change, it is only genetically-based costs that are relevant to future adaptation. In turn, when considering plastic, environmentally-driven responses within genotypes, 37 38 physiological costs are those that matter.

Four approaches have been used to illustrate costs of reproduction (Reznick, 1985): phenotypic correlations, experimental manipulations of reproduction, genetic correlations and selection experiments. Among these approaches, only genetic correlations and selection experiments are valid to test the assumptions of life history theory (Reznick, 1985). On the other hand, phenotypic correlations and, preferably, experimental manipulations illustrate physiological costs of reproduction provided that confounding factors such as environmental and genetic variability are controlled (Obeso, 2002).

46 The study of costs of reproduction in trees is more complex and variable than in short lived taxa, as investment in growth, maintenance and reproduction might change along time 47 (Thomas, 2011). Available evidence for tree species highlights a physiological trade-off 48 between reproduction and growth, with several recent papers (Sánchez-Humanes et al., 2011; 49 50 Alla et al., 2011; Sala et al., 2012; Barringer et al., 2013; Żywiec & Zielonka, 2013) and a review (Thomas, 2011) helping to define mechanistic links between both functions. 51 52 Reproduction has been related to reduced growth in terms of stem diameter and shoot elongation, but other traits like leaf size and chemistry (Leal & Thomas, 2003), and 53

54 photosynthesis (Obeso, 1998) have been also used to illustrate costs of reproduction in trees. 55 Despite reflecting a more basic trade-off than that between reproduction and growth, negative 56 correlations between current and future reproduction are not always explored, and evidence 57 in plants remains only marginal (Obeso, 2002).

Studies reporting on physiological costs of reproduction in trees are mainly based on 58 correlational studies (Thomas, 2011). The most basic approach consists on relating a variable 59 60 of interest, usually growth, with naturally-occurring variable levels of reproduction (or vice 61 versa) at different hierarchical levels, from shoots to whole the tree (Sánchez-Humanes et al., 2011), and in one or several populations or species (Barringer et al., 2013) along one or 62 63 several growth seasons (Pulido et al., 2013). Some studies have used manipulation experiments, helping to generate more variability than naturally available for water or 64 65 nutrient resources (Sánchez-Humanes & Espelta, 2011; Montesinos et al., 2012; Pulido et al., 66 2013). Moreover, direct manipulation of reproduction is widely considered a more straightforward method for highlighting costs of reproduction (Reznick, 1992; Obeso, 2002), 67 68 but it has so far been reported only once in a tree species, with non-conclusive results (Fox, 69 1995). Indeed, manipulation of reproduction in trees poses great practical challenges in such large organisms. 70

71 Genetically-based costs of reproduction in trees are also scarcely reported in the plant ecology literature given that genetic trials must be planted and cared after at least until trees 72 reach reproductive maturity, and then there is the issue of defining the variables that best 73 74 represent investment in growth and in reproduction. There are however some examples in the 75 forestry literature reporting on negative genetic correlations between reproductive and vegetative traits (Schmidtling, 1981; El-Kassaby & Barclay, 1992; Santos-del-Blanco et al., 76 77 2010). Non-significant or positive correlations between current reproduction and growth or 78 future reproduction often come from studies where environmental confounding factors like
79 tree size are not controlled (Almqvist et al., 2001; Hannerz & Aitken, 2001; Bilir et al., 2002). Moreover, the use of genotypes selected for enhanced growth in breeding programs 80 can incorporate a severe bias that limits a broad interpretation of results (El-Kassaby & 81 82 Barclay, 1992). Data from tree breeding programs could nonetheless be used to illustrate genetic costs of reproduction by comparing reproductive traits in progenies from trees 83 84 submitted and not submitted to phenotypic selection. This, however, has not yet been done and there is only indirect evidence for genetic costs of reproduction derived from artificial 85 selection (Varghese et al., 2009). 86

Overall, it seems that a comprehensive study of costs of reproduction in trees is 87 impaired mainly by logistic reasons. In this study we tried to increase our knowledge of costs 88 of reproduction both at the phenotypic and genetic levels by focusing on a precocious 89 Mediterranean pine whose reproductive ecology is becoming reasonably well understood, the 90 91 Aleppo pine (Pinus halepensis Ait.) (Ne'eman et al., 2004, 2011; Climent et al., 2008; 92 Santos-del-Blanco et al., 2013). We specifically (1) test the existence of a physiological 93 trade-off between current reproduction and subsequent growth and reproduction at the whole 94 tree level by experimentally manipulating reproduction in young *Pinus halepensis* trees, and (2) provide estimates of phenotypic and genetic correlations between vegetative and 95 reproductive traits from common garden experiments, indicative of costs of reproduction. 96

#### 97 MATERIAL AND METHODS

98 Study site

Our experiment was carried out in a provenance-progeny common garden planted in 1995
with one year old seedlings. Size and female reproduction was periodically registered for all
trees in the common garden since establishment. Details can be accessed in (Santos-delBlanco *et al.*, 2010) and www.genfored.es. .

103 Study species

104 Pinus halepensis has several features that make it a suitable species for the study of costs of (female) reproduction. First, it is a precocious species, producing female cones as early as 105 three years old. Second, it has a relatively low size at maturity, which at early ages enables 106 107 manipulation without need of special equipment like scaffolds or elevator platforms. Third, it 108 has a high reproductive allocation and produces cones regularly along its life, which 109 facilitates the detection of costs of reproduction (Grivet et al., 2013).. And fourth, it is a 110 wide-spread species, with populations putatively adapted to varying ecological conditions 111 that might reveal compromises between reproduction and other life history traits (Climent et 112 al., 2008).

*Pinus halepensis* has a two year female cone developmental cycle, thus at any time there are potentially two cohorts of developing female cones (Pardos *et al.*, 2003) (Figure 1). Female floral induction of vegetative buds takes place at the end of the summer and female cones emerge the following spring (Enescu, 1987), generally from vigorous branches of the upper half of the crown (Shmida *et al.*, 2000). Pollination then takes place in spring but ovule fecundation and seed and cone development are arrested until the following year. First year female cones then fully grow along the season completing their development two years after pollination. Developed cones are attached to their bearing branches through a very thick
peduncle so they rarely drop after final development (Ne'eman *et al.*, 2004). Female cones in *Pinus halepensis* trees typically precede male cones in one or several years at the onset of
reproductive life. Therefore female reproduction is clearly more relevant during early stages
in this species (Ne'eman *et al.*, 2011), unlike other related pine species like *P. pinaster*(Santos del Blanco et al 2012) or *P. pinea* (Shmida *et al.*, 2000).

#### 126 Manipulative experiment

In May 2010, when trees where 16 years old, a manipulative experiment was carried out. At 127 that moment, still about half of the trees in the common garden remained vegetative or had 128 produced a low number of cones (less than five). We imposed a minimum threshold of five 129 130 cones in the previous two years for the trees to be eligible for the experiment. The aim was to increase power in detecting significant differences. Among those with at least five cones, we 131 132 selected trees evenly in the range of 5-61 cones per tree by subdividing them into three 133 categories, low, medium and high reproduction. In total, 110 trees were selected, randomly 134 assigning half of them to the treatment group and leaving the other half as a control group (CTR group hereafter). We ensured that there were no statistical differences in size or female 135 reproduction between groups previous to treatment. 136

All developing female cones, both of current year (strobili) and previous year
(conelets) were counted and removed from trees in the treatment group (FCR group
hereafter). A whole-tree assessment of the treatment effects was preferred since
compensation between plant parts could mask processes at the branch level (Karlsson &
Mendez, 2005), as suggested by a previous correlational study in *P. halepensis* (Ne'eman *et al.*, 2011). Special care was taken when removing the cones as not to damage the branches.
Current and previous year female cones were also counted in trees from the control group.

We did not assess costs of male reproduction for two reasons. Firstly because of the above mentioned prevalence of female gender during early reproductive stages (Ne'eman *et al.*, 2011) and, second, because a similar removal of male cones in pines is not possible without severely damaging vegetative structures of the affected shoot

At the end of 2011autumn growth season, thus two years after treatment a radial wood 148 microcore was extracted at breast height from each tree. Wood microcores were mounted on 149 150 a frame, sanded and radial growth for years 2009, 2010 and 2011 measured with a tree-ring measurement station. Basal area for each tree and year was calculated from diameter at breast 151 height in late 2011 and yearly radial growth increments. Current year female cones were also 152 153 counted in 2011. Basal area and female reproduction provided information to assess costs of reproduction with a holistic approach, focusing on their phenotypic outcome but not on their 154 proximate causes. 155

156 Data analysis.

Differences in vegetative growth one and two years after treatment were evaluated with linear mixed models which included treatment as a fixed factor and basal area previous to treatment as a covariate. Model term significance was evaluated by means of Likelihood Ratio Tests. The interaction between treatment and covariate was non-significant and therefore it was removed from final models. The provenance (population of origin) of each individual was included as a random factor in the model in order to avoid pseudoreplication.

163 Count data for female cones was analysed with Poisson Generalised Linear Mixed 164 Models. We first tested for differences between treatment and control groups in absolute 165 number of cones produced after the treatment (counted in late 2011). As before, treatment 166 was defined as fixed but we used no covariate. Population was included as a random factor as 167 well as an observation-level random factor in order to account for overdisperssion in our data (Elston *et al.*, 2001). Finally, in order to check for differences in reproductive allometry
between treatments, we included the logarithm of basal area in 2011 as a covariate in the
model.

Complementarily, correlations between reproduction and size, indicative of 171 phenotypic and genetic costs of reproduction were calculated with measurements at 16 years 172 from the entire trial (28 populations 144 families, 1305 trees). We calculated correlations 173 between biomass in 2009 (as data were available for all individuals) and absolute 174 (reproductive investment) or relative (reproductive allocation) reproduction until 2009. 175 Reproductive allocation was calculated by directly dividing the number of cones into 176 biomass, at the individual level (Karlsson & Mendez, 2005). Finally, we also calculated 177 correlations between biomass growth between 2005 and 2009 and number of cones produced 178 during that period, each variable reflecting investment in vegetative growth or reproduction, 179 180 respectively. Phenotypic correlations were estimated as Pearson's correlation coefficients 181 between phenotypic values after correcting for population effects. As a proxy for genetic 182 correlations we calculated Pearson's correlation coefficients between family corrected means. 183 All models and tests were performed with R packages lme4 v. 0.999999-0 and lmerTest v.1.2-0. 184

#### 186 RESULTS

Manipulated trees, i.e. those with removed current and first year female cones, FCR group, grew slightly more than control trees after treatment, but differences were only marginally significant in 2010 (P  $X_1^2$  3.33 = 0.07) and this small significance disappeared the following year (Table 1, Figure 2).

By contrast, the female cone removal treatment showed a greater effect in subsequent reproduction. FCR trees produced, on average about 70% more female cones than control trees (Table 1, Figure 3). We found only additive effects of treatment on reproductive allometry, indicating a similar effect of experimental treatment across tree sizes (P  $X_{1}^{2} 6.05 =$ 0.014).

The phenotypic correlation between size and reproduction, either absolute or relative, was positive and significant, i.e. bigger trees, or those that grew more, yielded more female cones (Table 2). The correlation between size and number of cones per size unit (RA) was however negative. Among-family genetic correlations between size and reproductive investment were positive and significant but lower than phenotypic correlations, while those with reproductive allocation were negative and significant (Table 2).

#### 202 DISCUSSION

Experimental removal of developing female cones in *Pinus halepensis* resulted in enhanced production of female cones one year later, even when treatment had only a minor effect on vegetative growth. To our knowledge, this is the first account on somatic costs of reproduction in a forest tree as assessed by a manipulative experiment.

207 Induction of vegetative meristems into meristems giving rise to female cones takes place after summer (Enescu, 1987), and induction depends on nutrient content on branches, 208 such that vigorous branches with higher nitrogen content are more likely to bear female cones 209 (Ne'eman *et al.*, 2011). Then, fruit development takes place partially at the expenses of 210 stored N and P, that in the case of the masting species *Pinus albicaulis* can be even depleted 211 212 (Sala et al., 2012) but with no apparent effect on vegetative growth. By removing developing 213 female cones, trees in our experiment were probably left with a surplus of readily available 214 resources that could be allocated to other functions or stored. Thus, cone induction in the 215 following year could be influenced in two ways. First, indirectly by the investment of those 216 resources into vegetative growth that could help to acquire and store more resources. And second, directly by readily stored-extra resources the previous year. Measurement of nutrient 217 218 (Ne'eman et al., 2011; Sala et al., 2012) and carbohydrates (Ichie et al., 2013) at the branch 219 and tree level could help elucidate between those alternatives. Our results suggest that manipulation of reproduction can have greater carry-over effects on reproduction compared 220 to immediate effects on vegetative growth. 221

Experimental manipulation of reproduction proved critical in the detection of costs of reproduction in our study. Indeed, manipulative experiments have been recommended for the study of costs of reproduction in plants, while controlling for confounding factors like size and genetic background (Obeso, 2002). For example, large trees invest more in both vegetative and reproductive growth, leading to positive correlations (de Jong & Klinkhamer,
2005) which indeed are the base of allometric relationships between size and reproduction
(Niklas & Enquist, 2003). Such an example is also provided by our data at the phenotypic
level. Therefore only rarely a negative correlation between absolute size and reproduction is
described but see (Hansen *et al.*, 2013). Besides size effects, other environmental factors can
blur the detection of costs of reproduction, but also produce false positives (Knops *et al.*,
2007). Experimental manipulations are therefore recommended whenever possible.

233 Despite the high number of trees and contrasted origin of the families included in our study, we only found mixed support for phenotypic and genetic costs of reproduction in this 234 species, indicative of evolutionary constraint. A negative relationship of growth and 235 reproduction was only evident at the individual and family level when comparing absolute 236 237 size and reproductive allocation, also reported in (Santos-del-Blanco et al., 2010). Such 238 comparisons have raised concern about spurious correlations (Klinkhamer & de Jong, 1990), 239 however they also describe a biological reality, and the problem might be not the use of such 240 correlations but their straightforward interpretation as evidence for costs of reproduction.

In our common garden study, smaller trees invested a higher proportion of their 241 242 resources in reproduction, compared to larger trees (Table 3), a pattern also described in (Climent et al., 2008). This could be caused by costs of reproduction, dragging the growth of 243 those trees that started to reproduce at smaller sizes (Climent et al., 2008) or just by specific 244 allometric patterns that might determine a more intense relative reproduction at young ages. 245 To the extent that the allometry of reproduction is genetically determined for a species, one 246 247 could interpret the difference between the theoretical reproductive allocation and the actual reproductive allocation as driven by costs of reproduction, i.e. the predicted reproductive 248 allocation, or that of a control group, can provide a reference by which decide whether the 249

relationship between size and the actual reproductive allocation can or cannot be influencedby costs of reproduction. Also here, manipulative experiments would be a preferred tool.

Even if present, several factors could have affected our ability to detect such costs. 252 First, family estimates had high errors as the number of trees per family was relatively low. 253 254 Second, neutral genetic variability in Spanish populations is low, following recent range expansion (Grivet et al., 2011). Thus, although neutral and adaptive genetic variability need 255 not be correlated, and despite our comprehensive sampling effort, Spanish populations might 256 lack enough variation. And third, tree phenotypes are complex and lack of correlation is not 257 enough proof against it, also because there might be third variables indirectly influencing the 258 259 expression of trade-offs, like variability in resource acquisition (Friedman & Barrett, 2011).

Reproduction in trees has been an overlooked trait in studies addressing adaptation, while the relationship between fitness and reproduction is much more direct than that with vegetative traits. A deeper knowledge on tree reproductive ecology is thus still needed to understand adaptive processes in forest stands. Our experiment has shown that as long as reliable approaches and adequate model organisms are chosen, simple methods can be useful to illustrate costs of reproduction in trees.

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- 375 TABLES
- Table 1. Least square mean values per trait and p-value for the test of differences between
- 377 *Pinus halepensis* trees subjected to removal of developing cones (FCR) and control (CTR)
- 378 groups.

Trait	FCR	CTR	p-value
Basal area 2010 (mm <sup>2</sup> )	6188±43	6088±43	0.07
Basal area 2011 (mm <sup>2</sup> )	7638±112	7551±113	0.77
Female cones 2011	$16.8 \pm 2.0$	9.2±2.1	< 0.01

- 380 Table 2. Phenotypic and genetic (family) correlations between vegetative and reproductive
- 381 size in *Pinus halepensis*. RI correlation between absolute vegetative and reproductive sizes.
- 382 RA correlation between absolute vegetative size and reproductive allocation. R-G correlation
- between growth in the period 2005-2009 and the number of female cones produced.

	Phenotypic n= 1305	p-value	Family n=144	p-value
RI	0.46 (0.42-0.50)	< 0.001	0.38 (0.23-0.51)	< 0.001
RA	-0.40 (-0.440.35)	< 0.001	-0.43 (-0.560.29)	< 0.001
R-G	0.31 (0.27 – 0.36)	< 0.001	0.21 (0.05 – 0.36)	0.010

# 384 FIGURES

- Figure 1. Developmental stages of female strobili in *Pinus halepensis*. a) Female strobili in spring, soon after budburst. b) Female conelets in
- 386 spring, one year after budburst. c) Fully-developed and ripe two-year old female cones.
- 387



Figure 2. Scatterplots showing the relationship between basal area preivous to treatment
(2009) and a) one or b) two years after treatment. Red dots, control group; Blue dots, female
cone removal treatment.





- Figure 3. Boxplots of a) basal area in 2010, one growth season after experimental
- 394 manipulation, and b) number of female cones in 2011, one reproductive season after
- 395 experimental manipulation comparing control Pinus halepensis individuals (CTR) with others
- 396 subjected to experimental removal of developing female cones (FCR).



# APPENDIX VI

Guidelines for estimation of quantitative genetic parameters of reproductive traits. **Santos-del-Blanco L**, Climent J.

- 1 Guidelines for the estimation of quantitative genetic parameters for reproductive
- 2 traits in trees
- 3 Luis SANTOS-DEL-BLANCO<sup>1,2,3</sup>, José CLIMENT<sup>12</sup>
- <sup>4</sup> <sup>1</sup> Departamento de Ecología y Genética Forestal, CIFOR-INIA, Carretera de La Coruña
- 5 km 7.5, 28040 Madrid, Spain
- 6 <sup>2</sup> Instituto Universitario de Gestión Forestal Sostenible http://sostenible.palencia.uva.es
- 7 3 Department of Ecology and Evolution. University of Lausanne. Biophore Building.
- 8 CH-1015, Lausanne, Switzerland

#### 9 INTRODUCTION

10 Reproductive strategies lie at the core of adaptive evolutionary processes and are studied in depth in evolutionary ecology (de Jong & Klinkhamer, 2005). Reproductive 11 traits in forest trees are often neglected and the focus is instead placed on growth, 12 physiological and defence traits and survival. However, phenotypic selection on 13 14 putatively adaptive traits will be evolutionarily neutral unless traits covary genetically with fitness (Robertson, 1966; Price, 1970). Given a direct link between fitness and 15 16 reproduction, reproductive strategies are key to understand adaptive processes. Trees are long lived organisms and have a considerably long reproductive period 17 from maturity to senescence (Petit & Hampe, 2006). This enables trees to face variable 18 19 environmental conditions during many seasons, increasing the odds to find suitable 20 conditions for seedling establishment. Therefore, at any given time, it seems more rewarding in terms of fitness to focus on traits related to survival rather than to 21 22 reproduction. However, when thinking in adaptation to long-term changes like those predicted by climatic models, reproductive traits cannot be dismissed. In fact, enhancing 23 24 and anticipating natural recruitment has been suggested as a tool for forest adaptation to 25 climate change (Aitken et al., 2008), but we need a much deeper knowledge of 26 reproduction than we currently have in forest trees. Reproductive traits have been extensively studied in seed orchards from 27

breeding programs in a few commercially interesting forest trees (El-Kassaby & Cook,
1994; Burczyk & Chalupka, 1997; Nikkanen & Ruotsalainen, 2000; Bilir *et al.*, 2002;
Lindgren *et al.*, 2004; Prescher *et al.*, 2007). Since panmixia (equilibrated inter-crossing
among genotypes) is a basic assumption in seed orchard managent, deviations from
panmixia either via male of female sexual functions are a matter of concern as they
lower the genetic variability in the offspring (Kang *et al.*, 2003). As the final

contribution to the next generation will be determined by the absolute number of seeds
produced and sired by each tree, the effect of tree size in reproductive output is usually
neglected in seed orchard studies, not least because trees in seed orchards are usually
pruned, such that tree size can be artificially homogenised.

In natural stands, however, the relationship between size and reproduction, i.e. 38 the allometry of reproduction, determines key life history traits (Thomas, 2011). First, 39 40 the threshold size for first reproduction defines the onset of the reproductive stage for either male, female or both functions. Both under a migration scenario or when life 41 expectancy lowers (for example due to disturbance), threshold size for reproduction is a 42 43 key adaptive trait (Kozłowski, 1992). Second, during the adult stage, the relationship between Vegetative and Reproductive size defines the proportion of resources devoted 44 to reproduction. According to allocation theory, growth, reproduction and defence 45 46 trade-off against each other as a result of drawing resources from a common source (Reznick, 1992). Selection acting in a given environment will ultimately define which 47 48 optimal share of resources, and at which moment, will be devoted to each function in order to maximize fitness (Roff, 1992). For example, a recent a review has described a 49 global pattern towards earlier reproduction in plants as a result of climate change 50 51 confirming that reproductive traits are central to adaptive micro-evolution (Munguía-52 Rosas et al., 2011).

Thus considering or not the relationship between size and reproduction will ultimately depend on the questions to be answered. Here we define precocity either as the proportion of reproductive trees from a group (e.g. family or population) at a given time, irrespective of size. It is important to distinguish ontogenetic precocity along the developmental program from phenological precocity within years. Given that the onset of reproduction in plants is more strongly determined by size than by age (de Jong & Klinkhamer, 2005; Weiner *et al.*, 2009), it seems adequate to try to separate to which
extent onset of reproduction in a tree is determined by changes in size or in allometry.
This can be achieved simply by including size as a covariate in statistical models, and is
useful to standardise estimations of precocity between studies for which growth rates
might differ.

Similarly, we define fruitfulness as the total reproductive output of an individual 64 65 at a given time, irrespective of size. With cumulative reproductive investment we refer to the accumulated reproductive output throughout a certain period of the life of an 66 individual. Total reproductive output will be largely determined by size, but it is 67 68 possible for two equally sized trees to have a different investment in reproduction (Santos-del-Blanco et al., 2013). Reproductive allocation is defined as the amount of 69 70 resources devoted to reproduction relative to total resources (Karlsson & Mendez, 71 2005). For a given size, a greater investment in reproduction implies greater reproductive allocation. However, unless the relationship between reproduction and size 72 73 (as a proxy to available resources) has a slope of 1, reproductive allocation will not be constant but also dependent on size, and again, differences in reproductive allocation 74 between two individuals can be driven by size alone (de Jong & Klinkhamer, 2005). 75 76 Log (reproduction) – log (size) allometric models, commonly known as RV models, 77 have been recommended to model the relationship between reproduction and size due to their simplicity and flexibility (Klinkhamer et al., 1992). 78 Interestingly, many conifer species used in forest breeding are monoecious, 79 80 bearing independent male and female reproductive organs on single trees. This enables the separate study of both sexual functions, as well as the ratio of male to total 81

82 reproduction (sex allocation) within individual trees, and to test ecological theories

83 predicting differences in timing or intensity between sexual functions (Fox, 1993). For

example, it is commonly assumed that wind-pollinated plants will tend to have more
male-biased reproduction at larger sizes as tall plants are able to disperse pollen more
efficiently (Klinkhamer *et al.*, 1997).

87 However, an added difficulty in the analysis of reproductive traits is their discontinuous nature. In the case of precocity or threshold for reproduction, individual 88 trees either reproduce or not, following a binary variable. In the case of fruitfulness or 89 90 RV models, reproduction is sometimes coded as counts of fruits or seeds if not in weight, and is typically strongly asymmetrical. Also when analysing sex ratio data, 91 problems also arise from non-normal distribution of proportions (Wilson & Hardy, 92 93 2002). The analysis of discontinuous variables is easily accommodated by Generalised Linear Models (GLM) using logit or probit link functions for binary or proportions data 94 or logarithmic or inverse gamma link functions for count data. GLMs can also be 95 readily extended into Generalised Linear Mixed Models (GLMM) that include random 96 variables. GLMMs are suitable for the analysis of hierarchical designs while avoiding 97 pseudo-replication and also for variance partition, for example in genetic trials (Zuur, 98 2009; Nakagawa & Schielzeth, 2010). A major constraint in the use of GLMMs in the 99 estimation of quantitative genetic parameters has been that standard formulas need to be 100 101 modified, and have been rarely used until quite recently (Nakagawa & Schielzeth, 102 2010). However, these drawbacks should be solved if we are to calculate quantitative genetic parameters of reproductive traits, as cornerstones of evolutionary ecology 103 providing relevant information about the genetic basis of phenotypic variation, the 104 105 possibility for traits to be modified by natural or artificial selection, and the degree of current genetic differentiation among groups. 106

Along these guidelines we review the literature on quantitative genetics of
reproduction in forest trees and propose a methodology for its standardised study,
providing a case study with data from a *Pinus pinaster* provenance-progeny trial.

#### 111 MATERIAL AND METHODS

#### 112 Analysis of binary, proportion and count data

Binary data are frequent in field genetic trials and convey information about the 113 114 presence or absence of relevant features like survival (Aparicio et al., 2012), disease (Vivas et al., 2011) or reproduction (Santos-del-Blanco et al., 2013). Binary data are 115 116 considered to have a low information content, as under the assumption of a polygenic 117 basis of inheritance of such traits, we can only record two alternative phenotypes. The underlying continuous trait controlling the binary variable is known as liability, and 118 cannot be observed directly. Instead, values of the liability above or below a certain 119 120 threshold produce contrasting phenotypes, and thus they are also known as threshold traits (Falconer, 1989; Lynch & Walsh, 1998). 121

122 Due to their relevance, various methods for the analysis of binary data from 123 genetic trials have been available since early times. Here, for the analysis of reproductive data we advocate the use of Generalised Linear Mixed Models as they can 124 125 readily accommodate several explanatory covariates, like size, as well as confounding variables. In such models reproduction, coded as binary, is the response variable. In a 126 typical half-sib design, family needs to be defined as a random variable for the 127 128 estimation of additive genetic variance. Additive genetic variance in half-sib designs is estimated as four times the family variance, just as in models with Gaussian data. It is 129 also possible to fit an individual-level variable instead and define a pedigree, as to 130 obtain a matrix that is proportional to the variance-covariance structure of additive 131 genetic effects. These models are known as animal models, as they have been mainly 132 developed by animal breeders, and allow a direct estimation of additive genetic effects 133 from the model. Other variables can also be included in the model in order to account 134

for pseudoreplication e.g. block effects, to obtain variance estimates used in quantitativegenetic formulae e.g. population effects for Qst or to model overdisperssion.

An important difference of binary models with respect to Gaussian models is 137 that the error term is fixed to an arbitrary value depending on the statistical software 138 used. Although this avoids a direct comparison of the values of variance estimates 139 140 across studies, has little effect on variables that are derived from ratios of variance 141 components like repeatability, heritability or Qst. Total variance, will thus be 142 determined by the sum of variance components of the model (population, family, block, overdisperssion, error...), together with the fixed variance associated with the link 143 144 function. For the analysis of binary data it is sensible to choose between logit or probit link functions. The variance of a logit distribution is  $\pi 2/3$ , while that of a probit 145 distribution is one. Heritability in the transformed (latent) scale in GLMMs is therefore 146 147 calculated analogously to Gaussian models dividing additive genetic variance into total variance. Qst is calculated dividing population variance into the sum of population 148 variance plus two times the additive genetic variance. 149

150 Proportion data are a special case of binary data. Proportions are commonly used to report on, for example, germination rates or herbivore or pathogen damage. When 151 152 applied to the proportion of male relative to total reproduction (sex allocation or sex 153 ratios), they are a powerful tool to test ecological theories. Proportion data can be arcsin or square root transformed to achieve normality, but using a single estimate might be 154 statistically inefficient as important information regarding sample size is lost. For 155 156 example, a proportion of 0.25 can be achieved with the germination of one out of four seeds or of a hundred out of four hundred, being in this last case the estimation more 157 158 precise. GLMMs make use of all information by considering all untransformed data and

thus are less prone to bias. As argued before, within-individual errors can be estimatedfrom proportion data, and thus are not fixed in the model.

After the onset of reproduction, trees start producing a low number of 161 162 reproductive structures and there is typically strong asymmetry among individuals. Poisson or inverse gamma models with logarithmic error distribution are commonly 163 164 recommended to deal with such data. Quantitative genetic parameters derived from 165 Poisson models are however more difficult to obtain as they depend on the average 166 value of the dependent variable, e.g. average number of cones per individual in the data. Heritability in the latent scale with additive overdisperssion is calculated according to 167 the formula: 168

$$h^{2} = \frac{\sigma_{a}^{2}}{\sigma_{a}^{2} + \sigma_{e}^{2} + \ln(1/(\exp(\beta_{0}) + 1))}$$

169

170 Where  $\sigma_a^2$  is the additive genetic variance,  $\sigma_e^2$  corresponds to additive overdisperssion 171 variance and  $\beta_0$  is the model intercept on the link scale i.e. untransformed.  $\beta_0$  informs 172 about the average value of the variable, and thus heritability estimates depend on 173 average trait values. In models with covariates, e.g. reproductive allometry models,  $\beta_0$ 174 informs about the expected reproductive value for an individual of size equal to 0. In 175 this cases, it is possible to substitute  $\beta_0$  by the actual average reproductive value in the 176 population (Santos del Blanco et al in prep).

#### 178 A worked example with *Pinus pinaster*

Data for the present study were obtained from the *Pinus pinaster* provenance-progeny
common garden described in (Santos del Blanco et al 2012). The common garden
comprises 2,767 trees from 194 open-pollinated families and 23 natural populations. Of
the 2,767 trees initially planted, 2,240 trees were still alive at last measurement in 2010.
Reproductive and growth-related traits (Table 1) were recorded in 2007, 2009 and 2010,
covering the transitional period towards reproductive maturity in most of the trees.

General and Generalised Linear Mixed Models were fitted for size and 185 reproductive data, respectively (Table 2). Bayesian inference as implemented in 186 187 MCMCglmm R package, was preferred over restricted maximum likelihood methods as 188 it allows a more accurate and intuitive estimation of errors of parameters that are 189 derived from ratios of other parameters like heritability, Qst or the median threshold size for first reproduction. Animal models were used to estimate additive genetic 190 191 variance directly from the model. Pedigree was constructed assuming that mothers (dams) within population were unrelated and that progenies were true half sibs. For 192 vegetative size data (height, diameter at breast height and biomass) the inclusion of 193 194 block by family interaction resulted in a high correlation between animal (individual 195 tree) and error terms and model instability. Vegetative size data was instead modelled by substituting animal by family terms. Additive genetic variance was estimated as four 196 197 times the family variance.

Different models were run with varying chain lengths, as necessary to achieve a population of 1000 independent (autocorrelation < 0.1) estimates per parameter. Lack of trends and convergence of the model was checked graphically and by using the Gelman-Rubin statistic comparing the results of two models that had the same specification. Here we report modes and 95% credible intervals for model parameters (Table 3).

#### 203 Literature review

As a previous step to a meta-analysis on quantitative genetic parameters of 204 205 reproductive traits in trees, we surveyed the scientific literature in search of articles reporting broad-sense or narrow-sense heritability, genetic correlations, additive 206 coefficients of variation or Qst estimates. The search was aimed to be as through as 207 208 possible and also included cited references in relevant articles, mostly in the forestry literature. A total of 29 articles were retrieved, most of them reporting on broad-sense 209 210 heritabilities of conifer species, and particularly *Pinus spp.* obtained in seed orchards (Table 4). Studied traits were very diverse and included diverse estimates of male and 211 female reproductive investment (fruitfulness), reproductive allocation, reproductive 212 213 phenology and precocity.

# 214 TABLES

# 215 Table 1. Variables and measurement dates

Variable		Description
Н	Height (cm)	Total height of each tree in 2007, 2009 and 2010
DBH	Diameter at breast Height	Diameter at breast height (1.30 m) measured in 2007, 2009 and 2010
Biomass	Biomass	Inferred from DBH (REF)
F	Female reproduction	Presence or absence of female cones in each tree, yearly from 2007
Fq	Cuantitative female reproduction	Number of seed cones in each tree, yearly from 2007
M	Male reproduction	Presence or absence of pollen cones in each tree in 2009 and 2010
Mq	Cuantitative male reproduction	Number of male cone clusters per tree counted in 15 seconds and modified by a
-	-	factor proportional to cluster size. 2010

Table 2. Model specification

Response variable	Туре	Concept modelled	Fixed terms	Random terms
H, DBH, Biomass	Gaussian	Growth	-	Population, Family, Block, Family by Block
F, M	Binary	Precociousness relative to size	Size (H, DBH or Biomass)	Population, animal, individual tree
F (2008-2010), M (2009, 2010)	Binary	Precociousness at a given age	-	Population, animal
Fq,Mq	Poisson	Fruitfulness relative to size (R-V)	Size (Biomass)	Population, additive overdisperssion, animal
Fq,Mq	Poisson	Fruitfulness (accumulated until 2010)	-	Population, additive overdisperssion, animal
Fq,Mq	Proportion	Sex allocation relative to size	Size (H,DBH, Biomass)	Population, additive overdisperssion, animal

Table 3. Narrow-sense heritability (h<sup>2</sup>) and Q<sub>st</sub> estimates for several vegetative and reproductive traits in *Pinus pinaster*. 95% CI, 95% credible interval. CE, covariate effect. ns, not significant. 

	$h^2$		$Q_{st}$		CE
	mode	95% CI	mode	95% CI	
Growth					
Н	0.13	(0.01-0.30)	0.21	(0.08-0.64)	
DBH	0.13	(0.03-0.34)	0.14	(0.04-0.48)	
Biomass	0.12	(0.02-0.28)	0.12	(0.03-0.42)	
Precociousness relative to size	<u>.                                    </u>				
Female	0.43	(0.27-0.67)	0.13	(0.06-0.23)	+
Male	0.42	(0.28-0.53)	0.07	(0.02-0.14)	+
Precociousness at a given age	_				
F <sub>2008</sub>	0.38	(0.07-0.59)	0.25	(0.07-0.61)	
F <sub>2009</sub>	0.34	(0.16-0.54)	0.14	(0.05-0.35)	
F <sub>2010</sub>	0.32	(0.15-0.50)	0.10	(0.03-0.26)	
M <sub>2009</sub>	0.59	(0.41 - 0.74)	0.08	(0.04-0.20)	
M <sub>2010</sub>	0.51	(0.3-0.70)	0.08	(0.03-0.19)	
Fruitfulness relative to size	_				
FRV	0.34	(0.19-0.62)	0.15	(0.06-0.34)	+
M RV	0.38	(0.23-0.63)	0.09	(0.04-0.23)	+
Fruitfulness 2010	_				
F Fr	0.36	(0.17-0.56)	0.11	(0.04-0.29)	
M Fr	0.48	(0.30-0.70)	0.09	(0.05-0.22)	
Sex allocation					
Sex allocation H	0.29	(0.12-0.43)	0.17	(0.08-0.38)	+
Sex allocation DBH	0.30	(0.13-0.45)	0.20	(0.09-0.38)	ns
Sex allocation Biomass	0.28	(0.15-0.47)	0.16	(0.08-0.36)	+

Table 4. Summary of heritability (narrow-sense for family, F, trials; broad-sense for clonal, C, trials) estimates reported in the literature for
 several tree species.

Species	Trait and (age)	Trial	Heritability (range)	Reference
Picea abies	Cone production (6 / 14)	С	0.14-0.24 / 0.37-0.55	(Almqvist et al., 2001)
Picea abies	Male / female flowering abundance (not reported)	С	0.37 / 0.38	(Nikkanen & Ruotsalainen, 2000)
Pinus elliottii	Cone yield (14-17) / Flower production (4-6)	С	0.50 / 0.49	(Varnell <i>et al.</i> , 1967)
Pinus halepensis	Female flowering (3-4) / cone production (8-9)	C	0.86-0.83 / 0.43-0.41	(Matziris, 1997)
Pinus halepensis	Cone production (9-10)	F	0.68-0.57	(Matziris, 2000)
Pinus halepensis	Female reproductive allocation (11-15)	F	0.29-0.63	(Santos-del-Blanco et al., 2010)
Pinus koraiensis	Male / female strobili (15-18)	С	0.34-0.56 / 0.20-0.46	(Choi <i>et al.</i> , 2004)
Pinus nigra	Cone production (11-13)	С	0.82-0.88	(Matziris, 1993)
Pinus pinaster	Male / female cone abundance (8-11)	С	0.39-0.59 / 0.76-0.82	(Merlo & Fernández López, 2004)
Pinus pinaster	Female reproductive allocation (5)	F	0.32-0.47	(Santos-del-Blanco et al., 2010)
Pinus pinea	Logcone yield (1-10)	С	0.15	(Mutke <i>et al.</i> , 2005)
Pinus pinea	Logcone yield (9)	С	0.19	(Mutke <i>et al.</i> , 2003)
Pinus sylvestris	Cone production (several seed orchards, review)	С	0.24-0.78	(Prescher et al., 2007)
Pinus sylvestris	Log male flowers, log female flowers, log females cones (13-15)	C	0.70 / 0.52-0.54 / 0.37-	(Sivacioglu et al., 2009)
Pinus sylvestris	Male and female pollen and cone production (17-31)	C	0.36-0.54	(Savolainen et al., 1993)
Pinus sylvestris	Male and female strobili number (10-19)	С	(0.00-0.15)- (0-0.16)	(Bilir <i>et al.</i> , 2006)

Pinus sylvestris	Sex ratio (male/female)	C	0.56	(Burczyk & Chalupka, 1997)
Pinus radiata	Male and female onset, ending and duration of flowering phenology (3-5)	С	Male 0.92 / 0.82 / 0.32	(Codesido et al., 2005)
			Female 0.93 / 0.84 / 0.83	
Pinus taeda	Precocity (3-4) / Square root female cone production (2-8)	F	0.47-0.65 / 0.61	(Schmidtling, 1981)
Pinus taeda	Square root +0.5 Male and female cone production (6-18)	С	0.5-0.4	(Schmidtling, 1983)
Pinus taeda	Cone production (4-10), (8-20)	С	0.35-0.40	(Byram <i>et al.</i> , 1986)
Pinus contorta	Male and female flowering (11-29 / 10-11-29)	F	0.40-0.64 / 0.13-0.42	(Hannerz & Aitken, 2001)
Pinus virginiana	Fruitfulness	F	0.59-0.65	Bramlett 1971 cited in (Schmidtling, 1981)
Pseudotsuga menziesii	Cone / seed production (19)	С	0.38 / 0.40	(El-Kassaby & Cook, 1994)
Cryptomeria japonica	Male cones (7)	F	0.78-1.05	(Tsubomura <i>et al.</i> , 2012)
Eucalyptus cladocalyx	Precocity (3)	F	0.51	(Mora <i>et al.</i> , 2009)
Eucalyptus cladocalyx	Flowering intensity (8)	F	0.48	(Cané-Retamales et al., 2010)
Eucalyptus globulus	Precocity (2-4)	F	0.21-0.93	(Jordan <i>et al.</i> , 1999)
Eucalyptus globulus	Presence or absence of reproduction (4)	F	0.47	(Chambers et al., 1997)
Prunus avium	Advanced flowering / flower number / cherry number (6-7)	С	0.82 / 0.90 / 0.64	(Díaz & Merlo, 2008)

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# APPENDIX VII

Ecología evolutiva de la reproducción en dos pinos mediterráneos: *Pinus pinaster* y *Pinus halepensis*. **Santos-del-Blanco L**, Chambel R, Notivol E. Alía R, Climent J. Proceedings 6º Congreso Forestal Español. Sociedad Española de Ciencias Forestales, Vitoria, 2013.



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# Ecología evolutiva de la reproducción en dos pinos mediterráneos: Pinus pinaster y Pinus halepensis

SANTOS DEL BLANCO, L.<sup>1,2</sup>, CHAMBEL, R.<sup>1</sup>, NOTIVOL, E.<sup>3</sup>, ALÍA, R.<sup>1,2</sup> y CLIMENT, J.<sup>1,2</sup>

<sup>1</sup> Departamento de Ecología y Genética Forestal, INIA-CIFOR,

<sup>2</sup> Instituto Universitario de Gestión Forestal Sostenible Uva-INIA

<sup>3</sup> Unidad de Recursos Forestales, Centro de Investigación y Tecnología Alimentaria de Aragón

#### Resumen

La adaptación de los árboles a su entorno está ligada a una adecuada programación de la reproducción, que a su vez está interrelacionada con el crecimiento y mantenimiento. Los ensayos genéticos forestales permiten testar las hipótesis de diferenciación y plasticidad para caracteres adaptativos tan relevantes como el tamaño umbral de reproducción y la alometría de la reproducción. Los resultados de varios ensayos de Pinus pinaster y P. halepensis en condiciones ambientales contrastadas corroboran una diferenciación entre procedencias y la existencia de variabilidad genética para caracteres reproductivos en ambas especies. En general, los ambientes de origen con un alto grado de continentalidad se relacionaron con una reproducción femenina precoz y más intensa en relación al tamaño. A su vez, unas condiciones más limitantes del ambiente de ensayo también tendieron a acelerar la reproducción. En P. pinaster, además, encontramos marcadas diferencias en la asignación sexual entre procedencias. Estos resultados demuestran compromisos entre crecimiento y reproducción, constatados a nivel genético mediante correlaciones genéticas negativas y a nivel fisiológico mediante experimentos de retirada de conos. Esta información es clave para entender la adaptación local y orientar el uso y conservación de los recursos genéticos de ambas especies.

#### **Palabras clave**

Genética forestal, diversidad genética, asignación reproductiva, adaptación local, caracteres de historia vital.

## 1. Introducción

Los caracteres de historia vital son aquellos que definen los patrones de desarrollo en cuanto a crecimiento, reproducción y supervivencia de un organismo (ROFF 1992). Estos caracteres están estrechamente ligados a la adaptación de los organismos al medio, principal materia de estudio de la ecología evolutiva y de gran relevancia en la gestión forestal sostenible. En el caso de especies de árboles forestales, existe un patrón general de ciclo vital definido por altas tasas de mortalidad juvenil, edad de reproducción tardía y alta longevidad, pero dentro de este patrón, existe un amplio abanico de variaciones íntimamente ligado con el ambiente (THOMAS 2011).

Los pinos mediterráneos, y especialmente *Pinus halepensis* Mill. y *P. pinaster* Ait., constituyen un modelo de estudio muy adecuado tanto desde puntos de vista teóricos como aplicados, de cara a conocer los procesos de adaptación al medio, pasados y futuros, de las especies forestales en España (LEV-YADUN & SEDEROFF 2000). Ello



se debe a una serie de características de la biología de estas especies (NE'EMAN et al. 2004, 2011): edad temprana de reproducción, que permite evaluar caracteres reproductivos de forma relativamente precoz; asignación reproductiva elevada, que facilita la detección de costes de reproducción; escasa variabilidad interanual de la reproducción en comparación con otras especies más veceras, lo que posibilita hacer seguimientos con series de datos más continuas y con mejores propiedades estadísticas; mantenimiento variable de los conos femeninos cerrados (serotinia), lo que posibilita la reconstrucción de la reproducción en temporadas pasadas; separación entre reproducción masculina (estróbilos productores de polen) y femenina (conos o estróbilos), lo que hace posible testar hipótesis específicas respecto al papel de la reproducción masculina o femenina en estas especies; reproducción obligatoria por semillas, y no vegetativa, lo que facilita la asociación entre partes del árbol y funciones, e incrementando el valor adaptativo de la reproducción sexual; arquitectura de copa simple y por último, baja longevidad en comparación con otras especies forestales, mejorando las correlaciones entre la inversión reproductiva real a lo largo de toda la vida de un individuo y la estimada en una serie menor de años.

También son destacables las características ecológicas de estas especies, como su amplio nicho ecológico, sobre todo en *P. pinaster* (FADY 2012), que se traduce en un amplio rango de distribución y la posibilidad de testar la relación entre caracteres fenotípicos y ambientales. Otro factor importante de la ecología de estas especies es su relación con regímenes de perturbaciones, específicamente incendios forestales, que han dejado su impronta en adaptaciones como el espesor de corteza o la serotinia (KEELEY 2012).

Por último, la gran superficie ocupada por estas especies en España tanto de forma natural como por reforestación, pone de manifiesto su importancia ecológica. Estas masas, sin embargo, se ven afectadas por incendios forestales cada vez más virulentos y frecuentes lo que precisamente ha motivado varios estudios científicos sobre la ecología reproductiva de P. halepensis (DE LAS HERAS et al. 2007, ESPELTA et al. 2008, GONZALEZ-OCHOA et al. 2004, IRAIMA y ESPELTA 2004, MOYA et al. 2007, ORTIZ et al. 2011, VERKAIK y ESPELTA 2006). Desde el punto de vista de la gestión forestal, se pretenden conocer los tratamientos selvícolas y condiciones en las que se produce una mayor cantidad de piñas de forma más precoz. De esta forma, asumiendo que una gran proporción de piñas en edades jóvenes son serótinas, se construye un banco de semillas aéreo suficiente para garantizar la regeneración post-incendio en caso de corto periodo de recurrencia del fuego. Las principales conclusiones de estos estudios son que las condiciones que aceleran el crecimiento de los árboles como clareos o podas (DE LAS HERAS et al. 2007, MOYA et al. 2007), fertilización artificial o alta calidad de estación (MOYA et al. 2007, ORTIZ et al. 2011), aceleran también la producción de piñas. Sin embargo, una producción de piñas mayor por árbol en masas poco densas, puede verse compensada por un mayor número de pies en masas más densas (MATYAS & VARGA 2000).

A pesar del indudable valor para la gestión forestal proporcionado por los citados estudios, su aportación al campo de la ecología se ve limitado por varios factores. En primer lugar, se considera la producción de piñas en valor absoluto por árbol o por superficie independientemente del tamaño del árbol. Así los árboles de mayor tamaño son los que normalmente producen más piñas, sin embargo la producción de piñas por unidad



de biomasa es mayor en árboles pequeños (ORTIZ et al. 2011). El segundo problema se da en estudios que comparan diferentes procedencias de regeneración natural. En este caso, no es posible discriminar entre causas genéticas y/o ambientales que expliquen las correlaciones de la reproducción con variables climáticas. Para conocer de forma sistemática si existe variación genética entre procedencias en caracteres reproductivos, una muestra representativa de árboles de las procedencias a comparar deben crecer en condiciones ambientales lo más homogéneas posible.

Los ensayos genéticos forestales de ambiente común (*common gardens*) cumplen precisamente la función de proporcionar condiciones ambientales homogéneas, bajo las que se comparan entradas genéticas que pueden ser de rango poblacional (ensayo de procedencias), familiar (ensayo de progenies) o individual (ensayo clonal). Estos ensayos pueden llevarse a cabo en condiciones de campo o de invernadero. Los ensayos de progenies, además, permiten el cálculo de parámetros de genética cuantitativa que describan la arquitectura y grado de determinación genéticos de los caracteres estudiados.

Los caracteres más comúnmente estudiados en pinos mediterráneos en ensayos de ambiente común han sido por un lado el crecimiento (ALÍA et al. 1995, CHAMBEL et al. 2007) y por otro, caracteres fisiológicos de valor adaptativo, como la resistencia al frío (CLIMENT et al. 2009), a la sequía (ARANDA et al. 2010, VOLTAS et al. 2008) o fenología (CODESIDO y FERNANDEZ-LOPEZ 2009). Desde un punto de vista evolutivo, un carácter se considera adaptativo si tiene una influencia positiva en el número de descendientes producidos por un organismo. De ello se deriva la gran importancia de cuantificar también los caracteres reproductivos, situados de esta forma en un nivel jerárquico superior de relevancia.

La evidencia empírica sobre la estrecha relación entre condiciones ambientales naturales o influidas por el hombre y caracteres reproductivos es abundante en numerosos tipos de organismos (ROFF 1992, STEARNS 1992). Los pinos mediterráneos *P. pinaster* y *P. halepensis*, no parecen ser una excepción, y varios trabajos describen caracteres reproductivos de historia vital en ambas especies (CLIMENT et al. 2008, SANTOS-DEL-BLANCO et al. 2010, 2012, TAPIAS et al. 2004).

## 2. Objetivos

Los objetivos del presente trabajo son testar las hipótesis básicas de la teoría de historia vital aplicadas al caso de especies mediterráneas forestales como son *P. pinaster* y *P. halepensis* y hacer una lectura práctica de los resultados de cara a la gestión forestal. En concreto, pretendemos (1) cuantificar la diferenciación entre procedencias para caracteres de historia vital relacionados con la reproducción (tamaño umbral de reproducción, alometría reproductiva, esfuerzo reproductor), (2) comprobar la relación entre los citados caracteres y variables ambientales en términos de variación, (3) cuantificar la existencia de variación genética aditiva dentro de procedencias para caracteres de historia vital mediante herramientas de genética cuantitativa y (4) estudiar patrones de compensación (*tradeoffs*) entre caracteres de historia vital reproductivos y caracteres de crecimiento vegetativo



# 3. Metodología

#### 3.1 Ensayos de campo

Para *Pinus halepensis* utilizamos un ensayo de procedencias y otro de procedencias-progenies. El ensayo de procedencias fue instalado en 1997, estando replicado en seis localidades, dos de las cuales se incluyen en este trabajo (Valdeolmos, Madrid –P24OLM- y Rincón de Ademuz, Valencia-P24ADE-). En ellos están representadas 52 procedencias nativas de la especie a lo largo de todo su rango de distribución en España continental, Islas Baleares, Italia, Francia, Túnez y Grecia. (ver CLIMENT et al. 2008 para más detalles). Los sitios elegidos presentan condiciones ambientales contrastadas. El sitio de ensayo OLM tiene veranos cálidos e inviernos suaves, siendo el suelo arenoso y profundo. Por el contrario, en ADE los inviernos son fríos y el suelo es rocoso y poco profundo. Debido a la combinación de factores climáticos y edáficos, el sitio de ensayo de ADE es mucho más limitante para el crecimiento vegetativo que OLM.

El ensayo de procedencias-progenies fue instalado en 1995, estando replicado en dos localidades, ambas incluidas en este trabajo (Megeces, Valladolid —F24MEG—, y Montañana, Zaragoza —F24MON—). En ellos están representadas 148 familias de polinización abierta de 32 procedencias repartidas por el rango de distribución de la especie en la Península Ibérica e Islas Baleares. Aquí también existen diferencias entre ambos sitios: F24MEG está situado en un suelo calizo poco profundo y en ligera pendiente, mientras que F24MON está situado en un valle aluvial fértil y con disponibilidad hídrica durante el verano (ver SANTOS-DEL-BLANCO et al. 2010 para más detalles). Adicionalmente tanto en el ensayo de procedencias como en el de procedencias-progenies, se incluyeron progenies de tres rodales de origen desconocido plantados en la provincia de Valladolid que no han sido considerados en el presente trabajo.

Para *Pinus pinaster* hicimos uso de un ensayo de procedencias-progenies instalado en 2005 y replicado en cinco sitios de ensayo, utilizando uno de ellos en este trabajo (A Merca, Orense –F26MER). El ensayo contiene 194 familias de polinización abierta de 23 procedencias naturales distribuidas sobre gran parte del rango de la especie, incluyendo Península Ibérica Atlántica, Francia Atlántica, Córcega, España Mediterránea y Marruecos. Las condiciones ambientales del ensayo se consideran intermedias respecto al nicho ecológico de la especie, pues se sitúa en una zona de transición entre clima Mediterráneo y Atlántico (ver SANTOS-DEL-BLANCO et al. 2012 para más detalles). Todos los ensayos utilizados en este trabajo pertenecen a la red nacional de ensayos genéticos forestales GENFORED.

# 3.2 Toma de datos

Se tomaron datos de variables representativas del crecimiento vegetativo (altura total, diámetro normal) y reproductivas en los diferentes ensayos de campo a edades que variaron entre los 5 y los 15 años (Tabla 1). La altura total se midió con pértiga telescópica y el diámetro normal con forcípula. Dependiendo del ensayo, se discriminaron las piñas por cohortes en función de su tamaño y color: los estróbilos



femeninos formados en la primavera del año en curso son de color rojo y de pequeño tamaño (aprox. 1 cm), al comienzo de la primavera siguiente aumentan su tamaño y toman un color verde (aprox 2-3 cm) y al final de la segunda primavera han completado su desarrollo final pero conservan el color verde, que perderán poco a poco para ser color marrón brillante al comienzo de la tercera primavera. A partir de entonces, las zonas de las piñas más expuestas al sol irán degradando su color tomando colores grisáceos cada vez más claros (GIL et al. 2009). Dado el aún limitado desarrollo de los árboles en la mayoría de los ensayos (F26MER, F24MEG, P24OLM, P24ADE), fue posible distinguir las piñas por cohortes. En MON, sin embargo, el elevado porte y las copas adyacentes de los árboles impidieron distinguir entre cohortes y se estimó el número total de piñas por árbol contadas en 15 segundos (KNOPS & KOENIG 2012). La reproducción masculina sucede a la femenina en el desarrollo de Pinus halepensis (NE'EMAN et al. 2004) pero no así en P. pinaster (SANTOS-DEL-BLANCO et al. 2012), indicando un mayor interés de la reproducción masculina precoz en esta última especie. Se recogieron datos semicuantitativos de reproducción masculina en los ensayos F26MER y F24MEG. El método aplicado fue el conteo de agrupaciones de conos masculinos en 15 segundos y paralelamente aplicar una escala cuantitativa con valores del 1 al 5 dependiendo del tamaño y densidad de dichas agrupaciones. Esta valoración cualitativa se utilizó para ponderar los datos cuantitativos.

A partir de las coordenadas geográficas de las procedencias incluidas en los ensayos, se obtuvieron variables climáticas de temperatura mediante los modelos de clima de GONZALO-JIMÉNEZ (2010) para procedencias de España Peninsular y de HIJMANS et al. (2005) para el resto de procedencias.

Especie	Ensayo	Variables medidas (edad)
Pinus pinaster	F26MER	Altura total (6), diámetro normal (6), reproducción masculina (5) y femenina anuales cuantitativas (5)
Pinus halepensis	F24MEG	Altura total (15), diámetro normal (15), reproducción femenina anual cuantitativa (15)
Pinus halepensis	F24MON	Altura total (15), diámetro normal (15), reproducción femenina total cuantitativa (15)
Pinus halepensis	P24OLM	Altura total (13), diámetro normal (13), reproducción femenina anual cuantitativa (13)
Pinus halepensis	P24ADE	Altura total (12), diámetro normal (12), reproducción femenina anual cuantitativa (12)

Tabla 1. Ensayos de campo de Pinus pinaster y P. halepensis utilizados en este trabajo, y caracteres medidos conedad de medición entre paréntesis

La biomasa total de los árboles se calculó a partir de ecuaciones alométricas (MONTERO et al. 2005). El tamaño umbral de reproducción a nivel de procedencia se derivó combinando medidas de crecimiento vegetativo y presencia o ausencia de reproducción masculina o femenina en modelos binomiales (SANTOS-DEL-BLANCO et al. 2010, 2012).

# 3.3 Análisis de datos

Para testar la hipótesis de diferenciación entre procedencias para los caracteres de interés, ajustamos modelos mixtos generales para las variables continuas (altura total y diámetro normal) y generalizados para las discontinuas (binomial para reproducción en



forma categórica; Poisson para reproducción en forma cuantitativa). De estos modelos se extrajeron las estimaciones medias por procedencia y/o familia para el cálculo de correlaciones ambientales y entre caracteres.

La relación entre valores fenotípicos medios por procedencia y variables climáticas propias de las zonas de origen se comprobó mediante correlaciones de Pearson. Como variable integradora de la condiciones ambientales, se eligió el índice de continentalidad o rango anual de temperaturas (temperaturas máximas – temperaturas mínimas). Un índice de continentalidad alto está asociado a un periodo de crecimiento vegetativo corto y viceversa (SANTOS-DEL-BLANCO et al. 2012).

En aquellos ensayos con representación de progenies (familias), se cuantificó la existencia de varianza genética aditiva dentro de procedencias incluyendo en modelos mixtos el factor familiar además de corregir por las posibles diferencias medias entre procedencias. Para los caracteres de crecimiento se incluyó en la parte fija la estructura de bloques del diseño para reducir el ruido ambiental. Este factor fue eliminado para el cálculo de parámetros genéticos para tener estimaciones más cercanas a las que serían observables en la naturaleza. Para caracteres reproductivos se incluyó como covariable la biomasa de los árboles. De esta forma los modelos reflejan la variación en reproducción entre familias y procedencias debidas a causas diferentes a la variación en tamaño. A partir de los componentes de varianza de modelos mixtos aplicados a datos de procedencias-progenies se calculó la heredabilidad en sentido estricto ( $h^2$ ), diferenciación entre procedencias ( $Q_{ST}$ ) y coeficiente de variación de varianza genética aditiva ( $CV_A$ ) (FALCONER 1989).

La compensación a nivel fisiológico y genético entre caracteres de crecimiento vegetativo y reproductivos (*tradeoffs*) se calculó a tres niveles: individual dentro de procedencias y familias (correlaciones fenotípicas), familiar dentro de procedencias (correlaciones genéticas) y a nivel de procedencia mediante modelos mixtos y correlaciones de Pearson.

# 4. Resultados

# 4.1. Diferenciación entre procedencias

Tanto los caracteres de crecimiento como los reproductivos mostraron diferencias generalizadas entre procedencias para ambas especies, aunque estas diferencias fueron algo menores para caracteres de crecimiento en *P. halepensis* (Tabla 2).

# 4.2. Correlaciones ambientales

Los caracteres reproductivos, en especial el tamaño umbral de reproducción femenino, mostraron correlaciones positivas con el índice de continentalidad mientras que en los caracteres de crecimiento vegetativo las correlaciones fueron negativas en *P. pinaster* y en el ensayo de *P. halepensis* con los árboles más desarrollados (P24MON). En el resto de ensayos de *P. halepensis*, las correlaciones no fueron significativas. En resumen, valores crecientes del índice de continentalidad se relacionaron con menor crecimiento vegetativo y mayor inversión en reproducción femenina (Tabla 3).



# 4.3. Parámetros genéticos

Se hallaron valores altos de heredabilidad para caracteres de crecimiento y reproductivos en el ensayo de *P. pinaster*, mientras que para el ensayo menos desarrollado de procedencias-progenies de *P. halepensis* (F24MEG), los valores fueron moderados. En el ensayo más desarrollado (F24MON), la variación sólo fue significativa entre procedencias y no entre familias de una misma procedencia (Tabla 4).

## 4.4 Compensación entre funciones (tradeoffs)

Las correlaciones entre crecimiento vegetativo y reproducción femenina en ambas especies fueron positivas a nivel fenotípico individual  $(r_{ph})$  y negativas a nivel genético  $(r_a \ y \ r_{pr})$ . En éste último caso, la correlación a nivel de procedencia fue de mayor magnitud.

Tabla 2. Valores de significatividad del efecto de la procedencia sobre diferentes variables relacionadas con el crecimiento vegetativo y la reproducción masculina y femenina para Pinus pinaster y P.halepensis en varios sitios de ensayo. H, altura total; DBH, diámetro normal; biom, biomasa; TSRf, tamaño umbral de reproducción femenino; TSRm, tamaño umbral de reproducción masculino; Rep.f., reproducción femenina cuantitativa; Rep.m. reproducción masculina cuantitativa. n.d., dato no disponible

Especie	Ensayo	Н	DBH	biom	TSRf	TSRm	Rep.f.
P. pinaster	F26MER	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
P.halepensis	F24MEG	< 0.001	0.318	0.048	< 0.001	0.004	< 0.001
P.halepensis	F24MON	0.013	0.002	0.012	n.d.	n.d.	< 0.001
P.halepensis	P24OLM	< 0.001	< 0.001	< 0.001	< 0.001	n.d.	< 0.001
P.halepensis	P24ADE	< 0.001	< 0.001	< 0.001	< 0.001	n.d.	< 0.001

Tabla 3. Coeficientes de correlación ente valores del índice de continentalidad y diferentes variables relacionadas con el crecimiento vegetativo y la reproducción masculina y femenina para Pinus pinaster y P. halepensis en varios sitios de ensayo. Valores en negrita indican correlaciones significativamente (P < 0.05) diferentes de 0. Ver abreviaturas en Tabla 2.n.d., dato no disponible

Especie	Ensayo	Н	DBH	biom	TSRf	TSRm	Repf	Repm
P. pinaster	F26MER	-0.80	-0.65	-0.70	-0.71	0.00	0.40	-0.50
P.halepensis	F24MEG	0.09	-0.40	0.33	-0.51	-0.16	0.48	0.10
P.halepensis	F24MON	-0.37	-0.43	-0.41	n.d.	n.d.	0.45	n.d.
P.halepensis	P24OLM	0.03	0.01	-0.04	-0.53	n.d.	0.53	n.d.
P.halepensis	P24ADE	-0.02	0.02	0.06	-0.40	n.d.	0.33	n.d.

Tabla 4. Parámetros genéticos  $(h^2)$ , heredabilidad,  $Q_{ST}$ , diferenciación ente procedencias  $CV_A$ , coeficiente de varianza genética aditiva) para altura (H), biomasa (biom) y reproducción femenina (TSRf) para Pinus pinaster y P. halepensis en varios sitios de ensayo. Ver abreviaturas en Tabla 2. n.d., dato no disponible

Especie	Ensayo	Н			biom			TSRf	
		h <sup>2</sup>	Q <sub>ST</sub>	CVA	h <sup>2</sup>	Q <sub>ST</sub>	CVA	h <sup>2</sup>	Qst
P. pinaster	F26MER	0.60	0.08	3.56	0.61	0.05	0.43	0.69	0.13
P.halepensis	F24MEG	0.12	0.15	1.05	0.11	0.12	0.89	0.14	0.26
P.halepensis	F24MON	n.s.	n.s.	n.s.	0.05	0.19	0.05	n.d.	n.d.



Especie	Ensayo	r <sub>ph</sub>	p.valor	r <sub>a</sub>	p.valor	$\mathbf{r}_{\mathbf{pr}}$	p.valor
P. pinaster	F26MER	0.23	< 0.001	-0.14	0.047	-0.45	0.031
P.halepensis	F24MEG	0.46	< 0.001	-0.15	0.073	0.14	0.468
P.halepensis	F24MON	0.31	< 0.001	0.01	0.946	-0.46	0.011
P.halepensis	P24OLM	0.33	< 0.001	n.d.	n.d.	-0.39	0.004
P.halepensis	P24ADE	0.36	< 0.001	n.d.	n.d.	-0.12	0.375

Tabla 5. Correlaciones fenotípicas a nivel individual  $(r_{ph})$  y genéticas a nivel familiar  $(r_a)$  y a nivel de procedencia  $(r_{pr})$ entre valores de biomasa y reproducción femenina después de descontar el efecto del tamaño individual. n.d., dato no disponible

## 5. Discusión

Los resultados basados en ensayos de campo para procedencias de todo el rango de distribución de *P. pinaster* (un ensayo) y *P. halepensis* (cuatro ensayos) demuestran una estrecha relación entre los caracteres reproductivos y el ambiente de origen de las procedencia así como ente caracteres reproductivos y de crecimiento entre sí.

La existencia de variaciones para caracteres adaptativos a nivel de procedencia dentro de especies es un hecho admitido para aquellas especies en las que se reconocen subespecies (e.g. *Pinus nigra*) o para las que presentan amplia diferenciación genética neutral ente procedencias (*P. pinaster*), pero es menos conocido cuando se trata de especies que no cumplen estas condiciones (*P. halepensis*). Sin embargo, incluso en la Península Ibérica, donde la variación genética de *P. halepensis* es menor debido a la historia de migración de la especie (GRIVET et al. 2009), la diferenciación entre procedencias también está presente (Tabla 2). Así, salvo casos excepcionales como *P. pinea* (MUTKE et al. 2010), la variación intraespecífica es un hallazgo común. Este hecho pone de manifiesto la relevancia de los datos obtenidos en ensayos de ambiente común para caracterizar los materiales de base de cara a la gestión forestal sostenible.

La mencionada variación intraespecífica descrita tanto para caracteres de crecimiento como reproductivos en *P. pinaster* y *P. halepensis* no es aleatoria sino que está relacionada con las condiciones ambientales en las que las procedencias han evolucionado, sugiriendo que los caracteres estudiados tienen valor adaptativo y que se han producido (y/o están produciendo) fenómenos de adaptación local. Los resultados obtenidos concuerdan plenamente con las predicciones derivadas de la teoría de historia vital, que postulan un mayor tamaño umbral de reproducción en aquellos ambientes favorables para el crecimiento (ROFF 1992).

Sin embargo, a pesar de que exista abundante evidencia de la influencia del ambiente local, por un lado, éste no es estático (LINDNER et al. 2010) y por otro, los mecanismos de adaptación de las especies suelen estar desfasados respecto a los cambios ambientales (REHFELDT et al. 1999). La selección natural actúa sobre fenotipos, que son el resultado de la interacción del genotipo y el ambiente. La influencia del ambiente es variable para diferentes caracteres, estando el crecimiento vegetativo más influenciado por la heterogeneidad espacial a pequeña escala (ZAS 2006) que los caracteres alométricos de reproducción (SANTOS-DEL-BLANCO et al. 2012). Dado que sólo las diferencias genéticas pueden promover el cambio evolutivo, una mayor correlación entre genotipo y fenotipo implica una mayor eficiencia de los fenómenos de selección en la naturaleza. En el presente caso de estudio, las heredabilidades ligeramente superiores



para el tamaño umbral de reproducción femenino que para caracteres de crecimiento (altura, biomasa) hacen pensar en una selección más eficiente para caracteres reproductivos que de crecimiento, aunque el criterio de selección natural sea el número de descendientes de un individuo que llegan a edad adulta (*fitness*). La mayor diferenciación entre poblaciones encontrada para caracteres reproductivos encaja con esta visión.

Sin embargo, ambos conjuntos de caracteres no son independientes entre sí sino que su expresión en los organismos depende de unos recursos finitos por los que las funciones compiten. Estas restricciones se conocen como compensaciones o tradeoffs. Para un nivel de recursos fijo, un aumento en la inversión en reproducción suele implicar una menor inversión en crecimiento (ROFF 2000). A nivel individual, las compensaciones pueden no ser evidentes puesto que no todos los árboles tienen acceso al mismo nivel de recursos. Así dentro de una población, un árbol con acceso a más recursos puede alcanzar un mayor tamaño y al mismo tiempo producir un mayor número de conos que otro. Esto a su vez, produce una correlación positiva entre crecimiento y reproducción para el conjunto de la población (DE JONG & KLINKHAMER 2005). De hecho, los tratamientos encaminados a aumentar la producción de piñas en una masa (claras, podas, fertilización), explotan esta relación entre mayor tamaño y mayor producción de piñas (DE LAS HERAS et al. 2007, GONZALEZ-OCHOA et al. 2004, ORTIZ et al. 2011). Por el contrario, a nivel familiar o de poblaciones, una correlación negativa indica incompatibilidades más o menos fuertes entre funciones a nivel genético. De esta forma, una selección o mejora genética para crecimiento implica una menor inversión en reproducción; aunque, dado que la correlación no es perfecta (menor que 11), sería posible la selección para el aumento en crecimiento y reproducción simultáneamente.

# 6. Conclusiones

Los datos sobre caracteres reproductivos en el inicio de la reproducción en poblaciones españolas y del resto del rango de distribución de *P. pinaster* y *P. halepensis* obtenidos en ensayos de ambiente común han permitido describir un importante grado de diferenciación entre poblaciones, normalmente ligada a las condiciones ambientales de origen y que ilustra la acción de procesos de selección natural. Bajo un nuevo paradigma de gestión enfocado a aumentar la resiliencia de las masas (DE LAS HERAS et al. 2007) tienen también cabida actuaciones por parte de la genética forestal, informando sobre las cualidades de los materiales de base, describiendo patrones ecotípicos y posibilitando la mejora genética para el aumento de la resiliencia (*breeding for resilience*).

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