# 1 Does the seed fall far from the tree?-- <u>wW</u>eak fine-<u>-</u>scale

# 2 genetic structure in a continuous Scots pine population

3 Alina K. Niskanen<sup>1,3\*</sup>, Sonja T. Kujala<sup>2</sup>, Katri Kärkkäinen<sup>2</sup>, Outi Savolainen<sup>1</sup>, Tanja

4 Pyhäjärvi<sup>3</sup>

5

- 6 <sup>1</sup>Ecology and Genetics Research Unit, University of Oulu, 90014 University of Oulu,
- 7 Finland
- 8 <sup>2</sup>Natural Resources Institute Finland (Luke), <u>Paavo Havaksen tie 3, 90570</u>Oulu, Finland
- 9 <sup>3</sup>Department of Forest Sciences, University of Helsinki, 00014 University of Helsinki,
- 10 Finland
- 11
- 12 \*Corresponding authors:
- 13 Alina Niskanen, alina.niskanen@gmail.com
- 14 Tanja Pyhäjärvi, tanja.pyhajarvi@helsinki.fi

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#### 16 Abstract

Knowledge of fine-\_scale <u>spatial</u> genetic structure, *i.e.*, the distribution of genetic diversity at short distances, is important in evolutionary research and <u>in</u> practical applications such as conservation and breeding programs. In trees, related individuals often grow close to each other due to limited seed and/or pollen dispersal. The extent of seed dispersal also limits the speed at which a tree species can spread to new areas.

22 We studied the fine--scale spatial genetic structure of Scots pine (*Pinus sylvestris*) in two 23 naturally regenerated sites located 20 km from each other located in continuous south-24 eastern Finnish forest. We genotyped almost 500 adult trees for 150k SNPs using a custom made Affymetrix array. While wWe detected some pairwise relatedness at short 25 26 distances, but the average relatedness was low and decreased with increasing distance, 27 as expected. Despite the clustering of related individuals, the sampling sites were not differentiated ( $F_{ST}$  = 0.0005). According to our results, Scots pine has a large neighborhood 28 size (*Nb* = 1680–3 $\pm$ 2 $\pm$ 210), but a relatively short gene dispersal distance ( $\sigma_g$  = 36.5–71.3 m). 29 30 Knowledge of Scots pine fine-scale spatial genetic structure can be used to define suitable 31 sampling distances for evolutionary studies and practical applications. Detailed empirical 32 estimates of dispersal are necessary both in studying post-glacial recolonization and 33 predicting the response of forest trees to climate change.

# 34 Introduction

35 Understanding the fine--scale spatial genetic structure of species is important in a wide 36 range of fields. It builds up, through gene flow, from an interplay between: i) external The 37 structure builds up from the interaction of gene flow and extrinsic factors, such as fragmented habitats and differences in growth conditions, which in turn affect selection, 38 39 population size and density, and ii) internal On the other hand, intrinsic factors, such as 40 dispersal ability and mating patterns of the species also have an influence (Loveless & 41 Hamrick 1984; Vekemans & Hardy 2004). Gene flow takes place between individuals in 42 physical space (Bradburd & Ralph 2019), often leading to an isolation-by-distance pattern 43 (Wright 1943; Málecot 1967). Thus, individuals close to each other are commonly expected to be more closely related than a random sample from within the species. In 44 sedentary species, likesuch as trees, spatial aggregation of related individuals often 45 46 results from limited seed and/or pollen dispersal (Hardy & Vekemans 1999). In wind-47 pollinated species, pollen can disperse long distances, from hundreds of meters to 48 hundreds of kilometers (e.g., Kremer et al. 2012; Desilva & Dodd 2021), whereas seed 49 dispersal has shorter average distances (Kremer *et al.* 2012). Pollen dispersal distances in 50 animal-pollinated species are commonly shorter-from a few meters to a few kilometers (e.g., Levin & Kerster 1974; Kremer et al. 2012; but see Ahmed et al. 2009). As a result of 51 52 the differences in dispersal distances, strong fine\_scale <u>spatial genetic</u> structure is common in animal-pollinated and rare in wind-pollinated tree species (Vekemans & Hardy 53 54 2004; Hardy et al. 2006; Born et al. 2008; Vakkari et al. 2020). Thus, tThe distribution of 55 genotypes at-a short distances is usually transient, largely impacted by life-history traits and can be evaluated through estimating relatedness between individuals. 56

In evolutionary research, spatial genetic information is valuable for inferring the strength of
 the evolutionary forces—genetic drift, selection, and gene flow—that participate in forming
 the genetic structure (Rousset 2003; Slatkin 1985). The <u>fine-scale spatial genetic structure</u>

60 of populations needs to be considered in many applications too. For example, in genotype-61 phenotype association analyses, spurious associations may arise due to correlation of 62 allele and trait frequencies in different populations, but also within a population, if the 63 underlying genetic structure is not corrected for (Pritchard & Rosenberg 1999; Persyn et al. 2018). Also in practical applications, wheren individuals are chosen from natural 64 65 populations for conservation, population management or breeding programs from natural 66 populations, it is essential to know how genetic diversity and, for example, rare alleles are distributed in space to maintain high genetic diversity, avoid inbreeding and, on the other 67 68 hand, <u>yunnecessarunintended</u> mixing of differentially adapted populations (Desilva & 69 Dodd 2021; Escudero et al. 2003; Smith et al. 2018). When the span of spatial 70 autocorrelation is known, sampling can be adapted to the needs of each application. 71 Knowledge of fine--scale spatial genetic structure can recursively be used to infer dispersal 72 distances (Málecot 1967; Rousset 1997 & 2003). The ability to disperse becomes increasingly important in the light of climate change, as when plant species and their 73 74 locally adapted populations may become maladapted to their current locations (Gougherty 75 et al. 2021). Dispersal information can be used, for example, in predicting species' potential for adaptation (Kuparinen et al. 2010; Kremer et al. 2012; Barton 1979; Slatkin 76 77 1973), bearing in mind that dispersal rates in, e.g., the open landscapes of colonization stage may be different than those estimated here. If the natural dispersal rate is estimated 78 79 to be too slow tofor responding to the challenges of climate change, human assisted 80 migration is one possible way to aid adaptation (Marris 2009; Aitken & Whitlock 2013). Scots pine (*Pinus sylvestris*) is a keystone conifer species in large parts of the forests of 81 Northern Eurasia and, thereforeus, important for the ecosystem functioning (Pyhäjärvi et 82 83 al. 2020 and references therein). As a major source of timber, paper and pulp, Scots pine 84 holds also has high economic value, especially in Fennoscandia (e.g., in Finland, 85 https://www.luke.fi/en/statistics/wood-consumption/forest-industries-wood-consumption-2021). Due to its continuous and wide distribution, wind-pollination and predominantly 86

87 outcrossing mating system (Muona & Harju 1989), Scots pine shows very weak <u>genetic</u> 88 population structure at the global scale; largely in the form of subtle isolation-by-distance 89 across its distribution range (Tyrmi et al. 2020). Less is known about within-population\_ 90 genetic structure as only small, fragmented populations have been studied forin this 91 arespect so far (Robledo-Arnuncio & Gil 2005; Sofletea et al. 2020). Although artificial 92 regeneration with genetically improved seedlings ishas becomeing more common in 93 forestry, natural regeneration has been the predominant regeneration method in forests, 94 making the patterns described here common inacross Fennoscandia. Since Scots pine 95 has such a dominant role in the boreal forest ecosystems, even small changes in its distribution or adaptationadaptive ability may have large consequencesroll-down effects. 96 Here we investigate the fine-scale spatial genetic structure of Scots pine in two naturally 97 98 regenerated sites that are located in south-eastern Finland. We use a genome-wide 400k 99 single nucleotide polymorphism (SNP) array (Kastally <u>& Niskanen</u> et al. 2022), which allows us to estimate relatedness in a large sample of 469 trees. We estimate the 100 101 parameters of the isolation-by-distance model and use this information to derive estimates 102 of gene dispersal distance. We T then we investigate the spatial spread and sharing of rare 103 alleles that can have a distinct profile of, for example, different fitness effects and average allele age compared to more common alleles. The knowledge of fine\_scale spatial genetic 104 structure and dispersal distance areis useful in practical applications of Scots pine 105 106 breeding and in associating genetic and trait variation in natural populations, but also for 107 modeling adaptation and making predictions on how widely- distributed wind-pollinated trees can respond to climate change. 108

## **109** Material and Methods

#### 110 Samples and genotypes

- 111 Our study population at <u>the</u> Punkaharju intensive study site (ISS,
- 112 https://www.evoltree.eu/resources/intensive-study-sites/sites/site/punkaharju) is a naturally
- 113 regenerated site in south-eastern Finland-and includes two sampling sites, Mäkrä
- 114 (61°50'<u>16.8"</u>N, 29°23'<u>39.5"</u>E) and Ranta-Halola (61°39'<u>19.7"</u>N, 29°17'<u>14.7"</u>E) located 21
- 115 km apart (Figure 1). <u>The landscape in south-eastern Finland has mostly continuous pine</u>
- 116 forest cover, lakes and some agricultural areas. The study site stands arose through
- 117 <u>natural regeneration after seed tree cuttings (i.e., retaining only part of the mature trees to</u>
- 118 provide seed for establishing the next generation) 60–70 years ago in Mäkrä and an
- 119 <u>unknown, but probably a few decades longer, time ago in Ranta-Halola.</u> We sampled 469
- adult (33–145 years) Scots pines <u>at approximately at 20 m distances</u>, with <u>the shortest</u>
- 121 within-\_sampling site distance between trees\_ofbeing 10 and 14 m and the longest 464 and
- 122 1164 m in Mäkrä and Ranta-Halola, respectively. <u>We selected trees that were similar in</u>
- 123 size by eye—113 trees from Mäkrä and 356 trees from Ranta-Halola. The mean age,
- 124 <u>estimated by counting the tree rings from a core sample at breast height, was 60.6 (range:</u>
- 125 33–112.5) and 90.3 (range: 43–144.5) years in Mäkrä and Ranta-Halola, respectively
- 126 (Figure S1). We could not estimate the age for two trees from Ranta-Halola.

Needle samples from the adult trees were genotyped on a custom-made Affymetrix SNP
array including 407 540 markersSNPs. Development of the SNP array and genotyping of
the samples is described in detail in Kastally & Niskanen *et al.* (2022). In short, of the 407
540 SNPs we used a dataset of 157 325 polymorphic SNPs with the ThermoFisher
conversion types Poly High Resolution (three well-separated genotype clusters) and No
Minor Homozygote (two well-separated genotype clusters, homozygous and
heterozygous) as a starting point for filtering the loci. We used four embryo samples and

134 their parents (described in Kastally & Niskanen *et al.* 2022) to estimate Mendelian errors



- 138 <u>seemingly heterozygous genotype in haploid megagametophyte samples (described in</u>
- 139 Kastally & Niskanen *et al.* 2022) to avoid SNPs in potentially paralogous genomic
- 140 regions. We filtered for Mendelian errors and errors likely arising from paralogy common in-
- 141 conifers (Neale *et al.* 2014).-Further filtering of the genotype data was done according to
- 142 the requirements of each analysis as described below.

143



Figure 1. Maps of the sampling sites in the Punkaharju intensive study site located in
south-eastern Finland. Sampled trees are indicated as red dots. Mäkrä and Ranta-Halola
are located 21 km apart. The maps were drawn in R using the package ggmap (Kahle &
Wickham 2013).

149 Spatial data and pairwise distances

150 We recorded the coordinates for each tree using a portable GPS locator in August-October 2020. The initial coordinates in the ETRS-TM35FIN geodetic coordinate system 151 were transformed to the coordinates in the EUREF-FIN-GRS80 geodetic coordinate 152 system using a Finnish map service on the web (https://kartta.paikkatietoikkuna.fi/). One of 153 the study trees from Ranta-Halola had died between the needle sample collection and 154 coordinate recording and was excluded from the spatial analyses. We estimated the 155 pairwise spatial distance matrix for 468 adult trees with coordinates using the R (v. 3.6.3, R 156 core team 2020) package fields (Nychka et al. 2017) function "rdist.earth" (Nychka et al. 157 158 2017).

159 **Population structure** 

To get an overall picture of the genetic structure of our study population, we conducted 160 161 principal component analysis (PCA) using the R package pcadapt (Privé et al. 2020). We ran PCA for two sets of individuals, first for all 469 individuals, and second for 332 162 individuals excluding: i) individuals used in the SNP discovery (Kastally & Niskanen et al. 163 164 2022), ii) individuals related to the SNP discovery individuals (pairwise relatedness, <u>genomic relationship matrix (GRM)</u>  $\geq$  0.044; Yang *et al.* 2011; see details below), and iii) 165 one individual from each pair with pairwise relatedness  $\geq$  0.044. We excluded closely 166 167 related individuals and SNP discovery individuals from the PCA to detect the underlying population structure without the signal of family structure or SNP ascertainment effects. 168 169 We used a set of 65 498 SNPs with the following characteristics: minor allele frequency  $(MAF) \ge 0.05$ , close to Hardy Weinberg equilibrium (HW; exact test p value  $\ge 0.001$ ), and 170 relatively lownot in high linkage disequilibrium (LD) r<sup>2</sup> < 0.9 with other SNPs in 10 kb 171

- 172 windows within a contig (using PLINK v. 1.9; Purcell et al. 2007), unless stated otherwise.-
- 173 We used four embryo samples and their parents (described in Kastally et al. 2022) to-
- 174 estimate Mendelian errors for each locus in PLINK and excluded SNPs with more than one
- 175 Mendelian error. Since Scots pine has a high proportion of repetitive and paralogous-
- 176 genome sequence (Wegrzyn et al. 2014), we excluded SNPs with more than one-
- 177 seemingly heterozygous genotype in haploid megagametophyte samples (described in-
- 178 Kastally et al. 2022) to avoid SNPs in potentially paralogous genomic regions.
- 179 We estimated pairwise  $F_{ST}$  (Weir & Cockerham 1984) between the two study sites using
- 180 the R package StAMPP (Pembleton *et al.* 2013) and performed 1000 bootstraps to
- 181 estimate its 95% confidence interval (CI). We also estimated  $\phi_{ST}$  (Excoffier *et al.* 1992),
- 182 which measures population differentiation at molecular level, using Nei's *D* pairwise
- 183 genetic distances (Nei 1972) between individuals and 10 000 permutations to gain a *p*-
- 184 value in StAMPP.
- 185 Fine-scale spatial genetic structure

We estimated pairwise relatedness between all 469 samples as a genomic relationship
 matrix (GRM; <u>Yang *et al.* 2011</u>) using <u>the PLINKGCTA (Yang *et al.* 2011)</u> command "–
 make-grm\_-make-grm-gz". GRM was estimated between individuals *j* and *k* over SNP loci
 from *i* to *N* using the formula

,

190 
$$A_{jk} = \frac{1}{N} \sum_{i=1}^{N} \frac{(x_{ij} - 2p_i)(x_{ik} - 2p_i)}{2p_i(1 - p_i)}$$

191 where  $x_{ij}$  and  $x_{ik}$  are the numbers of reference (major) alleles in an-individuals j and k, and 192  $p_i$  is the reference allele frequency. Since inclusion of closely related individuals may 193 inflate the relatedness estimates, we used allele frequencies estimated for 387 unrelated 194 adult individuals (GRM < 0.0625, the mean of relationship class for, e.g., first cousins once 195 removed) as reference allele frequencies in the estimation of GRM. We classified each 196pair of individuals into family relationship classes (e.g., see Manichaikul *et al.* 2010 for.197kinship relationship estimates (F), which are equal to half of the corresponding relatedness198estimates) based on pairwise GRM: second degree between [0.177,0.354), e.g., half-199siblings, third degree between [0.088, 0.177), e.g., first cousins, fourth degree between200[0.044, 0.088), e.g., first cousins once removed, and unrelated below 0.044. We also201estimated the genomic inbreeding coefficient ( $F_{GRM}$ ) for each individual based on genomic202relationship matrix in GCTA (Yang *et al.* 2011).

The Mantel test (Mantel 1967) is a traditional test of spatial autocorrelation where the 203 204 relationship of two dissimilarity (i.e., distance) matrices is investigated. In spatial genetics, 205 the null hypothesis of a Mantel test is that genetic distance, or similarity when measured 206 as relatedness or kinship, and spatial distance are not correlated. To study the relationship 207 between relatedness and spatial distance, we estimated their correlation using 208 Spearman's correlation coefficient ( $\rho$ ) separately for the two sampling sites Mäkrä and Ranta-Halola. We conducted the Mantel test separately for both sites using the R package 209 210 ecodist (Goslee & Urban 2007) with 10 000 permutations. The use of the Mantel test in 211 spatial genetics has been criticized because of the lackreguirements of homoscedasticity and linear correlation between genetic and spatial distances for the data (Legendre et al. 212 2015). These problems are, however, less severe in the Mantel correlogram analysis 213 where samples are divided into pre-\_\_defined spatial distance classes and each distance 214 215 class is compared separately to joint data from other distance classes. We therefore also 216 used also Mantel correlograms (10 000 permutations) to evaluate the correlation between relatedness and distance within distance classes. 217

218 Neighborhood size and dispersal distance

219 We estimated neighborhood size (*Nb*), the <u>effective</u> number of <u>-potentially mating</u>

220 individuals belonging to a <u>within-population neighborhood</u>-(Wright 1946)panmictic-

221 breeding unit, and gene dispersal distance ( $\sigma_g$ ) using an iterative approach implemented in

222 SPAGeDi (Hardy & Vekemans 2002). To <u>attainobtain</u>  $\sigma_g$ , SPAGeDi first estimates <u>a</u> 223 <u>starting value for</u> neighborhood size using the formula

224 
$$Nb = -(1 - F_N)/blog$$
,

where  $F_N$  is the mean kinship (Loiselle *et al.* 1995) of the first distance class (0–50 m in Mäkrä and 0–60 m in Ranta-Halola) and *blog* is the regression slope of the regression of kinship on the natural logarithm of spatial distance over all distance classes (Rousset 2000; Hardy & Vekemans 2002). Kinship was estimated with 28 378 SNPs with MAF  $\geq$ 0.20 (due to computational limitations), using the formula

231 
$$F_{ij} = \frac{\sum_{l} \left[ \sum_{a} (p_{ila} - p_{la}) (p_{jla} - p_{la}) + \sum_{a} \frac{p_{la}(1 - p_{la})}{(n_{l} - 1)} \right]}{\sum_{l} \sum_{a} (p_{la}(1 - p_{la}))}$$

where  $p_{ila}$  and  $p_{jla}$  are is the frequency of allele *a* at locus *l* in individuals  $f_i$  and  $j_i$ ,  $p_{la}$  is the reference allele frequency of allele *a* at locus *l*, and  $n_l$  is the number of gene copies defined in the sample at locus *l* (Loiselle *et al*. 1995; Hardy & Vekemans 2002). Then gene dispersal distance was estimated using the formula

237 
$$\sigma_g = [Nb/(4 \pi D_e)]^{1/2}$$
 ,

where *D<sub>e</sub>* is the effective population density that accounts for differences in the
reproductive success of individuals (Hardy & Vekemans 2002). The *Nb* estimation
procedure was repeated using *blog* from kinship~distance regression up to the distance of
the chosen maximum σ value. Census density (*D*) estimates of unmanaged Finnish Scots
pine forests vary between 608–4470 trees per hectare (Lönnroth 1926), and Based on the
census density (*D*) estimate in commercial forests is about-of-2 000 trees/ha-per hectare

244	(200 000 trees per km² <del>) in commercial forests (;</del> Fahlvik <i>et al</i> . 2005; Väisänen <i>et al</i> . 1989),
245	Our study sites have regenerated naturally after cutting to an unknown, but likely lower
246	than 2000 trees/ha density of seed trees. However, as pollen and seed dispersal from
247	surrounding forests (Jiménez-Ramírez et al. 2021) likely increases the effective density,
248	we used 2000 trees/ha as our starting point, and we estimated $\sigma_g$ assuming ratios of
249	effective to census density of 0.25 ( $D_e$ = 500 trees/ha) and 0.5 ( $D_e$ = 1 000 trees/ha)
250	Census density estimates of unmanaged Finnish Scots pine forests vary between 608-
251	4470 trees/ha (Lönnroth 1926); moist forests have lower densities than dry forests at the
252	same age due to faster growth. Gene dispersal distance can reliably be estimated within a
253	distance that is assumed to be in mutation-drift-equilibrium that should be reached in a few
254	generations within the distance $\sigma_g/(2\mu)^{1/2}$ ( $\mu$ = mutation rate; Rousset 1997 & 2000). Based
255	on a 10 <sup>-3</sup> mutation rate of microsatellites, this distance is approximately 20* $\sigma_g$ . Since the
256	SNP mutation rate (10 <sup>-9</sup> ; Willyard <i>et al</i> . 2007) is lower than the microsatellite mutation rate,
257	we estimated Nb (and thus also sigma) using both 20* $\sigma_g$ and a higher value of 60* $\sigma_g$ .
258	However, since the maximum distances within our study sites are shorter than the
259	estimated 20* $\sigma_g$ distance, increasing the maximum distance does not affect our estimates
260	of sigma nor <i>Nb</i> . To measure the strength of the fine <u>-scale spatial</u> genetic structure, we
261	estimated the intensity of spatial genetic structure (Sp; Vekemans & Hardy 2004) using the
262	formula

- 263  $| Sp = -blog/(1 F_N) |$ .
- 264 We used 1 000 permutations of individual locations in estimating *blog* to test how probable
  265 it is to get higher *Sp* by chance.
- 266 Rare alleles

To study the spatial spread of rare alleles, we investigated sharing of rare alleles with MAF <br/> < 0.01 (23 623 SNPs after removing singletons) between individuals and correlated <br/>
(Spearman's  $\rho$ ) this with pairwise relatedness and spatial distance. To avoid the SNP

- discovery ascertainment bias in rare allele sharing, we estimated the correlation using only
  426 individuals excluding: i) individuals used in the SNP discovery and ii) individuals with
  pairwise relatedness (GRM) >= 0.044 with the SNP discovery individuals. We used within
  sampling site Mantel correlograms to study the correlation between rare allele sharing and
  spatial distance within distance classes. *P*-values for the correlations were
  constructedobtained with Mantel's test using 10 000 permutations. For illustrative
  purposes, we fitted a local (LOESS) regression in R for the proportion of shared rare
- alleles on relatedness and included sampling site as a fixed predictor.

#### 278 **Results**

#### 279 Spatial genetic analyses

On the population scale with all 469 adult trees, we identified <u>a</u>very weak structure 280 between Mäkrä and Ranta-Halola based on PCA (Figure 2). The majority of the samples in 281 both Mäkrä and Ranta-Halola clustered together along principal components (PC) 1 and 2 282 (Figure 2a). However, we detected one outlying cluster of samples from Ranta-Halola on 283 PC1 and one on PC2, and a group of samples from Mäkrä dispersed along PC2. These 284 285 outlying samples consisted of related individuals, and the outliers disappeared when we conducted PCA only for 332 individuals that were unrelated (GRM < 0.044) and not used 286 287 in the SNP discovery (Figure 2b). Concordantly, we found a low F<sub>ST</sub> of 0.0005 (95% CI: 288 0.0004–0.0005) between Mäkrä and Ranta-halola and a low but significant between-study-289 site variance component ( $\phi_{ST}$ ) of 0.0019 (p < 0.001).



Figure 2. Principal component analysis (PCA) for Mäkrä and Ranta–Halola for a) all 469
 adult trees and b) only 332 trees; excluding individuals used in the SNP discovery
 (Kastally <u>& Niskanen et al.</u> 2022), individuals with <u>pairwise relatedness</u> higher than 0.044 pairwise relatedness with the SNP discovery individuals, and one individual of each pair
 with pairwise relatedness ≥>= 0.044.

295

296 We found that pairwise relatedness was low within <u>the</u> study sites (Figure 3; <u>Table S1</u>).

297 Even in the class with <u>the</u> shortest distance between individuals, the mean GRM was <u>only</u>

298 **0.0004** in Mäkrä and 0.00218 in Ranta-Halola. However, relatedness still\_decreased with

299 increasing distance in both sampling sites withat a similar rate (Figure 3: see Figure S2 for 300 distance restricted to Mäkrä's maximum distance). We used relatedness (GRM) or kinship 301 (F, Loiselle) to estimate pairwise relatedness depending on the analysis, and both 302 methods gave similar estimates of relatedness (Pearson's r = 0.842, p < 0.0001; Figure S3). The spatial genetic structure was evident individuals in closer distance classes were 303 304 more related to each other compared to individuals in other distance classes until ~65 m in 305 Mäkrä and ~200 m in Ranta-Halola. In these distance classes, there was a negative 306 correlation between relatedness and distance as shown by the Mantel correlograms (Figure 4: Figure S4). The Mantel test indicated a subtle but significant decay of 307 308 relatedness withby spatial distance at Ranta-Halola (Spearmann's  $\rho = -0.044$ , one-tailed p 309 < 0.0001) but not at Mäkrä (Spearmann's  $\rho$  = -0.010, one-tailed p = 0.216). We also estimated the intensity of spatial genetic structure and found that, despiteeven when there 310 311 wasis evidence for spatial genetic structure, its intensity wasis low in both study sites 312 (Ranta-Halola Sp = 0.0008, one-tailed p < 0.001, and Mäkrä Sp = 0.0005, one-tailed p < 0.0010.001). The Llow intensity wasis caused by the smallmild decrease in the pairwise kinship 313 with distance over each sampling site and by the low average kinship in the first distance 314 class. 315

Among the 69 163 pairwise comparisons, 24 closely related pairs (GRM  $\ge$  0.177) were 316 identified (Figure 5). All closely related individuals (GRM  $\ge$  0.044) were growing  $\Theta$  in the 317 same sampling site. The highest relatedness (GRM) for a between sampling site pair was 318 0.031, whereas the highest within site relatedness was 0.332 in Mäkrä and 0.349 in Ranta-319 Halola. When the trees were divided into family relationship classes, the median distance 320 of the most related family relationship class found here (GRM = 0.177–0.354, indicating 321 322 second-degree relatedness) was 51 m in Mäkrä and 59 m in Ranta-Halola, compared to the respective median distances of 166 m and 357 m offor unrelated individuals (GRM < 323 0.044; Figure 5). This illustrates that the spatial aggregation of closely related individuals is 324 325 comparablesimilar in both the smaller and the larger study area. Family relationship

- 326 classes categorized using relatedness vs. kinship estimates were very similar, with a few
- 327 pairs categorized in the neighboring classes (Table S2). We did not find any sign of close
- 328 inbreeding in our sample; the highest inbreeding coefficient ( $F_{GRM}$ ) was 0.035 in Mäkrä and
- 329 0.080 in Ranta-Halola.

330





- 336
- 337 Depending on the effective population density  $(\underline{D}_e)$  estimate we used, the mean
- neighborhood size (*Nb*) over the two Scots pine sites was 3210 (with  $D_e = 500$  trees/ha;
- Table 1) or 1680 trees (with  $D_e = 1000$  trees/ha). We estimated the mean gene dispersal
- 340 distances ( $\sigma_g$ ) togetherjointly with *Nb* and found that the mean  $\sigma_g$  were 71.3 m ( $D_e$  = 500
- 341 trees/ha; Table 1) and 36.5 m ( $D_e = 1000$  trees/ha).

342



Figure 4. Mantel correlogram showing the €correlation (measured as Mantel *r*) between
 pairwise relatedness and distance within each distance classes estimated as Mantel
 correlogram in a) Mäkrä (circles, turquoise) and b) Ranta-Halola (squares, orange). Filled
 circlesshape indicates a *p*-value smaller than≤ 0.05. Two of the longest distance classes
 from Mäkrä and one from Ranta-Halola have been left out due to including less than 100
 pairwise comparisons.

**Table 1.** Gene dispersal distance ( $\sigma_g$ ) and neighborhood size (*Nb*) estimates for Mäkrä and Ranta-Halola for two different <u>ratios of</u> effective ( $D_e$ ) and <u>to</u> census (D) populationdensit<u>yies</u>. In this study,  $D_e/D$  ratio of 0.5 equals  $D_e = 1_000$  trees/ha and 0.25 equals  $D_e =$ 500 trees/ha. -Examples of *Nb* and  $\sigma_g$  for animal<u></u> and wind-pollinated tree species from previous studies <u>below</u>.

	D	/ <i>D</i> = 0.5	De	/ <i>D</i> = 0.25		
Study site (Pinus sylvestris)	Nb	$\sigma_{_g}$ (m)	Nb	$\sigma_{_g}$ (m)	Pollinator	Reference
Mäkrä	1404	33	3673	76	wind	this study
Ranta-Halola	1957	39	2747	66	wind	this study
Species						
Dicorynia guianensis	116	222		203	insects	Hardy <i>et al</i> . 2006
Moronobea coccinea	20	134		195	birds	Hardy <i>et al</i> . 2006
Milicia excelsa			370	3755	wind	Bizoux et al. 2009
Pinus pinaster*	97	30			wind	De-Lucas <i>et al.</i> 2009
Araucaria angustifolia	381	140			wind	Sant'Anna <i>et al</i> . 2013
Thuja occidentalis*	65	55			wind	Pandey & Rajora 2012

\*Only core or continuous populations included.



355 Figure 5. The relation Association between family relationship classes and spatial 356 distance. The pPairwise distances of the second degree related individuals in a) Mäkrä and b) Ranta-Halola are shown as red lines. The distribution of all pairwise distances of 357 individuals in different family relationship classes are shown in c) for Mäkrä (turquoise) and 358 359 Ranta-Halola (orange). Boxplots show the median (central vertical line), the lower and 360 upper guantiles (boxes), and up to 1.5 interguartile range (whiskers) distances. The Feamily relationships are classified based on pairwise GRM: second degree between 361 362 0.177–0.354 (e.g., half-sibling; n = 7 in Mäkrä and n = 17 in Ranta-Halola), third degree between 0.088–0.177 (e.g., first cousin; n = 12 in Mäkrä and n = 38 in Ranta-Halola), 363 364 fourth degree between 0.044–0.088 (e.g., first cousin once removed; n = 33 in Mäkrä and 365 <u>n = 164 in Ranta-Halola</u>), and unrelated below 0.044 (n = 6276 in Mäkrä and n = 62 616 in 366 Ranta-Halola).

## 367 Rare alleles

368 In line with overall relatedness, we found that more related individuals also shared a higher proportion of rare alleles than unrelated individuals (n = 426,  $\rho$  = 0.049, one-tailed  $\rho$  < 369 0.0001 estimated using Mantel's test; Figure 6), but this relationship iswas only visible in 370 higher relatedness values. Rare allele sharing and distance had a weak negative 371 372 relationship within study sites ( $\rho = -0.010$ ,  $\rho = 0.057$  in Ranta–Halola and  $\rho = -0.028$ ,  $\rho =$ 373 0.084 in Mäkrä; Figure S15). Mantel correlograms showed that pairwise sharing of rare alleles decreased with increasing spatial distance-similarly to the decrease of 374 relatedness with spatial distance (Figure S<sup>26</sup>). 375



Figure 6. The rRelationship between proportion of shared rare alleles of all rare alleles
and relatedness (GRM). Pairwise comparisons between individuals (dots) and LOESS
curve fitted to the proportion of shared rare alleles on relatedness (GRM).

#### 380 **Discussion**

381 Large continuous <u>Finnish</u> population of Scots pine has weak fine-<u>-</u>scale
 382 genetic structure

383 Species with <u>a very subtle structure betweenat the among populations level</u> are often 384 considered referred as panmictic, which indicates i.e., mating completely randomly mating 385 patternsalso within populations. While Scots pine's genotypic frequencies follow HW 386 frequencies equilibrium at the adult stage similarly to a panmictic species (Muona & Harju 1989, Pyhäjärvi et al. 2020), itsthe mating patterns are not completely random in space 387 (Robledo-Arnuncio & Gil 2005; Torimaru et al. 2012). In concordance Concordant with 388 389 spatially restricted mating patterns, we showed that fine-scale spatial genetic structure-390 albeit very weak—is maintained in adult Scots pine stands (Figures 3 & 4). This was 391 evident in the spatial proximity of individuals with higher pairwise relatedness (Figure 5). Scots pine has partial selfing (5–10% of mature seeds), but only a part of the selfed 392 offspring survive to the mature seed stage (Koski 1971, Kärkkäinen & Savolainen 1993), 393 394 and even fewer survive to the adult stage (Koelewijn et al. 1999). The average mortality of 395 the selfed seeds is 75–85%, compared to 20–30% for the open-pollinated seeds (Kärkkäinen et al. 1996, Koelewijn et al. 1999). In our samples, we did not find high 396 397 inbreeding estimates or pairwise relatedness values close to 1, which are expected in selfed progeny. Thus, relatedness patterns of adult trees found here are not a result of 398 399 selfing. Fine scale genetic structure has previously been found in smaller fragmented 400 Scots pine populations (Robledo-Arnuncio & Gil 2005; Sofletea et al. 2020), but here we show this pattern for the first time within a large continuous population for the first time. 401 402 We found that the spatial genetic structure reached somewhat longer distance in Ranta-Halola than Mäkrä (Figure 4; Figure S4). This may partly be caused by differences in the 403 404 shape and size of the sampled areas. In addition, each distance class includes 405 considerably more pairwise comparisons in Ranta-Halola, which results in more statistical

406 power. The mean age of Ranta-Halola is higher (Figure S1), which allows multiple

407 generations of dispersal events after the seed tree cuttings were done and, therefore, a

408 longer extent of spatial genetic structure. On the other hand, Mäkrä had proportionally

409 more pairs in the second degree family relationship class, which could be caused by the

410 more recent or possibly more intense seed tree cutting. There are likely differences in the

411 <u>fecundity of individual trees (Torimaru et al. 2012), which can cause differences between</u>

412 the sites. Nevertheless, the patterns of relatedness (Figures 3 & 5) and spatial genetic

413 <u>structure were very similar between the study sites, which suggests that the dispersal</u>

414 patterns are similar between the sites.

415 Long-distance pollination events and the continuous distribution range, allowing constant gene flow, are likely major contributors to the extremely low intensity of spatial genetic 416 417 structure we detected in our study (Sp from 0.0005 to 0.0008). In addition, the high 418 population density increases the neighborhood sizenumber of potential breeders, and all of these factors so thatkeep the fine-scale spatial genetic structure stays weak even 419 with when the relatively short average gene dispersal distance is relatively short. In 420 421 contrast, when populations are far from each other and disjunctnon-continuous, the chance of pollination from nearby trees is higher, which leads to stronger spatial genetic 422 423 structure. This effect of population fragmentation is evident in previous Scots pine studies 424 where Sp has ranged from up to 0.0098 in Scotland (González-Díaz et al. 2017) to themean of as high as 0.02071 onin the Carpathian Mountains (Sofletea et al. 2020). Our 425 426 study sites showed low intensity of spatial genetic structure also when compared to other conifers (Sp = 0.001–0.0349; Desilva & Dodd 2021; Kitamura et al. 2018; Sant'Anna et al. 427 2013; De-Lucas et al. 2009; Vekemans & Hardy 2004) and other tree species (Sp = 428 0.002–0.075; Bizoux et al. 2009; Hardy et al. 2006; Vekemans & Hardy 2004). The Llow 429 intensity of population spatial genetic structure is in line with strong gene flow through 430 pollen. ndlt also indicates that Scots pine populations evenr example, fo at the northern 431 432 distribution edge receive ample gene flow, and do not suffer from decreased have high

433	genetic variation as long as they are connected to the main population. Varis et al. (2009)
434	showed that the Finnish Scots pine populations at different latitudes are receptive for
435	pollen before their own pollen shedding starts, but the pollen from more southern
436	populations is already available for fertilization. This potential for gene flow from southern
437	to northern populations (Varis <i>et al.</i> 2009) also aidsfacilitates adaptation to climate change.
438	However, the situation is different in populations that are more fragmented and isolated,
439	such as the Spanish and Scottish populations, where the pollen dispersal is more
440	restricted and, thus, the spatial genetic structure is stronger (Hampe & Petit 2005).
441	Relatively short dispersal distance of Scots pine and its implications for
442	selection
443	We estimated that the average gene dispersal distance in our Scots pine stands was
444	53.94 meters (average of 71.3 m and 36.5 m), which is relatively short compared to the
445	gene dispersal distance estimates of other wind-pollinated trees, typically between 30
446	and 3755 meters (Table 1; Pandey & Rajora 2012; Sant'Anna <i>et al.</i> 2013; Bizoux <i>et al.</i>
447	2009; De-Lucas <i>et al.</i> 2009), and of animal—_pollinated trees up to 1296 meters (Hardy <i>et</i>
448	al. 2006) <u>although, However, p</u> opulation density <u>(estimate)</u> plays a large role in estimating
449	the dispersal distances. While estimating population density in Scots pine stands is
450	straightforward, it is more complicated to assess the density of a forest in the past, i.e.,
451	during establishment of seedlings that resulted in the current stand. As an early
452	succession species, regeneration mainly occurs in pulses after disturbances (e.g., forest
453	fires, storms and loggings; Linder et al. 1997; Lundqvist et al. 2017). Studies on
454	regeneration of seed and shelterwood stands also suggest that seedling establishment of
455	Scots pine occurs in sparse stands (50–200 trees/ha; Beland et al. 2010; Rautio et al.
456	2023). In the estimation of gene dispersal distances, the effective density estimate should
457	also take into account the breeding contribution of adult trees surrounding the cut area. It
458	should also be noted that the effective density of the study sites has varied greatly over

459	time, and trees germinating 33 or 143 years ago have faced very different fertilization and
460	germination conditions. We used the effective densities of 500 and 1000 trees/ha. By
461	assuming lower effective density, the estimated dispersal distance would increase.
462	However, a very long average dispersal distance does not seem probable given the
463	relatively sharp decay of the mean relatedness with distance and the observed pairwise
464	distances of the related individuals (Figures 3, 4 & 5). Further, Eearlier Scots pine seed
465	(10–20 m; Debain <i>et al</i> . 2007) and pollen dispersal (47.6–53 m; Koski 1970 <u>, based on</u>
466	radioactively labelled pollen; Robledo-Arnuncio & Gil 2005) estimates are relatively close
467	to our dispersal distance estimate, bearing in mind that our estimate includes both seed
468	and pollen components. Most dispersal distance estimates—including ours—assume that
469	the distribution of dispersal distances (i.e., the dispersal kernel) is Gaussian. However,
470	especially pollen is able to disperse very long distances (Lindgren et al. 1995; Robledo-
471	Arnuncio 2011) leading to potentially more leptokurtic dispersal kernels that are
472	challengingdifficult to estimate. Theseand would result in higher mean dispersal distances
473	(Robledo-Arnuncio & Gil 2005; Debain <i>et al</i> . 2007). <u>Taken together, our gene dispersal</u>
474	estimates can be taken as minimum estimates given the potential for lower effective
475	density and leptokurtic pollen dispersal.
476	UnderstandingKnowing the dispersal distance of thea species' dispersal distance is crucial
477	for <del>understandingpredicting</del> how quickly it can spread to new suitable habitats but also for
478	estimating its ability to respond to selection and <u>to</u> adapt. The balance between the
479	amount of gene flow and the strength of selection defines the probability of local
480	adaptation as a response to spatially diversifying selection, (Lenormand 2002). When the
481	environment is heterogeneous, locally adapted individuals are more likely to produce
482	offspring which succeed in the proximity of their parents. However, as gene flow from
483	differentiated populations causes migration load and hinders local adaptation (gene
484	swamping; Lenormand 2002). <u>Selection can be very efficient in species with large N<sub>e</sub>, such</u>
485	as Scots pine, and phenotypic climatic adaptation to different latitudes is well known

486	(Mikola 1982; Aho 1994; Notivol et al. 2007; Kujala & Knürr et al. 2017). Early mortality is
487	very high, and even when it is largely random, it also provides much opportunity for
488	selection. However, the question is not just how strong the selection is but also on what
489	spatial scale the species can track the environmental differences and adapt through
490	changes in allele frequencies. According to Slatkin (1973), A-a population can-only
491	respond to selection only if the underlying environmental heterogeneity occurs over a
492	distance longer than the characteristic length ( $L = \sigma / \sqrt{s}$ , where $\sigma$ is the offspring mean
493	dispersal distance and s is the strength of selection; Slatkin 1973) that is determined by
494	the length of offspring mean dispersal distance ( $\sigma$ ) and the strength of selection (s). With
495	our refined estimates of mean gene dispersal distance ( <u>53.9 m</u> <del>36.5-71.3 m, depending on</del> -
496	the effective population density used in estimation) and hypothetical selection coefficients
497	s = 0.01 or $s = 0.001$ , the characteristic length would be 539 m and 1 705 m, respectively
498	strong selection (Thurman & Barrett 2016), which is possible in steep ecological gradients
499	(Scotti et al. 2023) but unlikely in a homogeneous landscape such as our study area.rather
500	spatial scale would thus require smallAdaptation to a very a very fine local scale
501	(environmental change over 100 meters), the selection coefficient should be 0.29 (s = ( $\sigma$ /-
502	$L$ ) <sup>2</sup> , where L = 100 m and $\sigma$ = 53.9 m). no a population to adapt to For instance, in order , it
503	is possible to estimate the size of the selection coefficient needed for a particular
504	characteristic length.
505	No evidence for fine-scale adaptation has been found in Scots pine. Reciprocal transplant
506	experiments showed no evidence of local adaptation to different soils at the scale of some
507	kilometers (Jimenez-Ramírez et al. 2023). Furthermore, a nine-year common garden study
508	with progeny of Punkaharju ISS showed that selection at the local population scale was
509	rather weak on the adaptive seedling traits, even if fitness was lower in populations from
510	further north and south (Kujala et al. 2023). As adaptive traits are often polygenic,
511	selection on individual loci would be expected to be rather weak, yielding little potential to

512 respond to different selection in close-by sites. In some other European and

513 Mediterranean conifers, considerable selection coefficients were reported for individual loci

514 across steep ecological gradients, with at least 1 km distance and often hundreds of

515 <u>meters of altitudinal difference (Scotti *et al.* 2023).</u>

516 Fitness and practical implications of the fine-<u>-</u>scale spatial <u>genetic</u> structure

517 Aggregation of relatives leads to a higher chanceprobability of inbreeding and inbreeding depression. The Scots pine carries a high number of lethal equivalents (Koski 1971; 518 Savolainen et al. 1992), which makes selfing and more distant forms of inbreeding 519 520 detrimental. Our results indicate that related individuals carry an excess of shared rare 521 alleles. A large number of loci in Scots pine have very low minor allele frequencies (Tyrmi et al. 2020). Rare alleles are typically young and also enriched for recessive deleterious 522 variants, Thus, spatial genetic structure may lead to more homozygosity and fitness 523 524 reduction than expected in a totally panmictic population, where these alleles would rarelyseldom appear as homozygotes. 525

526 Fine-scale spatial genetic structure also has practical implications, e.g., in tree breeding. 527 When closely located trees are also more likely to be related, determining a suitable 528 collection distance of potential breeding individuals is very important in order to avoid introduction of related individuals into breeding programs. Possible inbreeding also needs 529 to be avoided in <u>and</u> production and deployment populations. Due to strong inbreeding 530 531 depression, manifesting as lowered yield of viable seeds and reduced viability and growth of the seedlings, accidental selection of related individuals to seed orchards would likely-532 lead to lowered yield of viable seedscause problems. Furthermore, information on spatial 533 534 genetic structures can help to define a minimum distance between trees to be used for collecting seed or cuttings in practical gene conservation work. Knowledge on the extent 535 536 and intensity of fine-scale spatial genetic structure is of importance also tofor forest management on of naturally regenerating sites as it can guide the optimization of distance 537 538 between spared seed trees during harvesting.

## 539 Conclusions

540 Here, we carefully described in detail the extent of fine-scale spatial genetic structure and average dispersal distance in a large population of Scots pine from a continuous part of 541 the distribution. We demonstrated that even a wind-pollinated widely distributed species 542 with large effective population size can have detectable, although weak, fine-\_scale 543 544 population spatial genetic structure. Our estimates of dispersal distance are relevant for 545 understanding the balance between gene flow and other evolutionary factors, especiallyselection, practical applications, as well as in predicting responses to environmental 546 changes, and understanding the balance between gene flow and other evolutionary 547 548 factors, especially selection.

#### 549 Acknowledgements

This project was funded by: the European Union's Horizon 2020 research and innovation 550 program, under grant agreement no. 773383 (to UOULU and Luke); the Seventh 551 framework program for research and development, under grant agreement no. 211868 (to 552 UOULU and Luke); the EU Network of Excellence EVOLTREE grant no. 016322 (to 553 UOULU and Luke); the Academy of Finland grants 287431, 293819 and 319313 (to 554 TP), 307582 (to OS), 307581 (to KK), and 309978 (to STK); NoE EVOLTREE (to STK). 555 We thank Natural Resources Institute Finland staff in Loppi and Punkaharju for collecting 556 557 samples and recording the GPS positions of the trees, and Soile Alatalo for molecular laboratory work at the Ecology and Genetics Unit of the University of Oulu, and Coen 558 559 Westerduin for comments on the previous version of the manuscript. We are grateful for 560 the PCI Evol Biol recommender Myriam Heuertz and the three reviewers Jean-Baptiste 561 Ledoux, Roberta Loh and Joachim Mergeay for their comments that improved the paper substantially. 562

#### 563 Conflicts of interest

The authors declare that they comply with the PCI rule of having no financial conflicts of interest in relation to the content of the article. The authors declare the following nonfinancial conflict of interest: Tanja Pyhäjärvi is a managing board member and a recommender of PCI Evol Biol.

#### 568 Supplementary Data

- 569 The following supplementary data is available as a separate document:
- 570 **Figure S1.** The distribution of Scots pine age in Mäkrä (turquoise; *n* = 113) and Ranta-
- 571 | <u>Halola (orange; *n* = 354) sampling sites.</u>
- 572 **Figure S2.** Decay of pairwise relatedness with distance in Mäkrä (turquoise) and Ranta-
- 573 Halola (orange) sampling sites. Mean (circles and squares) and standard deviation

574 (vertical lines) of relatedness is plotted for each distance class; number of pairwise

575 <u>comparisons in each distance class are shown in Table S1. The estimates of Ranta-Halola</u>

576 <u>have been moved two metres forward in the plot to avoid overlap with the estimates of</u>

577 <u>Mäkrä.</u>

578 **Figure S3.** Pairwise relatedness (GRM; Yang *et al.* 2011) plotted against pairwise kinship

579 (Loiselle; Loiselle et al. 1995) estimated for 468 Scots pines in the Punkaharju research

580 <u>area. The red line shows the expected relationship of 2:1 for these estimates.</u>

581 **Figure S4.** Correlation (measured as Mantel *r*) between pairwise relatedness and distance

582 within each distance class estimated as Mantel correlogram in a) Mäkrä (circles, turquoise)

583 and b) Ranta-Halola (squares, orange). Filled shape indicates a *p*-value smaller than 0.05.

584 Ranta-Halola and Mäkrä are divided into equally long distance classes. Two of the longest

585 distance classes from Mäkrä and three from Ranta-Halola have been left out due to

586 including less than 100 pairwise comparisons.

587 **Figure S5.** Decay of the proportion of shared rare alleles with spatial distance in Mäkrä

588 (turquoise) and Ranta-Halola (orange) sampling sites. Mean (circles and squares) and

589 <u>standard deviation (vertical lines) of relatedness is plotted for each distance class.</u>

590 **Figure S6.** Mantel correlogram for rare allele sharing and pairwise distance in a) Mäkrä

591 and b) Ranta-Halola. Filled circles indicate *p*-value smaller than 0.05.

592 **Table S1.** The mean and standard deviation (SD) of pairwise relatedness (GRM) in each

593 distance class for Ranta-Halola and Mäkrä sampling sites (Figure 3; Figure S2). The last

594 two and three distance classes for Ranta-Halola (upper panel) and Mäkrä, respectively,

595 are combined so that each class has at least 100 comparisons. The lower panel for Ranta-

596 Halola shows the relatedness values for Ranta-Halola, when distances are classified

597 according to Mäkrä's 14 distance classes.

- 598 **Table S2.** Comparison of family relationship classes estimated for 468 Scots pines in the
- 599 Punkaharju research area using relatedness (GRM; Yang *et al.* 2011) or kinship estimate
- 600 (Loiselle; Loiselle et al. 1995). Estimates on the darker green background show the same
- 601 degree of family relationship and ligher green shows one degree difference in the
- 602 <u>estimated class. Relatedness was estimated using 65 498 SNPs with MAF  $\geq$  0.05 and</u>
- 603 kinship using 28 378 SNPs with MAF  $\geq$  0.20 due to computational reasons.

#### 604 Data availability statement

- 605 The data supporting the findings of this study are available in Figshare (DOI:
- 10.6084/m9.figshare.23531142; the data will be open to public after this manuscript has
- 607 been accepted for publication).

## 608 References

- Ahmed A, Compton SG, Butlin RK & Gilmartin PM (2009) Wind-borne insects mediate directional pollen
   transfer between desert fig trees 160 kilometers apart. Proceedings of the National Academy of Sciences
   of the United States of America 106: 20342-20347. https://doi.org/10.1073/pnas.0902213106
- Aho M-L (1992) Autumn frost hardening of one-year-old *Pinus sylvestris* (L.) seedlings: Effect of origin and
   parent trees. Scandinavian Journal of Forest Research 9: 17-24.
   https://doi.org/10.1080/02827589409382808
- Aitken SN & Whitlock MC (2013) Assisted gene flow to facilitate local adaptation to climate change. Annual
   Review of Ecology, Evolution, and Systematics 44: 367-388. https://doi.org/10.1146/ANNUREV ECOLSYS-110512-135747
- 618 Barton NH (1979) The dynamics of hybrid zones. Heredity 43: 341-359. https://doi.org/10.1038/hdy.1979.87
- Beland M, Agestam E, Ekö PM, Gemmel P & Nilsson U (2000) Scarification and seedfall affects natural
   regeneration of Scots pine under two shelterwood densities and a clear-cut in southern Sweden.
   Scandinavian Journal of Forest Research 15: 247-255. https://doi.org/10.1080/028275800750015064
- Bizoux JP, Daïnou K, Bourland N, Hardy OJ, Heuertz M, Mahy G & Doucet JL (2009) Spatial genetic
  structure in *Milicia excelsa* (Moraceae) indicates extensive gene dispersal in a low-density windpollinated tropical tree. Molecular Ecology 18: 4398-4408. https://doi.org/10.1111/j.1365294X.2009.04365.x
- Born C, Hardy OJ, Chevallier MH, Ossari S, Attéké C, Wickings EJ & Hossaert-Mckey M (2008) Small-scale
  spatial genetic structure in the Central African rainforest tree species *Aucoumea klaineana*: A stepwise
  approach to infer the impact of limited gene dispersal, population history and habitat fragmentation.
  Molecular Ecology 17: 2041–2050. https://doi.org/10.1111/j.1365-294X.2007.03685.x
- Bradburd G & Ralph P (2019) Spatial population genetics: it's about time. Annual Review of Ecology,
   Evolution, and Systematics 50: 427-449. https://doi.org/10.1146/annurev-ecolsys-110316-022659
- Debain S, Chadœuf J, Curt T, Kunstler G & Lepart J (2007) Comparing effective dispersal in expanding
   population of *Pinus sylvestris* and *Pinus nigra* in calcareous grassland. Canadian Journal of Forest
   Research 37: 705-718. https://doi.org/10.1139/X06-265
- De-Lucas AI, González-Martínez SC, Vendramin GG, Hidalgo E & Heuertz M (2009) Spatial genetic structure
   in continuous and fragmented populations of *Pinus pinaster* Aiton. Molecular Ecology 18: 4564-4576.
   https://doi.org/10.1111/j.1365-294X.2009.04372.x
- Desilva R & Dodd RS (2021) Patterns of fine-scale spatial genetic structure and pollen dispersal in giant
   sequoia (*Sequoiadendron giganteum*). Forests 12: 1–14. https://doi.org/10.3390/f12010061
- Escudero A, Iriondo JM & Torres ME (2003) Spatial analysis of genetic diversity as a tool for plant
   conservation. Biological Conservation 113: 351–365. https://doi.org/10.1016/S0006-3207(03)00122-8
- Excoffier L, Smouse PE & Quattro JM (1992) Analysis of molecular variance inferred from metric distances
   among DNA haplotypes: Application to human mitochondrial DNA restriction data. Genetics 131: 479 491.
- Fahlvik N, Ekö P-M & Pettersson N (2005) Influence of precommercial thinning grade on branch diameter
   and crown ratio in *Pinus sylvestris* in southern Sweden. Scandinavian Journal of Forest Research 20:
   243-251. https://doi.org/10.1080/02827580510008266
- González-Díaz P, Jump AS, Perry A, Wachowiak W, Lapshina E & Cavers S (2017) Ecology and
   management history drive spatial genetic structure in Scots pine. Forest Ecology and Management 400:
   68-76. https://doi.org/10.1016/J.FORECO.2017.05.035
- Goslee SC & Urban DL (2007) The ecodist package for dissimilarity-based analysis of ecological data.
   Journal of Statistical Software 22: 1-19.

Gougherty AV, Keller SR & Fitzpatrick MC (2021) Maladaptation, migration and extirpation fuel climate
 change risk in a forest tree species. Nature Climate Change 11: 166-171. https://doi.org/10.1038/s41558 020-00968-6

- Hampe A & Petit RJ (2005) Conserving biodiversity under climate change: the rear edge matters. Ecology
   Letters 8: 461-467. https://doi.org/10.1111/j.1461-0248.2005.00739.x
- Hardy OJ & Vekemans X (1999) Isolation by distance in a continuous population: Reconciliation between
   spatial autocorrelation analysis and population genetics models. Heredity 83: 145–154.
   https://doi.org/10.1046/j.1365-2540.1999.00558.x
- Hardy OJ & Vekemans X (2002) SPAGeDi: a versatile computer program to analyse spatial genetic structure
   at the individual or population levels. Molecular Ecology Notes 2: 618-620. https://doi.org/10.1046/j.1471 8286.2002.00305.x
- Hardy OJ, Maggia L, Bandou E, Breyne P, Caron H, Chevallier MH, Doligez A, Dutech C, Kremer A,
  Latouche-Hallé C, Troispoux V, Veron V & Degen B (2006) Fine-scale genetic structure and gene
  dispersal inferences in 10 Neotropical tree species. Molecular Ecology 15: 559–571.
  https://doi.org/10.1111/j.1365-294X.2005.02785.x
- Jiménez-Ramírez A, Grivet D & Robledo-Arnuncio JJ (2021) Measuring recent effective gene flow among
   large populations in *Pinus sylvestris*: Local pollen shedding does not preclude substantial long-distance
   pollen immigration. PLoS ONE 16: e0255776. https://doi.org/10.1371/journal.pone.0255776
- 671 Jiménez-Ramírez A, Solé-Medina A, Ramírez-Valiente JA & Robledo-Arnuncio JJ (2023) Microgeographic
   672 variation in early fitness traits of *Pinus sylvestris* from contrasting soils. American Journal of Botany 110:
   673 e16159. https://doi.org/10.1002/ajb2.16159
- 674 Kahle D & Wickham H (2013) ggmap: Spatial Visualization with ggplot2. The R Journal 5: 144–161.
   675 https://journal.r-project.org/archive/2013-1/kahle-wickham.pdf
- Kastally C, Niskanen AK, Perry A, Kujala ST, Avia K, Cervantes S, Haapanen M, Kesälahti R, Kumpula TA,
  Mattila TM, Ojeda DI, Tyrmi JS, Wachowiak W, Cavers S, Kärkkäinen K, Savolainen O & Pyhäjärvi T
  (2022) Taming the massive genome of Scots pine with PiSy50k, a new genotyping array for conifer
  research. The Plant Journal 109: 1337-1350. https://doi.org/10.1111/TPJ.15628
- Kitamura K, Nakanishi A, Lian C & Goto S (2018) Distinctions in fine-scale spatial genetic structure between
   growth stages of *Picea jezoensis* Carr. Frontiers in Genetics 9: 490.
   https://doi.org/10.3389/fgene.2018.00490
- Koelewijn HP, Koski V & Savolainen O (1999) Magnitude and timing of inbreeding depression in Scots pine
   (*Pinus sylvestris* L.). Evolution 53: 758-768. https://doi.org/10.1111/J.1558-5646.1999.TB05370.X
- Koski V (1970) A study of pollen dispersal as a mechanism of gene flow in conifers. Metsätieteellisen
   tutkimuslaitoksen julkaisuja 70(4).
- Koski V (1971) Embryonic lethals of *Picea abies* and *Pinus sylvestris*. Metsätieteellisen tutkimuslaitoksen
   julkaisuja 75(3).
- Kremer A, Ronce O, Robledo-Arnuncio JJ, Guillaume F, Bohrer G, Nathan R, Bridle JR, Gomulkiewicz R,
   Klein EK, Ritland K, Kuparinen A, Gerber S & Schueler S (2012) Long-distance gene flow and adaptation
   of forest trees to rapid climate change. Ecology Letters 15: 378-392. https://doi.org/10.1111/J.1461 0248.2012.01746.X
- Kujala ST, Knürr T, Kärkkäinen K, Neale DB, Sillanpää MJ & Savolainen O (2017) Genetic heterogeneity
   underlying variation in a locally adaptive clinal trait in *Pinus sylvestris* revealed by a Bayesian
   multipopulation analysis. Heredity 118: 413–423. https://doi.org/10.1038/hdy.2016.115
- Kujala ST, Avia K, Kumpula TA, Kärkkäinen H, Heikkinen J, Kärkkäinen K & Savolainen O (2023) Within and between-population comparisons suggest independently acting selection maintaining parallel clines
   in Scots pine (*Pinus sylvestris*). Evolution Letters grad054. https://doi.org/10.1093/evlett/grad054
- Kuparinen A, Savolainen O & Schurr FM (2010) Increased mortality can promote evolutionary adaptation of
   forest trees to climate change. Forest Ecology and Management 259: 1003-1008.
   https://doi.org/10.1016/j.foreco.2009.12.006
- Kärkkäinen K & Savolainen O (1993) The degree of early inbreeding depression determines the selfing rate
   at the seed stage: Model and results from *Pinus sylvestris* (Scots pine). Heredity 71: 160-166.
   https://doi.org/10.1038/hdy.1993.120
- Kärkkäinen K, Koski V & Savolainen O (1996) Geographical variation in the inbreeding depression of Scots
   pine. Evolution 50: 111-119. https://doi.org/10.1111/j.1558-5646.1996.tb04477.x

- Legendre P, Fortin MJ & Borcard D (2015) Should the Mantel test be used in spatial analysis? Methods in
   Ecology and Evolution 6: 1239–1247. https://doi.org/10.1111/2041-210X.12425
- Lenormand T (2002) Gene flow and the limits to natural selection. Trends in Ecology & Evolution 17: 183 189. https://doi.org/10.1016/S0169-5347(02)02497-7
- Levin DA & Kerster HW (1974) Gene flow in seed plants. In *Evolutionary Biology*, eds. Th. Dobzhansky, M.
   K. Hecht, W. C. Steere, New York: Plenum 7: 139-220.
- Lindgren D, Paule L, Xihuan S, Yazdani R, Segerström U, Wallin J-E & Lejdebro ML (1995) Can viable pollen
   carry Scots pine genes over long distances? Grana 34: 64-69.
   https://doi.org/10.1080/00173139509429035
- 716 Linder P, Elfving B & Zackrisson O (1997) Stand structure and successional trends in virgin boreal forest
   717 reserves in Sweden, Forest Ecology and Management 98: 17-33, https://doi.org/10.1016/S0378-
- 717
   reserves in Sweden. Forest Ecology and Management 98: 17-33. https://doi.org/10.1016/S0378 

   718
   1127(97)00076-5
- Loiselle BA, Sork VL, Nason J & Graham C (1995) Spatial genetic structure of a tropical understorey shrub,
   *Psychotria officinalis* (Rubiaceae). American Journal of Botany 82: 1420–1425.
- Loveless MD & Hamrick JL (1984) Ecological determinants of genetic structure in plant populations. Annual
   review of ecology and systematics 15: 65-95.
- Lundqvist L, Ahlström MA, Axelsson EP, Mörling T & Valinger E (2019) Multi-layered Scots pine forests in
   boreal Sweden result from mass regeneration and size stratification. Forest Ecology and Management
   441: 176-181. https://doi.org/10.1016/j.foreco.2019.03.044
- Lönnroth E (1926) Untersuchungen über die innere Struktur und Entwicklung gleichaltriger naturnormaler Kiefernbestände: Basiert auf Material aus der Südhälfte Finnlands. Acta Forestalia Fennica 30: 1-269.
- Malécot G (1967) Identical loci and relationship. In *Proceedings of the Fifth Berkeley Symposium on Mathematical Statistics and Probability*, eds. L.M. LeCam and J. Neyman, University of California Press,
   Berkeley 4: 317–332.
- Manichaikul A, Mychaleckyj JC, Rich SS, Daly K, Sale M & Chen WM (2010) Robust relationship inference in
   genome-wide association studies. Bioinformatics 26: 2867-2873.
   https://doi.org/10.1093/bioinformatics/btq559
- Mantel N (1967) The detection of disease clustering and a generalized regression approach. Cancer
   Research 27: 209–220.
- 736 Marris E (2009) Planting the forest of the future. Nature 459: 906–908. https://doi.org/10.1038/459906a
- Mikola J (1982) Bud-set phenology as an indicator of climatic adaptation of Scots pine in Finland. Silva
   Fennica 16: 178–184.
- Muona O & Harju A (1989) Effective population sizes, genetic variability, and mating system in natural stands
   and seed orchards of *Pinus sylvestris*. Silvae Genetica 38: 221–229.
- Neale DB, Wegrzyn JL, Stevens KA, Zimin AV, Puiu D, Crepeau MW *et al.* (2014) Decoding the massive genome of loblolly pine using haploid DNA and novel assembly strategies. Genome Biology 15: R59.
   https://doi.org/10.1186/gb-2014-15-3-r59
- Nei M (1972) Genetic distance between populations. The American naturalist 106: 283-292.
- Notivol E, García-Gil MR, Alía R & Savolainen O (2007) Genetic variation of growth rhythm traits in the limits
   of a latitudinal cline in Scots pine. Canadian Journal of Forest Research 37: 540–551.
   https://doi.org/10.1139/X06-243
- Nychka D, Furrer R, Paige J & Sain S (2017) fields: Tools for spatial data. R package version 11.6.
   https://doi.org/10.5065/D6W957CT

Pandey M & Rajora O (2012) Higher fine-scale genetic structure in peripheral than in core populations of a
 long-lived and mixed-mating conifer - Eastern white cedar (*Thuja occidentalis* L.). BMC Evolutionary
 Biology. 12: 48. https://doi.org/10.1186/1471-2148-12-48

# Pembleton L, Cogan N & Forster J (2013) StAMPP: an R package for calculation of genetic differentiation and structure of mixed-ploidy level populations. Molecular Ecology Resources 13: 946-952. https://doi.org/10.1111/1755-0998.12129

- Persyn E, Redon R, Bellanger L & Dina C (2018) The impact of a fine-scale population stratification on rare
   variant association test results. PLoS ONE 13: e0207677. https://doi.org/10.1371/journal.pone.0207677
- Pritchard JK & Rosenberg NA (1999) Use of unlinked genetic markers to detect population stratification in association studies. American Journal of Human Genetics 65: 220-228. https://doi.org/10.1086/302449
- Privé F, Luu K, Vilhjálmsson BJ, Blum MGB & Rosenberg M (2020) Performing highly efficient genome scans
   for local adaptation with R package pcadapt version 4. Molecular Biology and Evolution 37: 2153–2154.
   https://doi.org/10.1093/molbev/msaa053
- Purcell S, Neale B, Todd-Brown K, Thomas L, Ferreira MAR, Bender D, Maller J, Sklar P, de Bakker PIW,
   Daly MJ & Sham PC (2007) PLINK: a tool set for whole-genome association and population-based
   linkage analyses. The American Journal of Human Genetics 81: 559–575.
   https://doi.org/10.1086/519795
- Pyhäjärvi T, Kujala ST & Savolainen O (2020) 275 years of forestry meets genomics in *Pinus sylvestris*.
   Evolutionary Applications 13: 11-30. https://doi.org/10.1111/eva.12809
- 769 <u>R Core Team (2020). R: A language and environment for statistical computing (version 3.6.3). R Foundation</u>
   770 <u>for Statistical Computing, Vienna, Austria. https://www.R-project.org/</u>
- Rautio P, Hallikainen V, Valkonen S, Karjalainen J, Puttonen P, Bergsten U, Winsa H & Hyppönen M (2023)
   Manipulating overstory density and mineral soil exposure for optimal natural regeneration of Scots pine.
   Forest Ecology and Management 539: 120996. https://doi.org/10.1016/j.foreco.2023.120996
- Robledo-Arnuncio JJ & Gil L (2005) Patterns of pollen dispersal in a small population of *Pinus sylvestris* L.
   revealed by total-exclusion paternity analysis. Heredity 94: 13-22. https://doi.org/10.1038/sj.hdy.6800542
- Robledo-Arnuncio JJ (2011) Wind pollination over mesoscale distances: an investigation with Scots pine.
   New Phytologist 190: 222-233. https://doi.org/10.1111/J.1469-8137.2010.03588.X
- Rousset F (1997) Genetic differentiation and estimation of gene flow from F-statistics under isolation by
   distance. Genetics 145: 1219-1228. https://doi.org/10.1002/ajmg.c.30221
- Rousset F (2000) Genetic differentiation between individuals. Journal of Evolutionary Biology 13: 58–62.
   https://doi.org/10.1046/J.1420-9101.2000.00137.X
- Rousset F (2003) Inferences from spatial population genetics, in: *Handbook of statistical genetics*, eds. D. J.
   Balding, M. Bishop, & C. Cannings. John Wiley & Sons.
- Sant'Anna CS, Sebbenn AM, Klabunde GHF, Bittencourt R, Nodari RO, Mantovani A & dos Reis MS (2013)
   Realized pollen and seed dispersal within a continuous population of the dioecious coniferous Brazilian
   pine [*Araucaria angustifolia* (Bertol.) Kuntze]. Conservation Genetics 14: 601-613.
   https://doi.org/10.1007/S10592-013-0451-5
- Savolainen O, Kärkkäinen K & Kuittinen H (1992) Estimating numbers of embryonic lethals in conifers.
   Heredity 69: 308-314.
- Scotti I, Lalagüe H, Oddou-Muratorio S, Scotti-Saintagne C, Ruiz Daniels R, Grivet D, *et al.* (2023) Common
   microgeographical selection patterns revealed in four European conifers. Molecular Ecology 32: 393-411.
   https://doi.org/10.1111/mec.16750
- 793 Slatkin M (1973) Gene flow and selection in a cline. Genetics 75: 733-756.
- 794 Slatkin M (1985) Gene flow in natural populations. Annual Review of Ecology and Systematics 16: 393-430.
- Smith JR, Ghazoul J, Burslem DFRP, Itoh A, Khoo E, Lee SL, Maycock CR, Nanami S, Ng KKS & Kettle CJ
   (2018) Are patterns of fine-scale spatial genetic structure consistent between sites within tropical tree
   species? PLoS ONE 13: e0193501. https://doi.org/10.1371/journal.pone.0193501
- Sofletea N, Mihai G, Ciocîrlan E & Curtu AL (2020) Genetic diversity and spatial genetic structure in isolated
   Scots pine (*Pinus sylvestris* L.) populations native to Eastern and Southern Carpathians. Forests 11:
   1047. https://doi.org/10.3390/f11101047
- 801 Thurman TJ & Barrett RDH (2016) The genetic consequences of selection in natural populations. Molecular 802 Ecology 25: 1429-1448. https://doi.org/10.1111/mec.13559-
- 803

- Torimaru T, Wennström U, Lindgren D & Wang X-R (2012) Effects of male fecundity, interindividual distance
   and anisotropic pollen dispersal on mating success in a Scots pine (*Pinus sylvestris*) seed orchard.
   Heredity 108: 312-321. https://doi.org/10.1038/hdy.2011.76
- Tyrmi JS, Vuosku J, Acosta JJ, Li Z, Sterck L, Cervera MT, Savolainen O & Pyhäjärvi T (2020) Genomics of
   clinal local adaptation in *Pinus sylvestris* under continuous environmental and spatial genetic setting. G3:
   Genes, Genomes, Genetics 10: 2683-2696. https://doi.org/10.1534/g3.120.401285
- Vakkari P, Rusanen M, Heikkinen J, Huotari T & Kärkkäinen K (2020) Patterns of genetic variation in leading edge populations of *Quercus robur*: genetic patchiness due to family clusters. Tree Genetics and
   Genomes 16: 73. https://doi.org/10.1007/s11295-020-01465-9
- Varis S, Pakkanen A, Galofré A & Pulkkinen P (2009) The extent of South-North pollen transfer in Finnish
   Scots pine. Silva Fennica 43: 717-726.
- Vekemans X & Hardy O (2004) New insights from fine-scale spatial genetic structure analyses in plant populations. Molecular Ecology 13: 921-935. https://doi.org/10.1046/j.1365-294X.2004.02076.x
- Väisänen H, Kellomäki S, Oker-Blom P & Valtonen E (1989) Structural development of *Pinus sylvestris*stands with varying initial density: A preliminary model for quality of sawn timber as affected by
  silvicultural measures. Scandinavian Journal of Forest Research 4: 223-238.
  https://doi.org/10.1080/02827588909382560
- Weir BS & Cockerham CC (1984) Estimating F-statistics for the analysis of population structure. Evolution
   38: 1358–1370. https://doi.org/10.2307/2408641
- Wegrzyn JL, Liechty JD, Stevens KA, Wu L-S, Loopstra CA, Vasquez-Gross HA, *et al.* (2014) Unique
   features of the loblolly pine (*Pinus taeda* L.) megagenome revealed through sequence annotation.
   Genetics 196: 891-909. https://doi.org/10.1534/genetics.113.159996
- Willyard A, Syring J, Gernandt DS, Liston A & Cronn R (2007) Fossil calibration of molecular divergence
   infers a moderate mutation rate and recent radiations for *Pinus*. Molecular Biology and Evolution 24: 90 101. https://doi.org/10.1093/MOLBEV/MSL131
- 829 Wright S (1943) Isolation by distance. Genetics 28: 114–138.
- 830 Wright S (1946) Isolation by distance under diverse systems of mating. Genetics 31: 39–59.
- Yang J, Lee SH, Goddard ME & Visscher PM (2011) GCTA: a tool for genome-wide complex trait analysis.
   The American Journal of Human Genetics 88: 76–82. https://doi.org/10.1016/j.ajhg.2010.11.011