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Genetics of *Pinus pinaster* Aiton with Cytoplasmic and Nuclear Markers

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Abstract

This thesis summarizes and discusses results of three studies in which biochemical and molecular techniques were used to study the genetic variation in *Pinus pinaster*. In particular, the investigation focused on: (i) the within- and among-population genetic diversity in the region hypothesised as a putative refugium for the species during the last glaciation; (ii) the comparison of nuclear and cytoplasmic estimates of diversity within and between two regions of the species; and (iii) the design of a test for provenance identification using knowledge about the levels of genetic variation between the two regions.

The distribution of the genetic variation of *P. pinaster* in Portugal, as revealed by chloroplast microsatellites (cpSSR), indicated that there are low levels of differentiation among populations and that the diversity is found mainly within populations. No discernible geographic pattern was found. Evidences of strong anthropogenic influence associated with extensive gene flow could explain these findings. Fossil, charcoal and palynological records supported the presence of the species in Portugal before and during the last glaciation; therefore, the hypothesis of a putative refugium in this country cannot be excluded.

The genetic variation of 24 populations from France and Portugal was investigated with amplified fragments length polymorphisms (AFLPs) and cpSSRs. Both types of markers could discriminate between the two provenances and the diversity of the French provenance was higher compared with that from Portugal. Similar differentiation estimates were found with nuclear and cytoplasmic markers. Extensive gene flow could account for this result, but higher mutation rates and homoplasy at cpSSR loci are not to be excluded. Despite the different modes of inheritance, a high correlation was found between the genetic distance matrices with both types of markers, which suggests that migration surpassed genetic drift in moulding the genetic structure of this species in the regions studied.

A provenance diagnostic test was designed, based on cpSSRs, to screen the putative origin of stands of *P. pinaster* in southwestern France and compared with the currently used terpene-based test. Five stands of unknown origin were diagnosed with both tests. The cpSSR-based test proved to be faster and more accurate to determine if stands were of French or northwest Iberian (Portugal and Galicia) origin. The result obtained was probably due to the higher capability of the DNA-based markers to discriminate between both provenances, compared to that of the terpene markers.

Key words: cpSSR, AFLP, terpenes, genetic variation, provenance identification, *Pinus pinaster*.

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**In memory of my father
to my daughter Filipa
and my mother**

SEXTO / D. DINIS

Na noite escreve um seu Cantar de Amigo
O plantador de naus a haver,
E ouve um silêncio murmuro consigo:
É o rumor dos pinhais que, como um trigo
De Império, ondulam sem se poder ver.

Arroio, esse cantar, jovem e puro,
Busca o oceano por achar;
É a fala dos pinhais, marulho obscuro,
É o som presente desse mar futuro,
É a voz da terra ansiando pelo mar.

Fernando Pessoa
in *Mensagem*

Abstract

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Contents

Introduction, 7

An insight into population genetics, 7

Pinus pinaster Ainton: origin, taxonomy and biology, 9

Distribution area, 10

Migration pathways of *Pinus pinaster* in Europe, 11

Human impact in the Mediterranean region forests, 14

Seed certification, 16

Objectives, 18

Methods, 19

Terpenes, 20

PCR-based markers, 21

Dominant markers, 21

Microsatellite markers, 24

Summary of the results, 27

Genetic variation of *Pinus pinaster*, 27

Variation within populations, 27

Variation among populations, 28

Nuclear versus cytoplasmic genetic variation, 29

Reproductive material identification and certification, 31

Closing remarks, 32

References, 34

Acknowledgments, 45

Appendix

This thesis is based on studies reported in the following papers, which will be referred to in the text by the corresponding Roman numerals:

- I. Ribeiro M.M., Plomion C, Petit R, Vendramin G.G. & Szmidt A.E. (2001) Variation of chloroplast simple-sequence repeats in Portuguese maritime pine (*Pinus pinaster* Ait.). *Theoretical and Applied Genetics*. 102(1):97-103.
- II. Ribeiro M.M.*, Mariette S.*, Vendramin G.G., Szmidt A.E., Plomion C. & Kremer A. (2001) Comparison of maritime pine (*Pinus pinaster* Ait.) diversity estimates using SSRcp and AFLP data. (Manuscript)
- III. Ribeiro M.M.*, LeProvost G.*, Gerber S., Vendramin G.G., Anzidei, M., Decroocq S., Marpeau A., Mariette S., & Plomion C. (2001) Origin identification of maritime pine stands in France using chloroplast simple-sequence repeats. *Annals of Forest Science*. (Accepted)

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* To be considered as joint first authors

Introduction

Pithys, a Greek nymph had two lovers, Boreas and Pan. Boreas became jealous and threw Pitys against a rock ledge. She turned instantly into a pine, and resin drops are her tears...
"Greek Myth"

An insight into population genetics

Population genetics, which studies the consequences of genetic information transmission, requires information about changes in the frequencies of genes within and among populations, and aims at understanding the influence and interaction of different factors of evolution in shaping genetic variation. Among the candidate explanations are a balance between natural selection, mutation, gene flow and drift (Stearns and Hoekstra 2000).

Population refers to a group of organisms of the same species living within a restricted geographical area where random mating is potentially possible. A precise definition of the term is difficult and varies from species to species due to the usual non-random pattern in the spatial distribution of the individuals, i.e., due to *genetic geographical structure*. Population subdivision can be caused by environmental patchiness. Therefore, local interbreeding units are defined as *local populations*, *sub-populations* or *demes*, and they constitute the units of population genetics. The level of *genetic differentiation* existing among local populations measures the genetic structure (or population subdivision) of a species, i.e., the frequencies of the alleles may differ from one local population to the next (Hartl and Clark 1997).

In any population, the *genotype frequencies* are determined largely by the patterns in which genotypes of the previous generation come together to form mating pairs. In *random mating*, genotypes form mating pairs in the proportions expected by chance alone. The expectations of genotype frequencies under random mating are described by the *Hardy-Weinberg equilibrium* (HWE). This equilibrium can be disturbed by a number of factors such as selection, mutation, migration and genetic drift, then allele frequencies will change over time (see e.g. Weir (1996) for details).

Several processes create new types of genetic variation or allow the reorganization of the previously existing variation either within genomes or among populations. The ultimate source of genetic variation is *mutation*, that is any heritable change in the genetic material. It includes a change in the nucleotide sequence or a chromosome rearrangement, such as an inversion or translocation. This process provides, by itself, a very weak possibility for changing allele frequencies. Mutation is a *rare event* (10^{-4} to 10^{-6} per gene and per generation for

wild-type genes), but in a large population there are many genes at risk of mutation. *Recombination* brings mutations of different parts of the genome together or conversely, splits portions of the chromosome that were previously together. *Gene flow* is the introduction of genes from one population to another by mating or migration. Like mutation, this process introduces new genetic variants into local populations. Gene flow enables mutations to spread among populations and sets the limit of genetic divergence among populations; moreover, it can prevent local differentiation if selection is weak and the distance that genes move in each generation is large (Stearns and Hoekstra 2000). Gene flow levels vary greatly among species, populations and seasons, but the isolation by distance is frequently not enough to prevent its homogenizing effect against genetic drift and diversifying selection (Ellstrand 1992). Nevertheless, a small amount of gene flow is usually not sufficient to disperse rare alleles among populations; therefore, rare alleles are often unique to one or few populations (Hartl and Clark 1997).

The random fluctuation of allelic frequencies in finite populations due to non-representative sampling of genes from one generation to another is called *random genetic drift*. This phenomenon changes the distribution of genetic variation in two ways: (i) the decrease of variation within populations (loss of heterozygosity and eventual fixation of alleles), and (ii) the increase of differentiation among populations. The effect of this process increases with decreasing population size and depends on allele frequencies of the parental populations (Ellstrand and Elam 1993). Wright (1931) predicted that genetic drift would substantially alter the organization of genetic variation of populations when it is much greater than mutation rate and selection. Populations founded by few individuals do not contain a representative sample of genes in the parental population (*founder effect*). Some alleles that may be completely absent or rare in the parental population can reach high frequencies in the new population simply because they were present in the founders. Similarly, if the composition of the population changes dramatically as it passes through a *genetic bottleneck*, many alleles are lost and others rise to high frequency (reviewed by Barrett and Kohn 1991).

According to the *neutral theory* (Kimura 1968) many mutations have so little effect on the organism that their influence in survival and reproduction is negligible. The frequencies of neutral alleles are not influenced by natural selection, but by genetic drift. Therefore, they have no particular role in the adaptation to new environments, which makes them particularly suitable to trace the geographical genetic structure of populations among other purposes.

The major questions of population genetics are: how to measure the genetic variation in populations and how much it affects individual reproductive success. In the middle sixties, molecular methods started to be used and they increased the possibility to obtain estimates of genetic variation at previously “invisible” loci. The amount of genetic variation is measured by *genetic diversity* or “heterozygosity”, defined as the probability that two alleles chosen at random at a locus in the population are different. *Molecular methods* do not solve the problem

of deducting the genetic variation from observed phenotypic variability: they circumvent the problem. They tell us how much genetic variation is present in a particular part of the genome, but they do not tell us how this genetic variation affects phenotypic variation (Stearns and Hoekstra 2000).

***Pinus pinaster* Ainton: origin, taxonomy and biology**

Pines belong to an old genus – the oldest in the whole family of *Pinaceae* – which, originated in the middle Mesozoic (190-136 My BP, million years before present) in middle latitudes. During the Cretaceous (136-65 My BP), the genus was already differentiated into two subgenera, and pines were widely distributed throughout the Northern Hemisphere. In general, pine populations were fragmented and displaced during the Eocene due to major climate changes (54-34 My BP). In the terminal Eocene, temperature and humidity decreased drastically leading to angiosperm taxa extirpation and expansion of pines and other cool and dry adapted taxa in middle-latitude locations. Fossil records indicate that pines rose in abundance throughout middle latitudes in North America, Europe and Asia in the Miocene (26-7 My BP); the direct ancestors of many modern pines can be traced to Miocene pines (Millar 1998 and references therein). There are indications that Mediterranean pines migrated to the region from eastern Asia along the mountain ranges that once extended north of and parallel to the Himalayas (Mirov 1967).

Pinus contains more species than any other genus of conifers (Little and Critchfield 1969), and they all share uniformity in chromosome number ($2n = 24$) (Sax and Sax 1933). More than 40 different classification systems have been proposed for this genus (Millar 1993) depending on the criteria used (e.g., Shaw 1914; Mirov 1967; Little and Critchfield 1969; Farjon 1984). This genus is usually divided into two subgenera *Strobus* (= *Haploxyton*, soft pines) and *Pinus* (= *Diploxyton*, hard pines), which are further divided into sections and subsections (Little and Critchfield 1969; Farjon 1984). According to Little and Critchfield (1969), *Pinus pinaster* belongs to the *Pinus* subgenus, subsection *Sylvestris*.

The evolution of the genus *Pinus* has been studied recently using different approaches, in particular molecular markers (e.g., Strauss and Doerksen 1990; Wang and Szmidt 1993; Farjon 1996; Wang *et al.* 1999; Wang *et al.* 2000). According to a study based on sequence divergence of chloroplast regions made with Eurasian pines (Wang *et al.* 1999), the Mediterranean pines formed one strongly supported clade within the subgenus *Pinus*, and within that clade, *P. pinaster* grouped with *P. canariensis* and *P. pinea*, but with a weak bootstrap support.

Pinus pinaster reaches a height of about 35 m. The needles are thick, rigid and shiny grey-green, with conspicuous rows of stomata on all sides. They are grouped in pairs, 15-20 cm long, with a smooth surface, and a half-circular cross-

section. Male strobili develop on the lower part of the new shoots, whereas female strobili form in the whorl around terminal buds. The cones are 10-20 cm long and remain in the branches for several years. Young trees 6-7 years old can start to produce cones on upper shoots. The seeds are about 1 cm long, with a 2-3 cm long wing (Farjon 1984).

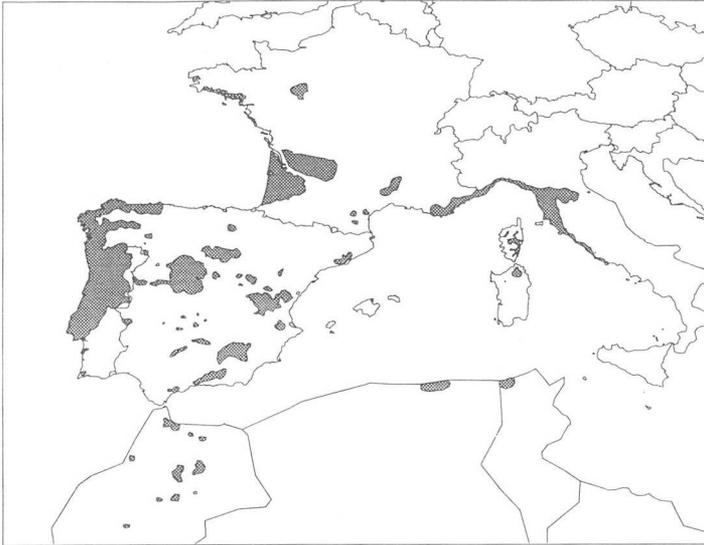


Figure 1. Natural distribution of *P. pinaster*. (After Baradat and Marpeau-Bezard 1988)

Distribution area

Some studies have shown that climatic changes during the Quaternary (2.4 My to present) in Europe have played a major role in shaping the phylogeography of European plant and tree species by contracting and expanding their natural ranges (Bennett *et al.* 1991; Hewitt 1996; Hewitt 2000). The successive glacial-interglacial oscillations of the Pleistocene (2.4-0.01 My BP) have moulded the range of plant species, by isolating refugia, which subsequently provided the source for new colonization (Comes and Kadereit 1998; Taberlet *et al.* 1998). By compiling several data sets available for various European tree species, Taberlet *et al.* (1998) were able to identify some general trends according to the location of their refugia in southern Europe, but they concluded that each species would exhibit a particular phylogeographic pattern. In the case of *P. pinaster*, it has been hypothesised that the actual distribution of this species is the result of events that occurred during the last glaciation (0.7-0.01 My BP) (Baradat and Marpeau-Bezard 1988). Nevertheless, the distribution of *P. pinaster* has also been greatly modified by human activities during historic times throughout the Mediterranean Basin until the recent expansion of its cultivated range (Barbero *et al.* 1998; Le Maitre 1998).

According to Scott (1962) and Farjon (1984) *Pinus pinaster* occurs naturally in southwest Europe and northwest Africa between latitude 31° and 46° N and longitude 9° and 13° E; from southwest Morocco to the mouth of the Gironde in France, and from the west coast of Portugal to the west coast of Italy. In France, Algeria, Tunisia and Italy the distribution is mainly coastal, but in Portugal, Spain, Morocco and Corsica this species grows from near the coast to far inland and high into the mountains. This species is distributed throughout its range area in a discontinuous way, due to geographic isolation of populations and to the ancient human impact in the Mediterranean Basin (Fig. 1). The areas occupied by the species in different countries are shown in Fig. 2. *Pinus pinaster* has been planted in many other places in Europe and it has often become well established outside its natural range, especially in coastal regions. *Pinus pinaster* is also found in Australia, South Africa and New Zealand as an exotic species, where it constitutes extensive and successful stands.

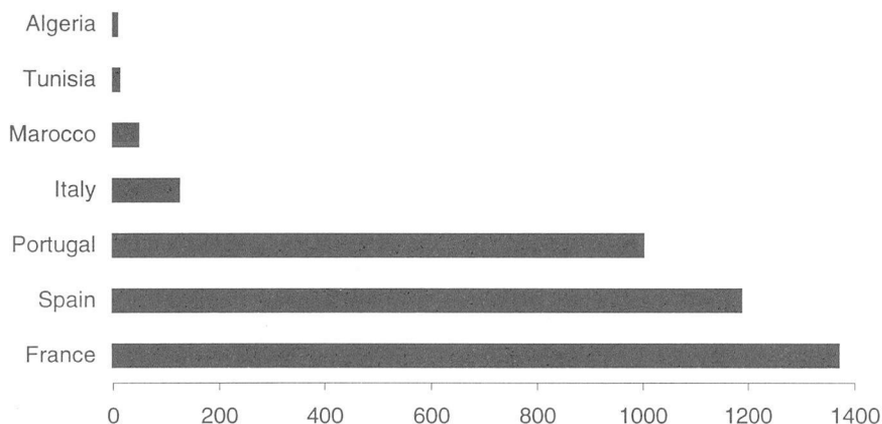


Figure 2. Areas occupied with *P. pinaster* in the countries of its natural range (x 1000 ha). (<http://pinus.dgf.min-agricultura.pt/estatistica/invent.htm>; Alía *et al.* 1996; Deroy 2000)

Migration pathways of *Pinus pinaster* in Europe

Tree species show different patterns in the distribution of genetic diversity within and among populations (Hamrick *et al.* 1981). As discussed earlier, several factors are responsible for moulding the genetic variation patterns: gene flow, selection, genetic drift, human activities and climatic changes. However, the relative significance of these factors is likely to vary greatly among populations and species (Stearns and Hoekstra 2000).

The genetic variation of *P. pinaster* has been studied using various ways. Intraspecific variation in the species has been investigated in numerous provenance trials established in different countries (Harfouche and Kremer 2000

and references therein), and these experiments showed that morphological and adaptive traits vary significantly among provenances. Several range-wide diversity studies have focused on terpenes, isozymes, denatured proteins and chloroplast microsatellites (Baradat and Marpeau-Bezard 1988; Bahrman *et al.* 1994; Petit *et al.* 1995; Vendramin *et al.* 1998). Some other studies have been undertaken at a regional level using isozymes, AFLPs (amplified fragment length polymorphisms) and nuclear microsatellite markers (Castro 1989; Salvador *et al.* 2000; González-Martínez *et al.* 2001; Mariette *et al.* 2001b).

Based on terpene markers, palynological and paleoclimatological records, Baradat and Marpeau-Bezard (1988) discriminated three major groups of *P. pinaster* populations: the Atlantic group, comprising populations from southwestern France, Portugal, and Galicia in Spain; the Mediterranean group, extending from central Spain to the Ligurian coast in Italy; and finally the North African group that includes stands from Morocco, Algeria and Tunisia. In another study, Bahrman *et al.* (1994) included eastern Spain in the Atlantic group. Using mitochondrial DNA markers, C. Burban (unpublished manuscript) discriminated three groups: the Moroccan (Rif and Atlas), the Occidental (Iberian Peninsula and southwestern France) and the Oriental group (southeastern France, Corsica, Sardinia, Italy, Pantelleria, Tunisia and Algeria).

The typically scattered distribution of this species may have prevented or limited gene flow among the different groups of populations, causing high genetic divergence among populations due to genetic drift (Baradat and Marpeau-Bezard 1988; Bahrman *et al.* 1994; Petit *et al.* 1995; Vendramin *et al.* 1998). Nevertheless, gene flow and human activity were probably responsible for the low differentiation found at a fine geographic scale (Castro 1989; Mariette *et al.* 2001b).

The presence of a centre of origin of *P. pinaster* in the southwest of the Iberian Peninsula at the end of the Pliocene (3 My BP) was hypothesised by Baradat and Marpeau-Bezard (1988) and supported by fossil findings (Teixeira 1945). The authors drew a different picture for the migration pathways of the species before and after the last glaciation. The preglacial hypothesis supposes the presence of three distinct pathways towards the north of Portugal, Spain and France, towards the south of Spain, France and Italy, and towards the north of Africa (Fig. 3). The postglacial hypothesis assumes that migration occurred only along the first two pathways. Moreover, the authors claim that the successive ice ages during the Pleistocene had several times stopped the northerly advance of *P. pinaster*, and eventually diminished its presence to scattered refugia in the south of the Iberian Peninsula, particularly during the Pleniglacial (0.35 My BP).

According to the study made by Vendramin *et al.* (1998) based on chloroplast microsatellites, a more complex picture of the possible migration pathways was drawn. Two main reservoirs of haplotypic diversity (Landes and Pantelleria) were identified (Fig. 3). The Pantelleria population (or North of Africa populations) might have represented a starting point of the migration process. Pantelleria could

represent an ancient population originating in the preglacial period from refugia located in the central part of North Africa, from which the migration took place towards West and East. The French area might have represented refugia from which the migration towards Italy began. Conversely, low levels of within-population diversity were observed in the Portuguese populations (Alcácer and Monção) with characteristics of recent establishment (possible founder effects). Such a reduction in genetic diversity with increasing distance from a refugium is a general phenomenon to be expected from repeated population bottlenecks at the advancing edge of a range in any species during postglacial expansion (Hewitt 1996).

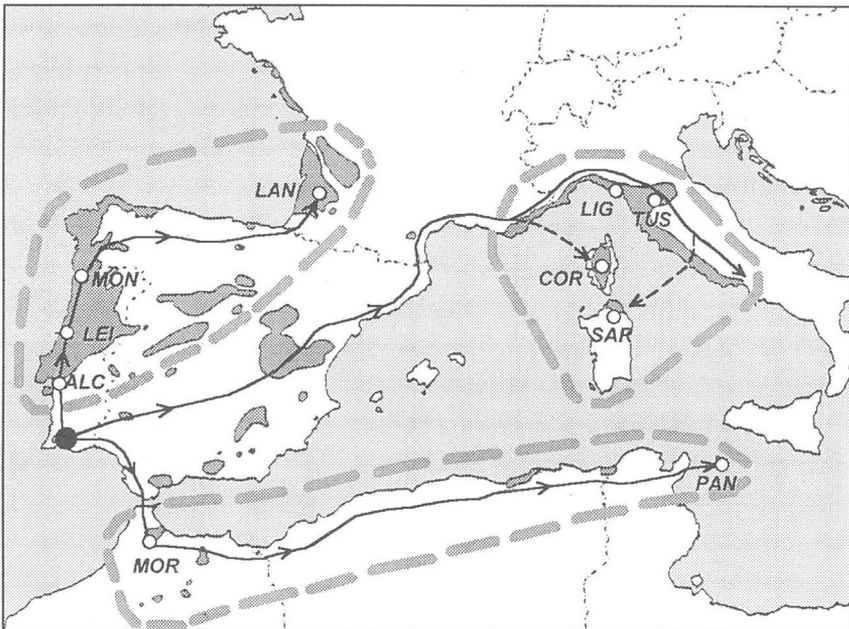


Figure 3. The three major groups of populations and preglacial migration pathways hypothesis considered for *P. pinaster*. ALC=Alcácer, LEI=Leiria, MON=Monção, LAN=Landes, PAN=Pantelleria, LIS=Liguria, TUS=Tuscania, COR=Corsica, SAR=Sardinia, MOR=Morocco. (Courtesy of Vendramin *et al.* 1998)

The Iberian Peninsula has been reported as glacial refugium for numerous plant species, (Hewitt 1996; Comes and Kadereit 1998)) and it is one of the most important native areas of *P. pinaster*. Salvador *et al.* (2000) observed a generally higher level of allozyme diversity in the Spanish populations than the Portuguese population included in their study, and hypothesised a migration pathway from Spain to Portugal. The authors used six other populations from Portugal from another study (Castro 1989) and recomputed the data by using the same loci. Afterwards, Salvador *et al.* (2000) concluded that the idea of refugia in Portugal could not be supported because no special variants were detected. In conclusion,

the authors suggested the disappearance of *P. pinaster* from that area during the last glaciation.

The phylogenetic analysis made with allozyme markers by González-Martínez *et al.* (2001) showed a high geographical structure in the Iberian Peninsula. The northwestern populations form a cluster and the southeastern populations another, in accordance with the study made by Baradat and Marpeau-Bezard (1988). The authors observed the highest levels of diversity in the eastern and the southern populations and an important reduction of gene diversity in the northwestern range of the species in the Peninsula. Nevertheless, a putative refugium in Portugal was not excluded, because *P. pinaster* could have survived during the last glaciation in sheltered areas at low altitude close to the Atlantic Ocean in Portugal, as suggested by Figueiral (1995) based on charcoal records.

Human impact in the Mediterranean region forests

The Mediterranean Basin is characterized by traditional human impacts on the forest (Thirgood 1981) and the modification of genetic diversity of species by human activity (Ledig 1992). The land use by humans over millennia has played a dramatic role in shaping the region's vegetation. In no other part of the natural range of pines has there been such a complex interplay between this genus and humans. The effects, however, have not been the same throughout the entire Mediterranean range, being more important in coastal areas and fertile soils. Various ecological factors, such as forest fires, drought, soils, etc., also played a major role in the adaptation of *P. pinaster*, in the isolation between stands, and, to some extent, in the genetic variation of the species (Barbero *et al.* 1998).

People and pines have had a long association. The first evidence of hominid habitation within the natural range of *Pinus* has been dated to about 1.4 My BP on the Eastern Shore of the Mediterranean Sea (Wood and Turner 1995). According to Le Maitre (1998) and references therein, by the time hominids encountered pines, the different species occupied most of their current range. Mediterranean-type climates, with their dry summers and recurrent fires, were also well established. Key events between humans and pines took place: the marked climatic fluctuations during the Pleistocene and Holocene (10 000 BP until present); the increasing use of fire that is essential for maintaining pine populations and reducing competition from hardwood species; the population growth that followed the domestication of agricultural crops; the copper and iron smelting that requires large amounts of wood; and the forest clearing necessary to produce open fields for agriculture and pasture (Fig. 4). Although population densities were low, human impacts during the prehistoric period were substantial and could have altered the abundance of pines, particularly in the Mediterranean Basin (Barbero *et al.* 1998).

The use of timber for the sea-borne trade and military protection further increased during the Graeco-Roman period. Also, the rapid population growth, particularly

around cities, caused higher wood demands for housing, domestic use, bridges and roadways (Fig. 4). The emergence of agriculture and biological sciences with higher control over plant cultivation accelerated forest depletion, but regulation of forest cutting was common both for wood conservation and religious purposes (Thirgood 1981; Le Maitre 1998). The earliest evidence of humans altering the distributions of pine species through cultivation dates from the Graeco-Roman period. Palynological studies suggest that pine populations expanded locally in degraded areas during that period, but humans also spread them, *P. pinaster* being a paradigmatic case (Willis 1992; Klaus 1989).

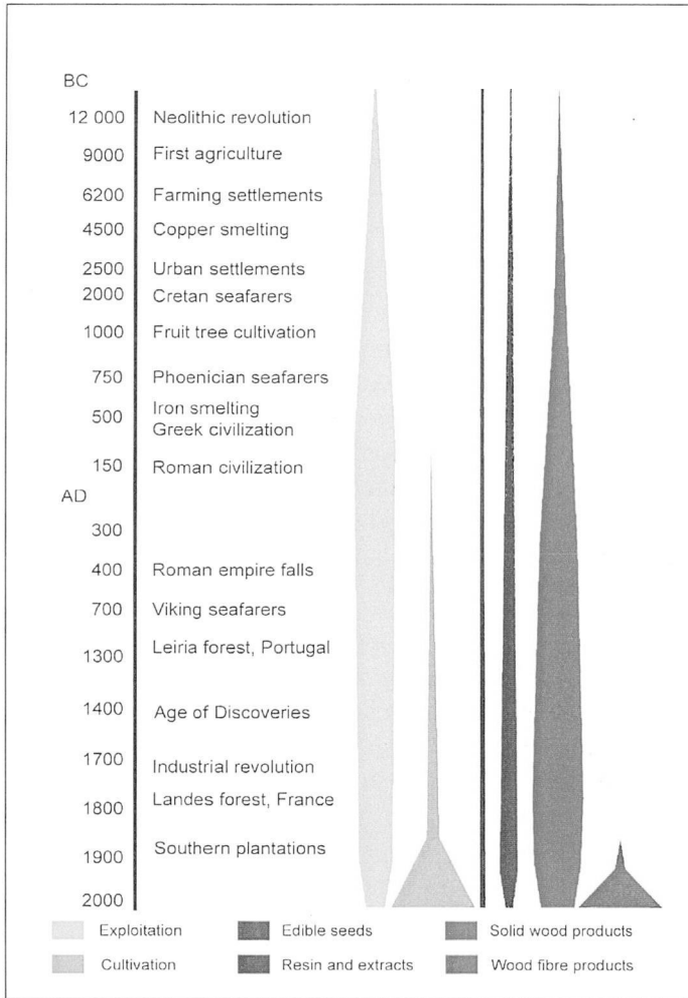


Figure 4. A diagrammatic representation of events in the history of cultivation of pines and the relative importance of the products or categories used by Man. (Adapted from Le Maitre 1998)

Growing concerns about forest depletion led to the first recorded large-scale reforestation. Natural forests in Portugal had suffered intense use before the 14th century due to fire, agriculture, grazing, mineral exploitation, wood consumption and shipbuilding; therefore, in 1310, King Don Dinis issued a law to protect pine forests (Scott 1962; Buting and Rego 1988). In the 13th century Cistercian monks established the “Leiria’s pineyard” to stabilise the sand dunes, but many others were established elsewhere for soil protection and to fight desertification of the country (Mattoso and Sousa 1993). From the 15th century onwards, the naval and merchant fleets of Portugal, Spain, France and Great Britain also consumed large quantities of wood (Thirgood 1981). According to Le Maitre (1998), global timber trade increased significantly in the 18th century and sawmill industry expanded, encouraging forest harvesting. Forest nurseries were established for large-scale planting in Portugal and by the end of the 18th century and the beginning of the 19th century, *P. pinaster* was clearly expanding its presence and supplanting other species or invading non-cultivated areas (N. Devy-Vareta, personal communication). In France, attempts at reforestation were made as early as 1500, and *P. pinaster* was used to stabilise the dunes near Bordeaux in 1713. Reforestation of the sandy coastal plains of Landes in southwestern France continued until a significant area of the region was covered (Scott 1962).

In Europe, the two World Wars were critical and devastating periods for the forests, but the development of petroleum-based synthetics has, to a great extent, replaced wood-based products. Nevertheless, paper consumption increased notably with the growing information demands of the 20th century. In the Mediterranean Basin, the primary concern has been the protection of soil and reforestation of degraded areas (Thirgood 1981). For example, in Portugal, about 400 000 ha of *P. pinaster* had been planted between 1900 and the 1960s (Devy-Vareta 1993).

A particular feature of *P. pinaster* is its ability to grow well on very poor sandy soils, low in nutrients, while tolerating summer drought, winter flooding, and sea winds. Under favourable climate, such soils give a good yield of wood for lumber or pulp, and also resin (Scott 1962; Farjon 1984). Those characteristics played a central role in its preferential use for reforestation. Therefore, since at least the 20th century, reforestation programmes in the *P. pinaster* range area have notably spread this species. The present situation, therefore, is a complex of indigenous populations and planted forests of unknown origin (Devy-Vareta 1993; Alía *et al.* 1996; Barbero *et al.* 1998; Salvador *et al.* 2000).

Seed certification

Human settlement and action over the centuries have influenced forests in Europe, particularly in the Mediterranean area as discussed in the preceding paragraphs, and planted forests constitute a major part of the resource of some countries. The

concern about the deterioration of forests throughout Europe led to an increasing awareness of their economic, ecological, social and cultural value.

Human impact on forests alters the genetic structure of tree species in several ways; an important example concerns the introduction of seeds from other regions. Because introduced seeds produce, in general, less adapted stands than native ones, introductions may provoke economical losses and further affect the productivity of autochthonous stands. The *origin* of an *indigenous stand* is defined as the place in which the trees are growing and the origin of a *non-indigenous stand* is the place from which the reproductive material was originally introduced. *Reproductive material* may comprise fruits, seeds, pollen, scions or tissues for tissue culture. Planting non-indigenous reproductive material may alter local patterns of variation by influencing adjacent indigenous stands due to pollen and seed dissemination. The consequences are expected to be more important where intensive plantings were made by using reproductive material from regions with very different environmental characteristics from those at the site of material introduction (Jones and Burley 1973).

According to Zobel and Taberlet (1984), seed certification has been a concern that started as early as the beginning of the 20th century in Japan with *Cryptomeria japonica*. There are many meanings for seed certification, and different methods for the collection and handling of seeds from forest trees. Some authors regard certification as the correct *labelling* in which the seed size, purity, germination and other information about the seed is given, others consider that the information about where the seed was obtained, *source certification*, should also be included (Barber *et al.* 1962; Zobel and Taberlet 1984). According to Jones and Burley (1973), *seed certification* is an official statement that a seed lot conforms to certain standards, which may include specific identity, origin, genetic characters and seed purity. For forestry, seed certification systems have been developed largely to provide labels and records that give officially authenticated details of *identification* of the seeds, which involves the step of identifying its origin, and also its *quality*, meaning the genetic superiority, when that information is available. Therefore, reproductive material identification and certification in forestry has become a relevant issue (Barber *et al.* 1962; Matthews 1964; Jones and Burley 1973).

Morphological data and biochemical markers (terpenes, isozymes and denatured proteins) have all been used for provenance identification and seed certification in forest trees (e.g., Falkenhagen 1985, Boisseaux 1986, Bahrman *et al.* 1994, Espinel *et al.* 1995). Molecular markers based on nuclear and organelle DNA analysis have also been used lately for those purposes (e.g., Szmidt *et al.* 1988, Aragones *et al.* 1997, Sinclair *et al.* 1998, Bucci and Vendramin 2000). For example, in France, a law under the supervision of the Ministère d'Agriculture et Forêts regulates the collection of commercial *P. pinaster* seed-lots in the Aquitaine region (Réglement technique récolte pin maritime, Arrête du 8 février 1990). Candidate stands for seed collection in the Aquitaine region must have their origin certified using a diagnostic test developed by Baradat and Marpeau-

Bezard (1988), based on a discriminant analysis of terpene profiles. The term *provenance* has different meanings, reviewed by Jones and Burley (1973); in this study, it refers to the original geographical source of a given lot of plant reproductive material.

Recently, consensus on reproductive material certification in forestry is under discussion in Europe, allowing regional differences and specificities. The Third Pan-European Ministerial Conference on the Protection of Forests in Europe, held in Lisbon, 2-4 June 1998, adopted criteria for sustainable forest management, one of them being the maintenance, conservation and appropriate enhancement of biological diversity in forest ecosystems. This resolution involves the establishment of standards for forest reproductive material certification. For reforestation, indigenous species and local provenances that are well adapted to site conditions should be preferred. Introduced species or provenances should be used only after the evaluation of the impacts on the ecosystem and on the genetic integrity of indigenous species or provenances. For further details the following web page should be consulted: <http://www.pefc.org/content.htm>.

Objectives

The purpose of this thesis was to study the genetic structure of *P. pinaster* at a regional level. In particular, the investigation focused on a practical application to be used in the forest management of this species.

The main objectives of the three studies included in this thesis were: (1) to assess the distribution of genetic diversity within and among populations of *P. pinaster* in the region hypothesised as a putative refugium for the species during the last glaciation; (2) to compare nuclear and cytoplasmic estimates of diversity within and between two regions of the species; and (3) to design a test in order to identify the origin of the stands in one region.

In studies reported in Papers **I** and **II**, cpSSR (chloroplast microsatellites) and AFLP (amplified restriction fragment polymorphism) markers were used to provide information on the level and distribution of genetic variation among and within populations of *P. pinaster* at the regional level. Two different approaches were used: the first involved analysis of chloroplast repetitive simple-sequence repeats (cpSSR) (Paper **I**) and the second involved a comparative analysis between the cpSSR and AFLP markers (Paper **II**). In the analysis described in Paper **III**, a cpSSR-based test was developed in order to determine the putative origin of maritime pine stands in the Aquitaine region (southwest of France) and to compare the cpSSR-based test with the test based on terpene profile analysis.

Methods

Knowledge about the genetic structure of a species can be obtained from polymorphic markers that allow the determination of gene and/or genotypic frequencies. Genetic variation of forest trees can also be inferred using the traditional quantitative analysis of morphological traits, but due to the environmental influence, the polygenic character of some traits, and the time and cost to retrieve the information, other methods have been sought to obtain the same type of information (Wang and Szmidi 2000). The differences among individuals can be traced using secondary compounds, such as terpenes and flavonoids, but they also fail to be good candidates due to difficulties of inferring the genotype from phenotypes and to a possible environmental influence (Crawford 1983; Hanover 1992). In the last decades, methods that look at protein polymorphisms (Strauss *et al.* 1992) and the direct analysis of polymorphisms at the DNA level have provided a diverse array of molecular tools for genetic analysis in forest tree populations (reviewed by Morgante *et al.* 1996; Wang and Szmidi 2000).

A *molecular marker* can be defined as a sequence of DNA or a protein which can be readily detected and whose inheritance can be monitored. It is the *polymorphism* of molecular markers that can be used to study genetic diversity. Polymorphism in proteins has been studied, e.g., through *allozymes*, i.e., different molecular forms of an enzyme coded by different *alleles* at one gene locus. Polymorphism can be identified in different types of DNA: nuclear and cytoplasmic or organellar DNA (in the chloroplast, cpDNA and in the mitochondria, mtDNA) (Mitton 1994; Vekemans and Jacquemart 1997; Parker *et al.* 1998).

The desirable properties of a marker are: polymorphic expression; *codominant* inheritance (the different forms of a marker should be detectable in diploid organisms to allow discrimination of homozygotes and heterozygotes); even distribution throughout the genome; easy, fast and inexpensive detection; and reproducibility within and between laboratories. No single molecular marker meets all those criteria; the choice of a particular molecular marker will therefore depend on the objectives of the study (reviewed by Karp and Edwards 1997; Parker *et al.* 1998; Szmidi and Wang 2000).

DNA-based markers have a great advantage over terpenes or isozymes, for they provide pure genetic information, since they are not the products of transcription or translation. Other advantages of these markers are the large variety of scales on which evolutionary processes can be studied and their great potential for

detecting variation in all kinds of organisms, in both living and dead tissues (Parker *et al.* 1998).

Markers mostly employed in the current studies are described in the following paragraphs.

Terpenes

Monoterpenes (C_{10} hydrocarbons) and sesquiterpenes (C_{15} hydrocarbons) are two classes of isoprenoid derivatives, which are elaborated from pyrophosphorylated precursors: respectively, geranyl (C_{10}) and farnesyl (C_{15}) diphosphates. These compounds are found in conifers, in particular in the genus *Pinus*, where they accumulate in the resin ducts of different tissues (needles, primary cortex, and conducting tissues). Terpenes constitute 20% or more of the volatile fraction of oleoresin (Baradat *et al.* 1995). Terpenoid biosynthesis and metabolism were further reviewed by Chappell (1995).

In the genus *Pinus*, single-gene inheritance of monoterpenes has been demonstrated in several species (Baradat *et al.* 1995 and references therein), including *P. pinaster* (Baradat *et al.* 1972). Marpeau *et al.* (1975) also demonstrated single-gene inheritance of sesquiterpenes in *P. pinaster*. The use of terpenes as markers may present difficulties, as their expression can be affected by the age of the tree and the sample source (position of the sample in the tree and type of tissue used) (Bernard-Dagan *et al.* 1971). In *P. pinaster* the cortex of young but completely lignified shoots, the oleoresin composition remains unchanged (Baradat *et al.* 1972), therefore such plant material is appropriate for extraction of terpenic compounds for analysis. Moreover, the terpenic composition in the cortex of that plant material remains stable as soon as the tree achieves 7-10 years of age (Baradat *et al.* 1991).

Terpene markers have been used to study geographical genetic differentiation in conifers, as reviewed by Müller-Starck *et al.* (1992) and Strauss *et al.* (1992), but they can also be used to study mating patterns, to assess the geographic origin of stands and to study seed orchard pollen contamination (Baradat *et al.* 1991; Coppen *et al.* 1993; Schiller and Genizi 1993; Baradat *et al.* 1995). Nevertheless, terpenoid composition, while strongly inherited, probably plays a role in tree's disease and insect resistance, and is thus undoubtedly subject to natural selection (Hanover 1992). Some terpenes, in particular, are believed to be involved in the resistance to the caterpillar *Dioryctria spendidela* that attacks the bark of *P. pinaster* (Baradat and Marpeau-Bezard 1988). Besides, for terpenoids, only genetic changes that substantially alter gene expression or biosynthetic enzyme activity will be detected; thus, products of many different mutational events will be confounded or undetected (Strauss *et al.* 1992).

PCR-based markers

Dominant markers

In past years, a new generation of markers, based on the *polymerase chain reaction* (PCR), has been developed. PCR is a technique to amplify specific DNA sequences by primer extension of complementary strands of DNA with the action of the thermostable DNA polymerase (Mullis and Faloona 1987; Saiki *et al.* 1988). This technique is very powerful for amplifying tiny amounts of DNA sequences several million times over only in a few hours, involving several cycles of heating and cooling, each time the newly synthesised strand becoming a template for the subsequent replication. Theoretically, the cycling of temperatures increases in an exponential way the amount of the specified sequence (Fig. 5).

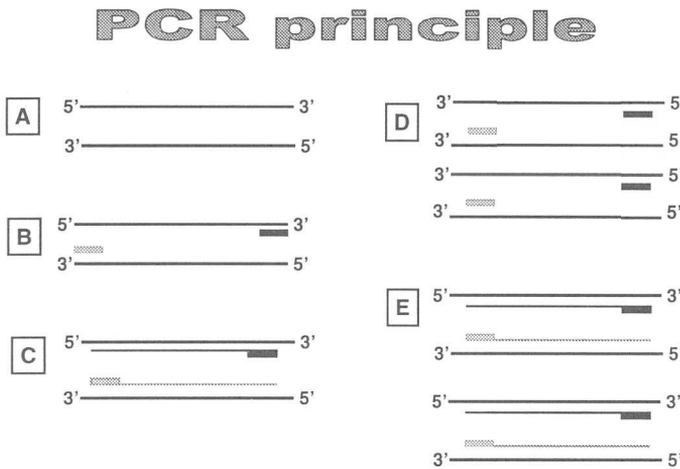


Figure 5. Diagrammatic representation of the PCR principle. A: In the denaturing step, the heat opens the double strand. B: In the annealing step, the cooling allows the primers (in grey and black) to bind the complementary regions. C: In the elongation step, the DNA polymerase synthesises complementary strands. D and E: The cycles of heating and cooling are repeated, each time the newly synthesised strand becoming a template for the subsequent replication.

In 1990, the use of short primers (usually 10 base-long) of arbitrary sequence was initiated to generate PCR amplification products, in low stringency conditions, from genomic DNA (Williams *et al.* 1990; Welsh and McClelland 1990). Depending on the specific conditions of the amplification or product separation and detection, different methods were termed: RAPD (*Random Amplified Polymorphic DNA*, Williams *et al.* 1990); AP-PCR (*Arbitrary Primer PCR*, Welsh and McClelland 1990); or, DAF (*DNA Amplification Fingerprinting*, Caetano-Annolés *et al.* 1991). RAPD analysis is performed at a low annealing

temperature (stringency), implying that the binding of the primer to the genomic DNA is partly non-specific. Therefore, in order to obtain reproducible results, the reaction conditions must be kept strictly constant and the analysis must be made in the same laboratory (Penner *et al.* 1993; Hallden *et al.* 1996; Jones *et al.* 1997; Rafalski 1997). Nevertheless, the method's speed, sensitivity and versatility make it suitable to survey large number of samples in population genetics of forest trees (e.g., Bucci and Menozzi 1995; Nesbitt *et al.* 1995; Schierenbeck *et al.* 1997; Gallois *et al.* 1998; Wu *et al.* 1999). RAPD markers usually show dominant Mendelian inheritance; amplification either occurs at a locus or not, leading to scores of band presence/absence. This means that the heterozygotes and homozygotes for the presence of the band cannot be distinguished (Isabel *et al.* 1995; Lu *et al.* 1995). Moreover, the amplified regions may represent both coding and non-coding sequences (Kazan *et al.* 1993; Lu *et al.* 1997).

Amplified Restriction Fragment Polymorphism (AFLP) is a powerful method for detecting polymorphism throughout the genome, based on a two-step amplification strategy that combines restriction enzymes and PCR (Zabeau and Vos 1993). This highly reproducible technique allows the simultaneous screening of a large number of molecular markers, randomly distributed throughout the genome (Vos *et al.* 1995; Zhu *et al.* 1998).

The genomic DNA is digested with two restriction enzymes, a frequent cutter and a rare cutter (in Fig. 6 *Mse*I and *Eco*RI). Afterwards, two adapters with 3' ends complementary to the sequences recognised by the enzymes are ligated to the DNA fragments, to provide known sequences for the first PCR amplification. In the pre-amplification step, primers with the 5' ends complementary to the adapters, but extended with one to several nucleotides at the 3' ends (two nucleotides in Fig. 6), are used to amplify a subset of the restricted fragments (PCR I). In the selective PCR amplification step, primers with the 5' ends, complementary to the adapters but extended with one to several nucleotides at the 3' ends (referred as selective nucleotides), are used to amplify the selected subset of restricted fragments (PCR II). Polymorphisms are detected by differences in the length of the amplified fragments by polyacrylamide electrophoresis. The amplified fragments are resolved in a sequencing gel and visualised by radioactivity, fluorescence or silver staining.

The AFLP method generates a large number of bands in a single reaction (Vos *et al.* 1995; Powell *et al.* 1996) and gives higher reproducibility compared with the RAPD method (Jones *et al.* 1997). The AFLP technique has been used for a wide range of purposes in tree species, among others, the investigation of genetic diversity (e.g., Winfield *et al.* 1998; Lerceteau and Szmidsztajn 1999; Cervera *et al.* 2000; Mariette *et al.* 2001b), the generation of linkage maps, and the identification of molecular markers linked to phenotypic traits and/or genetic loci (Cervera *et al.* 1996; Marques *et al.* 1998; Travis *et al.* 1998; Cato *et al.* 1999; Marques *et al.* 1999; Remington *et al.* 1999; Arcade *et al.* 2000; Costa *et al.*

2000; Lerceteau *et al.* 2000; Sewell *et al.* 2000). Similarly to RAPD markers, AFLP markers show predominantly dominant Mendelian inheritance (Paglia and Morgante 1998; Lerceteau and Szmidi 1999; Nikaido *et al.* 1999) and they detect variation in anonymous nuclear sequences.

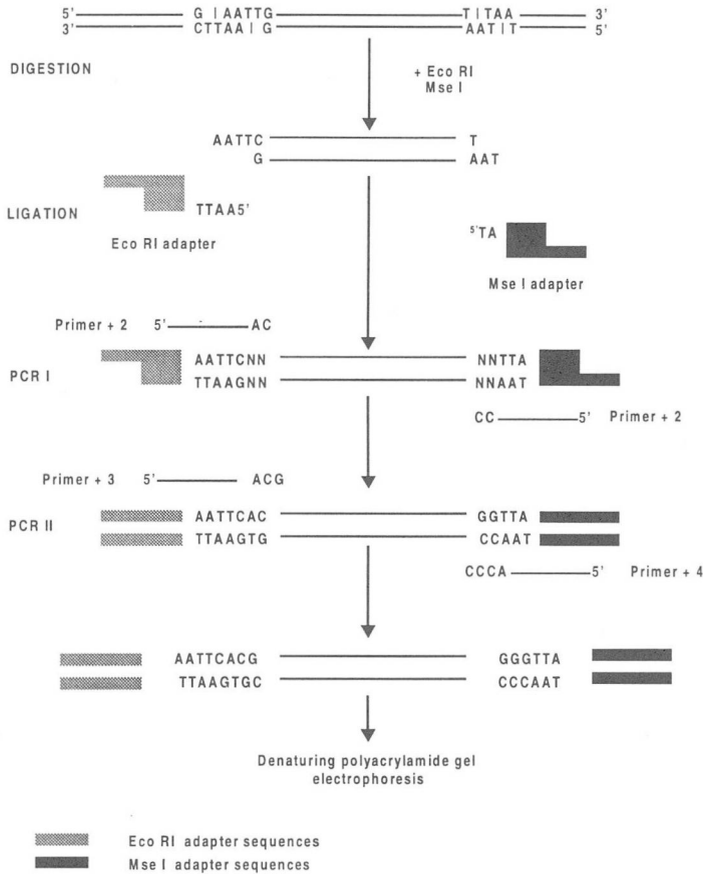


Figure 6. A diagrammatic representation of the AFLP procedure reported in this thesis. See text for explanation.

Only recently has this method been used in conifers because of the large size of their genome, which was responsible for the complicated band pattern, therefore the protocol had to be adapted in order to decrease the number of bands obtained per primer combination. The number of generated fragments can be restricted by changing the nucleotide extensions and/or the type of enzymes used in the digestion of the DNA (methylation sensitive or insensitive) (e.g., Paglia and Morgante 1998; Lerceteau and Szmidi 1999; Cervera *et al.* 2000; Costa *et al.* 2000). AFLPs are more technically demanding than RAPDs, but their automation

and the availability of kits (e.g., Lerceteau and Szmidi 1999) made possible their use on a larger scale.

PCR-based methods such as RAPD and AFLP markers are more easily obtained than most non-PCR alternatives and their analysis does not require sequence information or laborious cloning. However, since most of these markers are dominant, the genotypes are not unambiguously traced, therefore biases are introduced in the estimation of population-genetic parameters (Lynch and Milligan 1994; Isabel *et al.* 1995; Szmidi *et al.* 1996; Isabel *et al.* 1999; Krutovskii *et al.* 1999; Zhivotovsky 1999). According to Lynch and Milligan (1994), dominant markers can be used to estimate unbiased population genetics parameters, provided that the loci with low frequency of the null allele are pruned from the analysis. Also Zhivotovsky (1999) developed a new approach to overcome the dominance situation for these markers. Isabel *et al.* (1999) refers that the reliability of RAPDs fingerprints in estimating population structure can be improved if prior knowledge exists of the mating system and levels of populations structuring, and if the fragments meet some polymorphism criterion (Lynch and Milligan 1994).

Microsatellite markers

A new type of marker, known as *simple-sequence repeat* (SSR) or *microsatellite* has been developed based on DNA sequence variation. This marker is based on tandem DNA repeats characterised by short motifs (1 to 6 bp), repeated from two to many thousands of times (Tautz 1989). A different allele occurs at a SSR locus as a result of changes in the number of times a core element is repeated, altering the length of the repeated region. Differences in length at a SSR locus are detected with DNA amplification by PCR using two oligonucleotide primers that complement unique sequence flanking at the SSR locus. Polymorphism is detected by electrophoretic separation of fragment sizes that can differ by as few as two base pairs.

Current research suggests that the length variation between alleles at a SSR locus are created by slippage of DNA polymerase during the replication of tandem repeats followed by failure of DNA mismatch repair to restore the original sequences (Strand *et al.* 1993). Microsatellites are very useful because they are codominant and highly-polymorphic markers, but their identification is a very expensive and time-consuming process, which generally requires the construction and screening of a genomic library. Known primers are not likely to amplify the same locus across related taxa, unless the flanking regions were priming sites are located are highly conserved (Ellegren 1992), which happens, usually in closely related species (Kijas *et al.* 1995). Therefore, the success of cross-amplification diminishes with increasing species divergence (Whitton *et al.* 1997).

Single-locus SSR markers have been identified in conifers (Lefort *et al.* 1999 and references therein), but due to their large genome size, this task has proven to be difficult. Moreover, only a small fraction of SSR clones selected from genomic

libraries can be converted into informative SSR markers (e.g., Echt and Maymarquardt 1997; Pfeiffer *et al.* 1997; Mariette *et al.* 2001a). One strategy to increase the efficiency of the identification of microsatellite regions is to transfer SSR markers across genera. Nevertheless, the SSR information generally does not transfer across *Pinus* species (Echt and Maymarquardt 1997; Echt *et al.* 1999), e.g., 47 SSRs primer pairs developed in three *Pinus* species were tested in *P. pinaster*, but only one amplified at a single polymorphic locus (Mariette *et al.* 2001a). The SSR primer pair that cross-amplified was identified in *P. halepensis*, which is from the same subgenus *Pinus* and belongs to a strong support clade with *P. pinaster*, according to (Wang *et al.* 1999).

The availability of the entire chloroplast sequence of *P. thunbergii* (Wakasugi *et al.* 1994) allowed the identification of *chloroplast simple-sequence repeats* (cpSSR). Primers flanking a mononucleotide repeat located in the intergenic region between the *trnK* and *pbsA* genes were used to detect variation in different pine species (Powell *et al.* 1995b; Vendramin *et al.* 1996).

The evolution rate of cpDNA genes is estimated to be several times slower than of the nuclear genes (Wolfe *et al.* 1987), with a low average level of sequence variation (Clegg and Zurawski 1992). The genetic information contained in the chloroplast genomes of plants, including the arrangements of genes and intergenic sequences, is very conservative compared with the nuclear and mitochondrial genomes, which contain vast amounts of DNA of no apparent function (Birky 1988). Those properties confer to the cpSSR markers primer binding sites a high degree of conservation (Powell *et al.* 1995a), therefore the designed primers ought to work across taxa.

The high degree of conservation of sequences in the chloroplast genome of conifers and the universality of the primers was confirmed by several recent studies in conifers (Powell *et al.* 1995b; Vendramin *et al.* 1996; Vendramin and Ziegenhagen 1997; Sperisen *et al.* 1998). Moreover, the primers have been used with success in 110 different conifer species belonging to different taxa, in particular, *Pinaceae*, *Cupressaceae* and *Taxodiaceae* (G.G. Vendramin, unpublished results). The universality of cpSSRs allows the transfer of primers across taxa alleviating the cost involved in their identification for each species. The use of automated DNA sequencing apparatus and adequate software can increase the efficiency and allows obtaining a large set of data in a relatively short period of time.

The use of cpSSRs has allowed the investigation of the distribution of chloroplast haplotypes and haplotypic diversity in different conifers at the range level of the species (Lefort *et al.* 1999 and references therein), including *P. pinaster* (Vendramin *et al.* 1998), and that evidence appears to be associated with the migration processes from glacial refugia that occurred in the most recent postglacial period.

In contrast to nuclear genomes, plant organelle genomes are haploid and uniparentally inherited. For most angiosperms the chloroplast genome is

maternally transmitted, but in conifers it is generally paternally inherited (Neale *et al.* 1986; Szmidt *et al.* 1987; Neale and Sederoff 1989; Wagner *et al.* 1989; Stine and Keathley 1990; Dong *et al.* 1992; Ziegenhagen *et al.* 1995; Cato and Richardson 1996; Vendramin and Ziegenhagen 1997; Stoehr *et al.* 1998) including in *P. pinaster* (Plomion *et al.* 2001). Due to the uniparental mode of inheritance, the chloroplast genome behaves as a haploid single-locus and does not undergo recombination (Chiu and Sears 1985). Except for occasional mutations, this molecule is inherited unaltered with linked associated loci, therefore the sequences are a source of evolutionary history information.

Due to the uniparental inheritance and to the smaller effective population size, organelle genomes are more sensitive than nuclear DNA to severe reductions in the number of individuals in a population; they are expected to show different population dynamics and may be more sensitive to population subdivision (Birky *et al.* 1989). Studies in conifers have shown that cytoplasmic markers generally display higher values of differentiation compared with nuclear markers (e.g., Hong *et al.* 1993; Petit *et al.* 1993; Strauss *et al.* 1993; Ennos 1994; Wang and Szmidt 1994; Hong *et al.* 1995).

Patterns of population subdivision in conifers may be influenced by the contrasting mode of cpDNA and mtDNA inheritance (Hu and Ennos 1999). Gene flow of organelle genes distributed only through seed (e.g., maternal inheritance for mtDNA in pines) can be significantly less among wind-pollinated tree species compared to organelle genes distributed by pollen and by seed (e.g., paternal inheritance for cpDNA in pines). As a consequence, in wind-pollinated outcrossers such as pines, population subdivision can be weaker when cpDNA markers are used compared with mtDNA, because the wind-dispersed pollen is the main agent of gene flow (Dong and Wagner 1994; Ennos 1994; Latta and Mitton 1997).

Mutation rates are generally ignored because they are considered to be much lower than the migration rates, but this might not always be valid. Recent data show that mutation rates are higher at the cpSSR loci than substitution rates elsewhere in the chloroplast genome, and generally higher than in the nuclear genome sequences, except for the nuclear SSRs (Provan *et al.* 1999) and references therein), and in this case higher mutation rates could reduce population subdivision. Another possibility is size homoplasy, which has been observed at chloroplast microsatellites (Doyle *et al.* 1998). This could also lead to underestimates of differentiation when cpSSRs are used, by erasing some of the differences in haplotypes that have arisen in the past. Comparisons to other markers are needed to test whether or not those novel markers detect reliable estimates of genetic diversity and differentiation.

Summary of the results

Genetic variation of *Pinus pinaster*

Studies presented in Papers I and II provide information on the level and distribution of genetic variation among and within populations of *P. pinaster* at chloroplast and nuclear loci.

Variation within populations

In Paper I, some Portuguese populations of *P. pinaster* showed high within-population haplotypic diversity. The scattered distribution of *P. pinaster* in Portugal in ancient times, with different populations evolving separately and later mixed by human activity, could have been a source of haplotypic diversity. An additional hypothesis is the possible migration pathway originating from putative refugia in southeastern Spain, as defended by Salvador *et al.* (2000). When allozyme markers were used, Salvador *et al.* (2000) found higher values of within-population diversity of *P. pinaster* in the eastern region of the Iberian Peninsula than in the Atlantic region (which includes Portugal). Vendramin *et al.* (1998) also detected relatively low values of diversity in two of the populations from Portugal in their range-wide study of cpSSR variation in *P. pinaster*. The earlier hypothesis of Baradat and Marpeau-Bezard (1988), that Portugal was a refuge for the species during the last glaciation, was rejected by Vendramin *et al.* (1998) due to the low values for within-population diversity found in some of the Portuguese populations of *P. pinaster*. Those populations included in Vendramin *et al.* (1998) study were probably of very recent origin and experienced a severe reduction in population size, with consequent founder effect. Moreover, further comparisons (Paper I) based on a subset of common loci proved one population of Portuguese origin (Leiria) to be the most polymorphic of those used by Vendramin *et al.* (1998) and in Paper I.

Fossil, charcoal and palynological records indicate that *P. pinaster* was present in Portugal during the Middle Würm glaciation (55 000-25 000 BP), the late Plenigacial (25 000-15 000 BP) and terminal Pleistocene (12 000-11 000 BP) (Mateus and Queiroz 1993; Figueiral 1995 and references therein). In addition, pollen analyses have also indicated the presence of *P. pinaster* forests in the coastal area south of Lisbon, during the Atlantic period (7 580-6 550 BP). This species was probably able to survive the latest glaciation in Portugal in sheltered areas close to the Atlantic Ocean (Figueiral 1995), in contrast to what is suggested by Salvador *et al.* (2000). In their study, the latter also suggested the disappearance of *P. pinaster* in Portugal during the last glaciation. According to Bennett *et al.* (1991), the influence of the Atlantic Ocean causes western Europe to be anomalously warm for its latitude, and various evidence of southern European climate and topography from 18 000 BP suggest that sites within this

region should have been suitable for tree survival throughout the last glaciation, perhaps even at densities too low to disperse a detectable pollen rain.

In Paper **II**, the diversity of the French populations computed with cpSSRs and AFLPs (pruning those loci that showed less than 4 null homozygotes, as recommended by Lynch and Milligan 1994) was higher than the diversity of the Portuguese populations. This supports the trend found by Vendramin *et al.* (1998), i.e., a higher diversity in the French compared with the Portuguese populations, and the authors concluded that French populations represent one reservoir of haplotypic diversity in the range of the species. This finding does not exclude the possibility of a putative refugium in Portugal, and conclusions about the migration history of the species should be drawn in the context of historical, fossil and palynological information, due to the known human impacts experienced by this species in its range, as discussed in the foregoing and in Paper **I**. In particular, due to the intensive reforestation of this species in the southwest of France (Scott 1962), the high diversity found in the *P. pinaster* populations of that region could be a result of the mix of plant material coming from different places, representing a “melting pot” rather than a natural centre of diversity. It should be recognized that anthropogenic influences could have erased the fingerprints of migration pathways for *P. pinaster*, particularly in those regions where these effects were more intense, which was the case for Portugal and southwestern France.

Variation among populations

According to El-Kassaby (1991) and references therein, the majority of genetic variation for pine species is found within populations, with a small but significant component among populations. In contrast, in *P. pinaster* a high intraspecific variation and genetic differentiation has been found among populations across the range using different types of markers (phenotypic traits, terpenes, denatured proteins, allozymes and chloroplast microsatellites) (Baradat and Marpeau-Bezard 1988; Bahrman *et al.* 1994; Alía *et al.* 1995; Petit *et al.* 1995; Vendramin *et al.* 1998). The geographical isolation and the possibility of scattered refugia during the last glaciation have been offered as possible explanations for this pattern of differentiation (Baradat and Marpeau-Bezard 1988; Vendramin *et al.* 1998).

In studies **I** and **II**, both the Portuguese and French populations of *P. pinaster* showed a low level of among-population diversity. Nevertheless, those studies were made at a fine geographical scale, and both natural gene flow and human activities could have erased the differences among populations at a regional level. Castro (1989) using allozymes and Mariette *et al.* (2001b) using AFLP and nuclear microsatellites, both obtained a similar pattern of differentiation in groups of Portuguese and French populations. Results obtained with allozymes by Salvador *et al.* (2000) and González-Martínez *et al.* (2001) showed clinal trends of genetic variation and fine-scale spatial structure in the Iberian Peninsula, but orientation of the main mountain ranges and the scattered distribution of *P.*

pinaster in Spain constituted natural barriers preventing the genetic homogenizing effect of gene flow. Interestingly, one of the few stands that showed to be divergent from the others in Paper I is situated in a very isolated part of Portugal and is surrounded by a high range of mountains.

In Portugal, it was common in *P. pinaster* planting programmes since at least the beginning of the 20th century to use seeds from different parts of the country and even from abroad. The fact that no discernible geographical genetic pattern was found in study I reflects this practice and strong anthropogenic influences are confirmed by historical records (Devy-Vareta 1993). Good agreement was found between the results presented in Paper I and those obtained by Salvador *et al.* (2000), González-Martínez *et al.* (2001) and Burban *et al.* (1999). In the former two studies, the authors observed a lack of relationship between geographic and genetic distances among the northwestern populations of the Iberian Peninsula, in a set of populations from the Iberian Peninsula, using isozyme markers. In the latter study, the authors found a blurred geographic pattern for a specific pest of maritime pine, the bast scale *Matsucoccus feytaudi*, in Portugal.

In southwestern France, the human impact through reforestation with *P. pinaster* in that region, which started as early as the 18th century and continued until a significant area was covered (Scott 1962), should also be taken in consideration. The low differentiation exhibited by cpSSRs and AFLPs of the French populations detected in Paper II is strong evidence of the probable influence of this factor, together with gene flow, in obscuring the divergence among populations within this region.

In study II, the differentiation obtained with all populations considered together (Portuguese and French) was higher than within each provenance separately, and both provenances could clearly be distinguished with both types of markers (AFLP and cpSSR). Several earlier studies at the range level of the species and using different type of markers (terpenes and denatured proteins) showed that populations from southwestern France and Portugal clustered together (Baradat and Marpeau-Bezard 1988; Bahrman *et al.* 1994). In a more recent study the two provenances were discriminated with the results obtained using cpSSR (Vendramin *et al.* 1998).

Nuclear versus cytoplasmic genetic variation

In Paper II, the AFLPs revealed much lower within-population diversity than the cpSSRs. This could be explained by the nature of microsatellites that are usually highly polymorphic markers (Lefort *et al.* 1999 and references therein), and the cpSSR used in the study II were already known to show polymorphism in *P. pinaster* (Vendramin *et al.* 1998 and Paper I). Nevertheless, the same trend was found with both markers, i.e., a lower diversity in the Portuguese compared with the French provenance, after the AFLP loci with less than 4 null homozygotes were pruned from the analysis, as suggested by Lynch and Milligan (1994) (LM restriction).

The AFLP markers used in study **II** exhibited similar levels of among-population diversity compared with the cpSSR markers, within both the French and Portuguese provenances, when the LM restriction was used. In the absence of extensive gene flow, uniparentally inherited markers show, in general, less variation within populations and more among populations, than nuclear biparentally inherited markers, due to a lower effective number of genes and to differences in seed and pollen migration (Birky *et al.* 1983; Petit *et al.* 1993; Wade *et al.* 1994).

In pines, population subdivision can be weaker when cpDNA markers are used compared with mtDNA, because the wind-dispersed pollen is the main agent of gene flow (Dong and Wagner 1994; Mitton *et al.* 2000). Allozymes and cpDNA revealed little population structure in *P. flexilis*, whereas mtDNA showed high population differentiation (Latta and Mitton 1997). Additionally, use of the Mantel test in Paper **II** showed that the genetic distance matrix calculated with AFLP loci was highly correlated with that calculated with cpSSRs. Despite the fact that both types of markers have different modes of inheritance, it appears that the effect of gene flow through pollen surpasses the effect of genetic drift in shaping the genetic variation of the species, at least at the geographical scale studied in Paper **II**.

In Paper **I**, using cpSSR the results indicated that there is little or no geographic genetic pattern in Portuguese populations, due not only to the effect of human activity, but also to extensive gene flow among populations, as described in the previous paragraph. Extensive gene flow could also explain similar differentiation values found for both nuclear and cytoplasmic markers within the Portuguese provenance, by smoothing differences due to differences in effective population sizes and genetic drift. Moreover, in study **II** the genetic differentiation among the Portuguese populations with AFLP (after LM restriction) and cpSSR data that are similar to that reported by Castro (1989) using allozyme markers ($G_{ST} = 0.020$) in six populations of *P. pinaster* widely spaced across Portugal.

The differentiation estimates obtained with the cpSSR and LM-restricted AFLP data in Paper **II** for the French provenance are also similar to the estimate obtained with nuclear microsatellite markers in a study made by Mariette *et al.* (2001b), in the same group of populations. Again, the low differentiation showed with different type of markers is a strong evidence of the genetic homogenising effect among populations due to extensive gene flow.

Studies in conifers that report lower population subdivision with nuclear markers compared with cytoplasmic markers have not included chloroplast microsatellites (e.g., Birky 1988, Petit *et al.* 1993; Dong and Wagner 1994; Ennos 1994), with the exception of a study in *P. leucodermis* with cpSSR and allozymes reported by Powell *et al.* (1995b). Recent data showed that mutation rates are higher at the cpSSR loci than substitution rates elsewhere in the chloroplast and nuclear genome, except for the nuclear SSRs (Provan *et al.* 1999 and references therein),

and in this case higher mutation rates could decrease population subdivision. Another argument to be considered is the size homoplasy observable at cpSSR loci (Doyle *et al.* 1998 and references therein). Since cpSSRs are generated by mutations at a limited number of hotspots, they are prone to suffer identical mutations occurring independently in different populations, which, in turn, biases the estimates of differentiation downward by erasing the differences in haplotypes the populations would otherwise possess.

Reproductive material identification and certification

Following World War II, during which large areas of forest were burned, reforestation programs were undertaken in the southwest of France (the Aquitaine region) with *P. pinaster* seeds of northwestern Iberian origin (Portugal and Galicia), and the stands they formed suffered considerable frost damage (Boisseaux 1986). To overcome this problem, and to avoid further damage, a terpene-based test was developed to test the putative origin of adult stands in Aquitaine, before seeds could be distributed for commercial purposes in France (Baradat and Marpeau-Bezard 1988). In Paper III, we describe a new test employing cpSSRs, based on randomisation tests to facilitate identification of stand origin of stands and seed-lots of *P. pinaster* in the Aquitaine region in France, and thus providing an alternative to the former terpene-based test for reproductive material identification and certification.

The origin of five stands of unknown origin was determined with both the cpSSR and biochemical (terpene profile analysis) tests. The results obtained with terpenes proved to be less efficient than those obtained with the cpSSRs. The terpene test was initially inconclusive for two out of the five tested stands and it had to be repeated before a reliable answer was obtained. Thus, use of the terpene test risks the need for repetitions, increasing the amount of plant material, costs and time needed to get the same information as obtained from one cpSSR test.

The results obtained in Paper III can be partially explained by the fact the terpene markers discriminate less the Portuguese from the French provenances as indicated by the range-wide *P. pinaster* study that employed terpene analysis (Baradat and Marpeau-Bezard 1988). Chloroplast microsatellites revealed high levels of genetic differentiation among populations across the range due to differences in allele size (Vendramin *et al.* 1998), compared with the other markers (Bahrman *et al.* 1994; Petit *et al.* 1995). The cpSSRs data were used to design the test, due to the fact that they showed a homogeneous distribution of the polymorphism within groups and clear differentiation between the two groups of populations (French and Portuguese). Therefore, the cpSSR markers used in the Paper III were suitable for the principal purpose of that study, the design of a test to determine the putative origin of *P. pinaster* stands in the Aquitaine region of France.

Closing remarks

Studies **I** and **II** presented in this thesis illustrate the patterns of population genetic structure of the investigated populations of *P. pinaster*. Paper **I** presents strong evidence of Man's influence in shaping the genetic structure of a species and how the mixing of genetic material of unknown origin can affect the genetic resources. Little or no geographic genetic pattern was found in Portuguese populations of *P. pinaster* due to human activities during the last century, and to the extensive associated gene flow among populations. As a consequence of the blurred pattern found in the Portuguese populations of *P. pinaster*, the interpretation of the history of this species in this area may prove difficult if not impossible. In Paper **II** a similar population structure was observed in the southwestern region of France. The low levels of differentiation found among populations of that region was probably due to intense gene flow, but the reported human influence in the region's forest building is not to be excluded.

In future research, it is recommended to perform a range-wide study with a denser sampling of *P. pinaster* populations, to confirm the hypothesis drawn by Vendramin *et al.* (1998) to explain the history and pathways of the species since the last glaciation. The use of other markers is suggested, such as mtDNA markers, due to the lower scale dispersal of the seeds compared with the pollen in pine species. Gene flow through pollen could have blurred the information retrieved by paternally inherited chloroplast markers. Unfortunately, the polymorphism found so far with mtDNA markers (C. Burban, unpublished results) is very low, and further research needs to be pursued. Additionally, conclusions about the migration history of the species should also be presented in the context of historical, fossil and palynological information, due to the known anthropogenic influence in the forests of the Mediterranean Basin.

Similar levels of differentiation were found with AFLPs and cpSSRs markers, when the LM restriction was applied, in the two regions studied in Paper **II**. Moreover, the differentiation they exhibited was similar to that computed with codominant markers (nuclear microsatellites and allozymes) for the same regions. Despite the fact that the two kinds of markers have different modes of inheritance, the trends found with the genetic distances computed with both types of markers were similar. This observation supports the hypothesis that the effect of gene flow through pollen surpassed the effect of genetic drift in shaping the genetic variation of the species at the geographical scale studied.

In the genus *Pinus*, pollen flow among adjacent populations is generally high and lowers population subdivision, which may have smoothed the differences found here between estimates of differentiation by nuclear and chloroplast markers. However, it is likely that species that show low levels of differentiation on a

regional scale may reveal very high differentiation on a macro-scale. *Pinus pinaster* seems to be a good example of this kind of behaviour.

The test for provenance identification designed in Paper III can easily be applied to other commercial species, provided that there is a homogeneous distribution of the polymorphism within groups and clear differentiation among groups of populations. In particular, the cpSSR primers have already been shown to cross-amplify sequences from several species, which could be very advantageous given the long time and high costs involved in identifying markers (Powell *et al.* 1995b, Vendramin *et al.* 1996). The availability of reliable tests for identifying the origins of reproductive material will be very valuable for providing solutions to seed certification problems and also in the context of gene conservation.

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¹ It is worthwhile to read J.A. Estevão (1983) A florestação fos baldios, *Análise Social*: 1157-1260.

² According to J.Serrão (1977, *A Emigração Portuguesa*, Lisboa) until the beginning of the 50's the emigration in Portugal was negligible.

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