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Salmela, Matti J.; Cavers, Stephen; Cottrell, Joan E.; Iason, Glenn R.; Ennos, Richard A. 2013. Spring phenology shows genetic variation among and within populations in seedlings of Scots pine (Pinus sylvestris L.) in the Scottish Highlands.

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1	Spring phenology she	ows genetic variation	among and within	populations in s	eedlings of
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2 Scots pine (Pinus sylvestris L.) in the Scottish Highlands

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- 28 Abstract
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30 Background: Genetic differentiation in phenotypic traits is often observed among forest tree

- 31 populations, but less is known about patterns of adaptive variation within populations. Such
- 32 variation is expected to enhance the survival likelihood of extant populations under climate
- 33 change.
- 34 Aims: Scots pine (Pinus sylvestris) occurs over a spatially and temporally heterogeneous
- 35 landscape in Scotland. Our goal was to examine whether populations had differentiated
- 36 genetically in timing of bud flush in response to spatial heterogeneity and whether variation
- 37 was also maintained within populations.

38 Methods: Two common-garden studies, involving maternal families of seedlings from 21

39 native pinewoods, were established and variation in the trait was measured at the beginning of

- 40 the second growing season.
- 41 Results: Populations showed genetic differences in the trait correlated with the length of
- 42 growing season at their site of origin, but the majority of variation was observed within
- 43 populations. Populations also differed in their levels of variation in the trait; a pattern that
- 44 may be influenced by spatial variation in the extent of temporal climate variability.
- 45 Conclusions: Our findings suggest that populations have adapted to their home environments46 and that they also have substantial ability to adapt in situ to changes in growing season length.

47

- 48 Keywords: adaptation, adaptive potential, genetic differentiation, spatial heterogeneity,
- 49 temporal heterogeneity, variation within populations
- 50 51 52 53 54

When a species is distributed across a spatially heterogeneous landscape, natural selection is 58 59 expected to favour different trait optima in divergent environments. For example, growth in plants may continue furthest into the autumn in populations that experience the longest 60 61 growing seasons (e.g. Mikola 1982). This may lead to genetic differentiation in selected 62 phenotypic traits among populations and each population surviving and growing best at its 63 home site, i.e. local adaptation (Kawecki and Ebert 2004). Adaptations to local environments 64 have been described in many plant species (Linhart and Grant 1996), and in trees, numerous common-garden studies have demonstrated that, for example, growth phenology and timing 65 of cold hardiness are optimised to local climates (Howe et al. 2003; Savolainen et al. 2007). 66 Local adaptation has also been demonstrated in transfer trials in which populations have been 67 68 grown at home and foreign sites (e.g. Persson and Ståhl 1990); a possible reason for poorer survival and growth at foreign sites is a mismatch between annual climatic variation and 69 70 growth phenology (Eriksson et al. 1980).

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Ongoing rapid climate change is affecting ecosystems globally, which is evident as range 72 shifts and changes in timing of growth and reproductive events in several species (Parmesan 73 2006). Due to the commercial importance of many species and conservation issues, predicting 74 evolutionary responses to a changing climate has become an active area of research (Hendry 75 et al. 2011; Hoffman and Sgro 2011). For example, ecological modelling often considers 76 77 associations between species distributions and environmental factors: following environmental change, ranges may expand into new areas that become suitable for a species, 78 79 while extinction may take place at sites that become unfavourable (Elith and Leathwick 80 2009). In trees, common-garden experiments replicated in multiple environments have been used to estimate how environmental changes will affect contemporary populations and to 81 82 demonstrate that in the future, some populations might be exposed to non-optimal conditions 83 which might result in poorer growth (Rehfeldt et al. 2002; Reich and Oleksyn 2008). However, such approaches can only be used to examine how existing populations will 84 respond to environmental changes as they neglect an important feature of natural populations: 85 their capacity to adapt to environmental changes in situ (Hoffman and Sgrò 2011). This 86

87 capability means that, as well as range shifts, responses to environmental change should also 88 involve genetic changes between generations exposed to different environments, allowing 89 populations to adapt to new conditions. Indeed, taking adaptation and phenotypic plasticity 90 into account in models can greatly enhance the survival likelihood of current populations 91 (Aitken et al. 2008; Benito Garzón et al. 2011). Adaptation to novel environments has certainly taken place in many species when they expanded their ranges following the last ice 92 age (Davis and Shaw 2001), but it is possible that future changes will occur too rapidly for 93 94 populations to track them (Savolainen et al. 2004; Aitken et al. 2008) or that intensively 95 managed contemporary landscapes will be impermeable to migration.

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In situ adaptive responses to a changing environment are possible when phenotypes vary 97 98 within populations and when such variation is due to genetic factors (Hoffman and Sgrò 99 2011). However, research on adaptation has largely focused on examining how populations 100 differ from each other in terms of trait means and how environmental differences among the sites occupied by different populations contribute to such divergence (reviewed e.g. in 101 Savolainen et al. 2007). Thus, we currently have only a limited understanding of patterns of 102 adaptive trait variation within populations (Kramer and Havens 2009). When populations 103 104 adapt to their home environments, variation in traits affected by selection is expected to be 105 lost as individuals that differ too much from the local optimum have poorer chances of survival (e.g. Falconer and Mackay 1996). Still, it is commonly found across different kinds 106 of organisms that significant genetic variation can be preserved even in traits under the 107 strongest type of selection (Houle 1992; Merilä and Sheldon 1999; Barton and Keightley 108 109 2002). Also in trees, within-population variation in traits under selection has been widely 110 found in common-garden studies (Howe et al. 2003). A similar pattern can also be seen in 111 trees observed in their natural habitats: in a stand of *Betula pendula* Roth growing in south-112 eastern Finland, trees flushed at different times, among-tree differences being the smallest 113 during warm springs (Rousi and Heinonen 2007). The reasons for the persistence of such 114 variation in nature despite natural selection remain poorly understood, but a number of factors 115 might contribute to the maintenance of adaptive genetic diversity in forest trees. Due to their 116 longevity, trees are likely to experience a wide range of environmental conditions during their 117 lifespan (Petit and Hampe 2006), and mechanisms enabling individuals to modify their 118 phenotype according to the environment are expected to evolve under such conditions (Bull

119 1987). Indeed, phenotypic plasticity in initiation of growth allows trees to survive in 120 environments where temperature conditions in spring vary among years (Rousi and Heinonen 121 2007; Chmura et al. 2011). It is also possible that a population is found over a spatially 122 variable area so that selection pressure also varies across short distances (i.e., there is no 123 single trait optimum in a population; e.g. Campbell 1979), or that the environment varies between years so that different age groups within populations may have experienced differing 124 selection pressure. This may maintain diversity if selection acts only on specific age groups 125 (e.g. young seedlings) while having less effect on others (Ellner and Hairston 1994). Variation 126 127 may also be introduced by gene flow via pollen from environmentally different sites (Yeaman and Jarvis 2006). What the relative contributions of these factors in nature are remains largely 128 129 an unexplored topic.

130

Adaptive potential can be compromised especially in small and fragmented populations in 131 132 which random factors such as sudden population size changes may have shaped patterns of genetic variation more than natural selection (e.g. Willi et al. 2006). Scots pine (Pinus 133 sylvestris L.) is the only pine native to northern Europe and has an extensive distribution 134 across Eurasia (Critchfield and Little 1966). The Scottish populations of the species are 135 geographically separated on the north-western edge of this range and have been subjected to 136 137 heavy human interference in the past. Currently, 84 discrete native pinewood sites of variable size are recognised by the Forestry Commission of Great Britain which cover only about 1% 138 of their original postglacial maximum areal cover (Mason et al. 2004). Scots pine is a 139 140 foundation species upon which the persistence of many of the species in Scottish forests 141 depends. The native pinewoods are found over a geographically small but spatially highly 142 heterogeneous landscape, with steep gradients in temperatures and precipitation between the 143 oceanic west coast and the more continental east (Salmela et al. 2010). Despite a significant 144 decrease in abundance which has led to fragmentation, most of the populations are as diverse 145 at selectively neutral molecular markers as more continuous continental populations and show 146 very little differentiation for these neutral markers (Kinloch et al. 1986; Wachowiak et al. 147 2010). This suggests that at least historically, these populations have been connected by gene 148 flow. However, little is currently known about the patterns of adaptive trait variation in this 149 part of the species' distribution. In a recent experiment under natural climate conditions in 150 south-eastern Scotland, plants from eight populations were found to differ in their response to

151 winter and spring temperatures, which suggested environment-driven genetic differentiation

among some of the populations despite the small geographic scale (Salmela et al. 2011).

153 Similar differentiation was found for timing of growth initiation in spring, which was earlier

154 in populations from cooler, high-altitude locations.

155

In common with other parts of the world, increases in summer and winter temperatures and 156 changes in rainfall patterns are expected in Scotland in the coming decades (Ray 2008). 157 158 possibly leading to changes in selection pressures for traits related to timing of growth in 159 Scots pine. For current populations, these changes in climate have been predicted to be 160 detrimental (Ray 2008). However, the possibility of adaptation within populations has not been considered in these predictions which might result in their conclusions being too 161 162 conservative and potentially in management actions detrimental to the genetic integrity of 163 current populations. Considering how allowing for adaptation influences model predictions on 164 the effects of climate change in tree populations (Aitken et al. 2008), it is important for the conservation of the remaining native pinewood resources that the patterns of adaptive trait 165 variation among and within populations are investigated (Salmela et al. 2010). 166

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Due to the highly variable climate conditions that they are found in, adaptive genetic 168 differentiation is expected to have taken place among pine populations from different parts of 169 Scotland. In addition to spatial environmental variation among and within populations, the 170 171 Scottish climate is also characterised by temporal (among-year) fluctuations, for instance in 172 the length of the growing season (Perry and Hollis 2005) and winter severity (Harrison 1997). 173 Such fluctuations probably account for phenomena such as the observed temporal variation in 174 timing of bud flush under natural climate conditions in two birch species, Betula pubescens Ehrh, and *B. pendula* (Billington and Pelham 1991). The effects of climate fluctuations have 175 176 also been recognised in animals: sheep mortality on the island of St. Kilda has been found to 177 be higher in wet and warm winters which often coincide with positive phases of North 178 Atlantic Oscillation (Milner et al. 1999), a climatic phenomenon linked to the strength of 179 westerly winds across the northern Atlantic (Stenseth et al. 2002). Although temporally 180 variable selection has been suggested as one factor that may contribute to the maintenance of 181 adaptive diversity in long-lived trees (Howe et al. 2003; Westfall and Millar 2004; Yeaman

185 capacity. In this study, we used Scots pine as a model system to examine how a phenological 186 trait varied among and within populations sampled across a spatially and temporally 187 heterogeneous landscape. More specifically, we collected families from 21 environmentally 188 diverse sites in Scotland and grew their progeny in two separate glasshouse experiments to 189 address the following questions: 190 191 1) When grown under common-garden conditions, are 21 native Scots pine populations 192 differentiated for timing of growth initiation in spring at the beginning of their second 193 growing season? 194 2) Are observed population differences associated with environmental variation among 195 their home sites? 3) Is there significant variation within populations? If so, what is the pattern of this 196 variation? 197 4) Could the relative amounts of within-population variation be accounted for by within-198 site spatial and/or temporal variation in environment? 199

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- 202 Materials and methods

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Measuring genetic diversity in timing of bud flush 205

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- 207 Phenotypes of adaptive traits are determined by both genetic factors and the environment
- 208 (Falconer and Mackay 1996). Therefore, to reveal differences due to the genetic component,

and Jarvis 2006), its potential role is yet to be studied in more detail. If populations are

exposed to different levels of spatial and temporal environmental heterogeneity, they might

also differ in the level of genetic diversity in adaptive traits and consequently their adaptive

- 209 samples from different populations must be raised in a common-garden environment. To
- 210 estimate the levels of variation within populations, a family-structured design is needed so

211 that total variation in phenotype can be partitioned into among- and within-population

212 components. In tree populations, this can be accomplished by sampling multiple open-

213 pollinated seed from a number of mother trees in each population (e.g. White et al. 2007).

214 Due to high outcrossing rates (mother trees are generally pollinated by a large number of

215 pollen donors), such progeny are often assumed to consist mostly of half-siblings (i.e., family

216 members share only the maternal parent).

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219 Study populations

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A total of 21 native populations were sampled for this study, representing all parts of the
species' range in Scotland (Figure 1, Table 1). Cones were collected from 10 maternal trees in
each population in March 2007. Open-pollinated seed was extracted from cones and stored by
family.

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227 Provenance/progeny trials

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229 Sampled seed was used to establish two glasshouse-based common-garden trials located in 230 Edinburgh and Aberdeen (Figure 1) in late spring 2007. The two trials were set up by 231 independent investigators and they consequently had rather different germination conditions 232 and layout designs. In the Edinburgh trial located at the Centre for Ecology and Hydrology (55.86° N, 3.21° W), seed were sampled from four mother trees per population (i.e. 84 233 234 families in total) and sown on trays (75:25 compost type John Innes 1: sand) in June 2007 235 under common-garden glasshouse conditions. After germination, seedlings were transferred to 236 pots of size 0.62 l (diameter 11 cm, depth 9.6 cm) and kept under natural light conditions 237 (glasshouse was shaded to avoid excess light) with watering applied two or three times per 238 week during the growing season. No heating was applied during winter. Each family 239 consisted of 40 progeny (~3,360 seedlings in total). The trial was divided into 40 blocks, each

240 having one member from each of 84 families, and the order of the families within blocks was

randomised.

242

243 The trial in Aberdeen was located at the James Hutton Institute (57.13° N, 2.16° W). Seed 244 were sampled from 10 mother trees per population (i.e. 210 families in total). Cones were 245 placed in a warm room (30 °C) for two weeks so that they opened and seed could be extracted 246 for germination. Seed from the individual trees were kept separate and were soaked in water for 3 hours, then laid between sheets of damp paper towel placed in a cool room (3° C) for 247 248 several weeks to break dormancy. Seed were taken out of the cool room and left (wrapped in 249 damp paper) in the laboratory at room temperature until they germinated. Germination took 250 approximately 7 days and seed from all the sampled families germinated at this time. On 251 germination they were transplanted into potting medium in the glasshouse into $8 \times 8 \times 9$ cm 252 (0.4 l) pots. Each family consisted of eight progeny (~1,680 seedlings in total). The trial was 253 divided into 40 blocks with 42 plants per block, each block containing two plants from a different mother from each population. The 84 mother trees sampled in the Edinburgh trial 254 255 were a subset of those included in the Aberdeen trial. Watering was applied automatically and 256 no artificial light was used.

257

In both trials, growth initiation at the beginning of the second growing season was considered to have taken place when new green needle tips started to emerge from the apical bud (bud or needle flush), and this was measured as the number of days since the first scoring date. Bud flush was scored twice weekly in Edinburgh between March 23 and May 9 2008 and once weekly in Aberdeen between March 31 and May 27 2008.

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265 *Testing for genetic differentiation among populations and families*

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267 Data from the Edinburgh trial were analysed using nested analysis of variance (ANOVA),

268 with populations considered as fixed and families within them and blocks as random factors.

269 Unbalanced nested ANOVA was applied to the data from the Aberdeen trial. To examine the

270 relative contributions of different factors to total variation in the trait, variance components

271 due to populations, families, and blocks were estimated using the restricted maximum

272 likelihood (REML) approach. Correlation analysis was used to test whether similar trends

273 were observed in the two trials. All statistical analyses in this study were carried out using

274 GenStat Ver. 13.1.0.4470.

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- 277 Measuring the level of variation within populations
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To examine the variability of the trait within each population in more detail, standard 279 280 deviations (SD_{POP}) were examined separately in the two trials. SD's were calculated using raw values because of their strong correlation (r=0.94-0.99) with values adjusted for the effects of 281 282 individual blocks. Because timing of bud flush does not have fixed means (i.e., means vary depending on the date from which the timing is calculated), we used SD's instead of 283 284 normalised measures of dispersion (see e.g. Garcia-Gonzalez et al. 2012). Correlation analysis 285 was used to test whether similar patterns of variation among SD_{POP}'s were observed in the 286 two trials.

287

In a study design consisting of families grouped within populations, within-population 288 289 variation can arise from two components: among families and within families (residual 290 variation). Variation among families is considered to reflect the level of heritable (additive) 291 genetic variation (Falconer and Mackay 1996), and populations with higher levels of heritable 292 variation are expected to have better adaptive potential (Houle 1992). Because the amount of 293 additive genetic variation is directly proportional to the amount of variation among families 294 (Falconer and Mackay 1996), we only used estimates of among-family variation in the 295 analyses presented. In order to examine whether the level of among and within-family 296 variation differed among populations, the REML approach was used to calculate variance 297 components due to families ($V_{\rm AF}$) and individuals within families ($V_{\rm WF}$) separately within each 298 trial and population (i.e., variation within each population was divided into components due to 299 families, blocks, and residual variation). Variation within families may include a component

300	due to the gen	netic diversity	of pollen	donors sam	pled by r	nother trees.	SD's were also
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301 calculated for individual families (SD_{WF}).

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304 *Climate data*

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307 Long-term means at the sampled sites

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To investigate how the sampled sites varied in terms of temperature conditions, UK 309 Meteorological Office 40-year mean (1961-2000) climate data (Perry and Hollis 2005) were 310 used to create climatic profiles of the populations' origins. Data were extracted for the length 311 of the growing season (GSL), number of growing degree days (GDD), and mean February 312 and July temperatures (FMT and JMT; these represent on average, the coldest and warmest 313 months, respectively). GSL is defined as the period bounded by daily mean temperature above 314 5 °C for more than five consecutive days and daily mean temperature below 5 °C for more 315 than five consecutive days (after 1 July), while GDD expresses the sum of daily heat sum 316 accumulation above 5.5 °C. Exact details on how the climate data were generated are given in 317 Perry and Hollis (2005). Climate data are available in 5 km \times 5 km grids and are based on 318 319 interpolation of observations from the nearest weather stations. It is possible that due to 320 within-grid variation in the landscape, actual climate conditions experienced at our study sites 321 differ from the estimates, therefore, the climatic variables should only be considered as proxies. The range of altitude sampled at each site was used as a proxy for fine-scale (within-322 323 population) environmental variation.

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333 $km \times 5$ km grids in the period 1961-2000 were used. Using these 40-year data, mean absolute 334 deviations (*MAD*) of FT and JT, and coefficients of variation (*CV*) of GSL and GDD (CV_{GSL} , 335 CV_{GDD}) were calculated separately for each site. A combined estimate of monthly temperature 336 variability was calculated as the average of the MAD's of FT and JT. Annual means were also 337 calculated over all 21 sites. To test whether winter temperature in Scotland was associated with the North Atlantic Oscillation (NAO), linear regression was also used to test for an 338 association between annual FTs and February NAO indices provided by the Climate Analysis 339 340 Section, NCAR, Boulder, USA, at NAO Index 341 http://www.cgd.ucar.edu/cas/jhurrell/indices.html. Linear regression was used to explore 342 whether the level of temporal variation in climate varied along latitudinal, longitudinal, or 343 altitudinal gradients. 344 345 Associations between population means of timing of bud flush and climate at home site 346

To investigate patterns of temporal variability in temperature in Scotland and at the sampled

sites, annual estimates of GSL, GDD, February and July temperatures (FT and JT) for the 5

- 347
- 348 To investigate associations between variation in timing of bud flush, the locations of the
- 349 populations, and their climate, population means in the two trials were regressed against their
- 350 longitude, latitude, altitude, and long-term mean temperature estimates (GSL, GDD, FMT,
- 351 JMT) at origin.
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- Associations between the level of variation within populations and latitude, longitude, and
 altitude
- 356
- 357 Linear regression was used to explore whether the level of variation within populations

359 spatial gradients (longitude, latitude, altitude). We also tested for associations between the

360	locations of the sampled	families and an	nong and within-	family trait	variation ($(V_{\rm AF}, I)$	V _{WF} ,
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361 SD_{WF}) in each trial.

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- 364 **Results**
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367 Genetic differentiation in timing of bud flush among populations and families

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369 In the Edinburgh trial, population means ranged between 11 days after March 23 for AC, BB, 370 GA, GD, and GT, and 18 days after March 23 for AB, BE, SD, and RD. ANOVA provided 371 some evidence of differences among populations (*P*=0.058), and differences among families 372 within populations, and among blocks were significant (Table 2a). The variance component 373 due to differences among families within populations (15.39; 22% of total variation) was 374 approximately five times larger than that among populations (2.98; 4% of total variation).

375

In the Aberdeen trial, the range of population means was from 16 days after March 31 for AC 376 and GL to 22 days after March 31 for AB. Significant differences were observed among 377 populations, families within populations, and blocks (Table 2b). The variance component due 378 379 to families (5.88; 11% of total variation) was approximately four times larger than that of 380 populations (1.45; 3%) of total variation). Population means between the two trials were 381 significantly and moderately correlated (r=0.48, P<0.05). In both trials, the great majority of 382 the variation was residual, i.e. within families (Table 2a, b; 72% in the Edinburgh trial, 77% in 383 the Aberdeen trial).

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390 Temperature conditions varied greatly within Scotland: for instance, mean GSL ranged from 391 116 days in BB to 283 days in BE, and mean GDD from 446 to 1,329 dd at the same sites (Table 1). Spatial variation was also found in timing of bud flush, and results from the two 392 393 trials showed similar trends although there were differences in the absolute values. In the Edinburgh trial, when examining associations between population means and geographical 394 395 surrogates of environmental variation (latitude, longitude, altitude), means were best 396 associated with altitude at their site of origin. Low-altitude populations generally flushed later 397 than those from higher locations, and altitude explained 20% of the variation among 398 population means (β_0 =16.70, β_1 =-0.010, P<0.05). Altitude of the populations was negatively correlated with mean GSL (r=-0.80, P<0.001) and GDD (r=-0.65, P<0.01) at origin, and 399 higher R^2 's were obtained when using these climate variables instead of altitude. Earlier bud 400 flush occurred in populations from areas with shorter GSL ($\beta_0=5.87$, $\beta_1=0.038$, P<0.01, 401 R^2 =31%), fewer GDD (Figure 2; β_0 =8.14, β_1 =0.0066, P<0.01, R^2 =35%), lower FMT 402 $(\beta_0=12.62, \beta_1=1.13, P<0.01, R^2=29\%)$, and lower JMT $(\beta_0=0.35, \beta_1=1.13, P<0.010, R^2=27\%)$. 403 404

A similar trend with altitude was also found in the Aberdeen trial ($\beta_0=20.25$, $\beta_1=-0.0035$), but the association was not statistically significant (P=0.22, $R^2=3\%$). However, significant associations were obtained when temperature estimates were used instead of altitude, and sites with shorter GSL ($\beta_0=14.14$, $\beta_1=0.024$, P<0.001, $R^2=31\%$), fewer GDD (Figure 2; $\beta_0=15.70$, $\beta_1=0.0040$, P<0.01, $R^2=32\%$), lower FMT ($\beta_0=18.38$, $\beta_1=0.69$, P<0.01, $R^2=28\%$), and lower JMT ($\beta_0=9.68$, $\beta_1=0.79$, P<0.01, $R^2=35\%$) had earlier bud flush.

411

413 Differences in the level of variation within populations

414

415 Variation among SD_{POP}'s suggested that populations might have differed in the level of

416 variation in timing of bud flush (Table 3). In the Edinburgh trial, SD_{POP} 's varied between 6.25 417 in GT and 9.53 in AB, while in the Aberdeen trial with a larger number of families within 418 each population, SD_{POP}'s ranged from 4.29 in CG to 11.10 in GD. SD_{POP}'s across the two 419 trials were not significantly correlated (P=0.79). In the Edinburgh trial, SD_{POP} 's were not 420 associated with latitude, longitude, or altitude. However, in the Aberdeen trial, the pattern of 421 variation among SD_{POP} 's was related to the geographic location of populations and individual 422 mother trees: higher amounts of variation were observed at higher altitude sites (Figure 3; $\beta_0=5.14, \beta_1=0.0080, P<0.01, R^2=35\%$). The regression was strongly influenced by the three 423 high-altitude sites with large SD_{POP} 's. When excluding these, the linear regression remained 424 positive but became statistically non-significant ($\beta_0=5.59$, $\beta_1=0.0050$, P=0.112, $R^2=10\%$). 425

426

In the Aberdeen trial, differences among families were generally larger at higher altitudes, and 427 altitude explained 16% of the variation among V_{AF} 's (β_0 =-0.65, β_1 =0.024, P<0.05); however, 428 429 eight populations had V_{AF} estimates of 0. A positive correlation was observed between V_{AF} 's and the range of family means within each population (r=0.77, P<0.001). R^2 increased to 31% 430 when the range of family means within each population was used instead of V_{AF} (β_0 =6.36, 431 $\beta_1=0.019$, P<0.01). In the Edinburgh trial, higher SD_{POP}'s were associated with higher V_{AF}'s 432 (r=0.69, P<0.001). The association between altitude and V_{AF} 's was only suggestive of higher 433 levels of among-family variation at higher altitudes (β_0 =6.074, β_1 =0.040, P=0.11, R²=9%) 434

435

Similarly to among-family differences, there was some evidence of larger V_{WF} 's at higher 436 altitude sites in the Aberdeen trial ($\beta_0 = 27.39$, $\beta_1 = 0.070$, $R^2 = 13\%$, P = 0.062). This pattern was 437 also reflected in variation among SD_{WF}'s which ranged between 0 in nine families from seven 438 439 populations and 17.92 in a family from AC. Altitude explained 4% of the variation among SD_{WF} s (β_0 4.76, β_1 =0.0053, P<0.01), but the association was non-significant (P=0.53) when 440 the three highest-altitude sites were excluded. In the Edinburgh trial, there was no association 441 442 between altitude and $V_{\rm WF}$'s, but they were positively and significantly correlated with $SD_{\rm POP}$'s 443 (r=0.78, P<0.001). SD_{WF}'s varied between 4.16 in a family from GT and 11.04 in a family 444 from CR, but they were not associated with altitude (P=0.48).

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- 446

450 Mother trees within populations were sampled at different altitudes, and consequently the

451 altitudinal range sampled at each site varied from 23 m at RM to 179 m at GC in the

452 Edinburgh trial, and from 34 at RM m to 199 at GC m in the Aberdeen trial. However,

453 increasing altitudinal range sampled within populations did not account for larger SD_{POP} 's

454 (the Edinburgh trial: $\beta_0 = 8.42$, $\beta_1 = -0.0060$, $R^2 = 4\%$, P = 0.20; the Aberdeen trial: $\beta_0 = 7.010$,

455 $\beta_1 = 0.001, R^2 = 0\%, P = 0.95$).

456

457

458 Temporal climate variation

459

Climate differed markedly from year to year. For example, annual GSL and GDD of the sites 460 occupied by the 21 pinewoods showed extensive temporal fluctuation in the period 1961-2000 461 (Figure 4a). GSL varied between 174 days in 1968 and 271 days in 1989, while the lowest 462 GDD (756 dd) was reached in 1974 and the highest (1,167 dd) in 1995. Temporal variability 463 was also found in monthly winter and summer temperatures. The range of JTs was 9.90 °C in 464 1965 and 14.88 °C in 1983, and annual JTs were significantly correlated with GSL (r=0.46, 465 466 P < 0.01) and GDD (r = 0.64, P < 0.001) in the same year. FTs varied between -2.38 °C in 1963 and 5.94 °C in 1998, and were found to be associated with the NAO, with colder temperatures 467 468 coinciding with lower NAO indices (Figure 4b; $\beta_0=1.29$, $\beta_1=0.58$, P<0.0001, R²=33%). 469

470 Populations from different parts of Scotland experienced different levels of temporal variation 471 in these climate features. The combined *MAD* of FT and JT increased with ascending altitudes 472 (Figure 4c; β_0 = 1.032, β_1 =0.00072, *P*<0.0001, R^2 =77%), while for GSL and GDD, temporal 473 variability increased very little from altitudes of 48 to 343 m, but was higher at the three sites 474 located above 450 m (Figure 4d).

475

477 Discussion

478

479 In this study, we combined phenotypic and climate data to examine the patterns of variation in 480 a phenological trait among and within native Scottish populations of Scots pine. Under 481 common-garden conditions, populations sampled across a spatially highly heterogeneous 482 landscape were found to differ in timing of bud flush at the beginning of the second growing, 483 season which generally was earlier in populations from cooler locations. This suggests 484 environment-driven genetic differentiation. However, significant amounts of variation were 485 also found within populations. In addition, the data suggested that populations may differ in 486 their level of adaptive variation: in the Aberdeen trial, we found some evidence of higher 487 levels of such variation in populations from high-altitude sites that experience the most 488 among-year variation in temperature conditions.

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490

491 Populations are differentiated in timing of bud flush

492

The annual cycle of temperate trees is divided into two phases: active growing period in 493 494 summer and winter dormancy (Howe et al. 2003). Due to differences in the length of the 495 frost-free period in the Northern hemisphere, phenological differences are common among 496 tree populations (Howe et al. 2003; Savolainen et al. 2007). However, such patterns have 497 mainly been examined across wide geographic areas, and less is known about genetic 498 differences among populations separated by shorter distances. Despite the small geographic 499 area (maximum distance between two native pinewoods is less than 200 km), spatial 500 heterogeneity in climate within Scotland is extensive (Salmela et al. 2010). Thus, conditions 501 are ideal for development of local adaptation. Indeed, population differences observed in our 502 study suggest that adaptive differentiation in response to environmental variation has 503 occurred. Under glasshouse conditions, bud flush generally took place earlier in populations 504 from the coolest high-altitude locations in the eastern Highlands and later in those from the 505 maritime west coast. However, possible home site advantage of these populations cannot be

506 inferred without reciprocal transplant experiments (Kawecki and Ebert 2004). Also note that 507 due to the format of the partially interpolated climate data ($5 \text{ km} \times 5 \text{ km}$ grids) and spatially 508 complex landscapes in Scotland, it is possible that the home site conditions of the populations 509 differ from those described in Table 1. Weather station coverage in the UK is especially sparse 510 in the Scottish Highlands which is likely to result in inaccuracies in the climate variables 511 (Perry and Holliss 2005).

512

In spring, growth is initiated from stem units formed in buds during the previous growing 513 514 season after genetically-determined chilling and heat sum requirements have been fulfilled 515 (Aitken and Hannerz 2001; Howe et al. 2003). The patterns observed in our study could reflect longer chilling and higher heat sum requirements of populations from warmer 516 517 locations so that growth initiation is prevented under mild winter conditions (e.g. Leinonen 518 1996). Due to the strong dependence of the trait on temperature, these patterns of variation 519 may differ among years (e.g. Sagnard et al. 2002, but see Beuker 1994). However, corresponding spatial patterns of variation in this trait have been found in provenance studies 520 on adult trees of the same species sampled across Eurasia (Steiner 1979) and along a 521 latitudinal gradient in North Europe (Beuker 1994). Also, in an outdoor trial consisting of a 522 523 small subset of seedlings from eight populations included in the Edinburgh trial, timing of bud 524 flush at the beginning of the fourth growing season was slightly earlier in populations from cooler high-altitude locations (Salmela et al. 2011). Nonetheless, in accordance with earlier 525 526 findings also in other species (Aitken and Hannerz 2001), population differences in growth initiation appeared small. Although similar overall trends were found in the two trials, there 527 528 were differences in absolute values. This may be due to differences in winter and spring temperatures between the experimental sites. 529

530

531 It is possible that phenotypic variation is influenced not only by genetic variation due to 532 segregating genes among seedlings, but also by differences in seed maturation conditions 533 experienced by different mother trees in their home environments. These effects have been 534 shown to be strong for instance in *Picea abies* (L.) H.Karst (Johnsen et al. 2005). Although 535 we cannot exclude the possibility of such effects influencing the observed patterns of 536 variation also among Scottish pine populations, earlier studies suggest that in Scots pine, such 537 effects are not of the same magnitude as in *P. abies*. For example, Ruotsalainen et al. (1995)

538 found that seed maturing conditions did not have major effect on variation in another

539 phenological trait, timing of bud set, under common-garden conditions. Further, in addition to

540 young seedlings, evidence of adaptive differentiation among populations in timing of bud

541 flush has been observed when examining Scots pine trees aged over 10 (Steiner 1979) or

542 approximately 60 years (Beuker 1994). Thus, in further discussion we assume that the

543 differences observed in this study reflect mainly genetic variation among seedlings.

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546 Populations consist of genetically and phenotypically diverse individuals

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548 Despite the evidence of population differentiation, a much larger proportion of variation was 549 due to differences among families and individual seedlings within populations. This observation is akin for instance to the one found by Sagnard et al. (2002) among Abies alba 550 Mill. populations from the south-western Alps and indicates that the trait is heritable and that 551 552 populations maintain considerable internal potential to adapt to changing conditions (e.g. Kramer and Havens 2009; Hoffman and Sgrò 2011), such as variable growing season length. 553 Furthermore, our data suggested that populations differed in levels of internal variation: 554 555 family differences were larger and families more variable at high-altitude locations. This pattern was found only in the Aberdeen trial which may be due to the larger number of 556 557 families sampled within each population. Evolutionary biology models predict the loss of 558 genetic variation in adaptive traits due to selection favouring only optimal phenotypes in each 559 population (Falconer and Mackay 1996), but significant levels of within-population variation 560 are often documented in adaptive traits that are differentiated among populations (Howe et al. 561 2003, Alberto et al. 2011; Savolainen et al. 2011). Still, population differences in the amount of variation have been assessed only on few occasions (Savolainen et al. 2004; Notivol et al. 562 563 2007; Alberto et al. 2011).

564

565 Several factors could contribute to the population differences in the level of trait diversity 566 along an altitudinal gradient observed in our study. Although the areas covered by the 21 567 pinewoods vary in size (Mason et al. 2004), differences in population size are an unlikely 568 explanation, as earlier work using selectively neutral molecular markers in the nuclear 569 genome has shown no significant differences in molecular diversity across Scotland (Kinloch 570 et al. 1986; Wachowiak et al. 2010). Gene flow among heterogeneous sites may increase 571 variation within populations (Howe et al. 2003) and at least historically, Scottish populations have been linked by gene flow (Kinloch et al. 1986). Whether population differences in the 572 573 extent of long-range gene flow contribute to the patterns observed here requires further 574 exploration. However, studies of pollen flow suggest that the great majority of the fertilising 575 pollen usually comes from local trees (Smouse and Sork 2004), suggesting that a large proportion of within-population diversity can also arise from matings between local parents. 576 577 Thus, in a common-garden study design sampling open-pollinated seed from natural stands, 578 gene flow from other populations might contribute more to variation within than among families because offspring from matings between parents located in different populations 579 might not be as well adapted to their home site as those with local parents and might not 580 581 become established in a population (Burczyk et al. 2004). The high levels of residual variation found in both our trials might have resulted from effective outcrossing, while population 582 583 differences in the level of variation within families may reflect variation in the extent of longdistance pollen flow and/or the genetic diversity of local pollen donors. However, the current 584 585 study design does not allow the separation of genetic effects from other possible causes of 586 residual variation.

587

Expression of phenotypic variation is strongly influenced by the environment and 588 589 consequently, artificial growing conditions in glasshouses may induce the expression of 590 variation that would normally be 'hidden' in nature (e.g. Hoffmann and Merilä 1999). 591 Although growth conditions are known to affect population means in trees (Oleksyn et al. 592 1998; Mimura and Aitken 2010), their effects on trait variances remain poorly characterised. 593 It is possible that seedlings have expressed different levels of their total potential if the 594 growing conditions in the glasshouses were more novel to some populations. This possibility 595 could be investigated further by experiments in additional growing environments. In addition 596 to differences in within-population sampling, discrepancies between the spatial patterns of 597 variation among SD_{POP} 's between the trials could be due to different growing environments or 598 scoring intervals.

600 Adaptive diversity may be increased in environments that are highly variable across space or 601 time (Yeaman and Jarvis 2006). Assuming that the altitudinal range sampled at each site also 602 reflects the level of fine-scale environmental variation within populations, there was no 603 evidence of increased spatial heterogeneity at high altitudes. Finer-scale climate data are 604 needed to explore how environmental conditions vary across short distances in complex 605 landscapes like the Scottish Highlands. However, there is significant temporal heterogeneity in climate in Scotland, and for instance mid-winter temperatures were found to be associated 606 607 with the NAO phenomenon which is known to influence a variety of biological events in both 608 plants and animals (Stenseth et al. 2002). The effect of the NAO in Europe is known to be 609 particularly strong in winter, but positive phases of the NAO during spring (February-April) have also been demonstrated to be associated with elevated temperatures and earlier 610 611 phenological events in plants (Chmielewski and Rötzer 2001). Such variation may also partly explain the observed temporal variation in the length of the growing seasons and the number 612 613 of growing degree days in Scotland.

614

We also found that the extent of temporal variation varied spatially within Scotland, and 615 higher-altitude locations were characterised by more variable climates. Interestingly, our 616 617 common-garden data suggest that genetic diversity at least in timing of bud flush may be higher in populations found at the environmentally most variable sites. The association of 618 temporal climate variability with altitude most likely arises from the fact that in our sampling 619 sites, the highest-altitude sites are located at the most continental sites in the eastern 620 621 Highlands, while the lowest-altitude sites are found on the maritime west coast (see Vasseur 622 and Yodzis 2004). Although the climate data were based on interpolation across a temporally 623 and spatially variable number of weather stations and the precision of estimates varies 624 depending on the variable being estimated (Perry and Hollis 2005), patterns of variation in 625 latewood density chronologies among five Scottish pinewoods provide indirect but 626 corresponding evidence of site differences in temporal variability at least in summer 627 temperatures (Hughes et al. 1984).

628

629 Temporal variation in temperatures across the whole year suggests that phenotypic optima in

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631 in trait optima within Scotland may be more pronounced at sites with more among-year variation in climate. Large-scale climate fluctuations are known to have shaped the distributions of many tree species (Westfall and Millar 2004) and among-year variation for instance in summer temperatures has been documented to decrease the likelihood of good seed years especially in harsh conditions at high latitudes (e.g. Hilli et al. 2008), but so far, the role of temporal variation in factors likely to drive adaptation in plant populations has received only little attention. The contribution of temporal heterogeneity in maintaining adaptive diversity has often been considered weak (Hedrick 1986), but on the other hand, variation in environmental factors has not been studied in such detail as variation in phenotype. The role of temporally variable environment might be important especially in long-lived trees with overlapping generations and low climate-related mortality in adults (see Persson and Ståhl 1990; Ellner and Hairston 1994). Trees aged over 400 years have been found in Scottish pinewoods (Fish et al. 2010), and considering the evidence for substantial temporal heterogeneity in the European climate since 1500 (Luterbacher et al. 2004) and the age structure of populations, it is possible that different age cohorts have experienced differing selection pressure during their sensitive early life stages and that the current patterns of diversity reflect adaptations to a range of past environments. Accordingly, a more variable environment might also support higher levels of genetic variation and more diverse phenotypes. This possibility could be explored further by examining genetic differences among age groups from sites that differ in temporal heterogeneity: larger differences among age groups in more variable environments would provide stronger evidence for a positive association between the levels of environmental and genetic diversity. Phenotypic plasticity is also expected to evolve in heterogenous environments (Valladares et al. 2007); whether differences in plasticity contribute to our observations could be investigated in more detail by 654 655 examining the range of phenotypic variation expressed across multiple growing environments.

phenological traits in populations vary from year to year. Furthermore, among-year variation

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662 environments

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Genetic variation in adaptive traits has important biological implications. When assumed to 664 have no capacity to evolve, changing environmental conditions may result in reduced survival 665 666 or extinction of current populations in some areas. When the internal capacity of populations to adapt is accounted for, the chance of survival is increased (Aitken et al. 2008). Thus, 667 natural populations inhabiting heterogeneous habitats should not be treated as fixed and 668 669 independent entities whose responses to changes will be determined solely by the environment. Also, studies on adaption in tree and other plant populations would benefit from 670 more thorough examinations of environmental factors likely to drive adaptation processes 671 672 both within and among populations. Despite the evidence of temporal heterogeneity being a 673 general feature of natural environments (Vasseur and Yodzis 2004), local climates in 674 evolutionary studies have generally been considered to be rather static and non-overlapping, and the potential role of environmental heterogeneity in maintaining genetic diversity has not 675 676 been extensively explored.

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In this study, we have shown that the environmental variability natural populations are 678 679 exposed to may differ even across short geographic distances and that this may influence the amount of adaptive diversity found within populations. Predicting the future of natural 680 populations is complicated by possible correlations between adaptive traits, complex effects 681 682 of the environment on the expression of phenotypes, and many non-genetic factors, but 683 clearly, the fluctuating behaviour of environmental factors and the ubiquitous finding of 684 adaptive potential at least in some key traits needs to be taken into account in further studies 685 which aim to predict the effects of global change on natural populations. More familystructured trials grown across a range of sites are also needed to characterise within-686 687 population variation and its causes in more detail. Thanks to the long-history of common-688 garden studies with appropriate study designs in forest trees, existing data from a large 689 number of completed studies can be used to test for similar patterns in other species.

694 The authors wish to thank Scottish Forestry Trust for funding (MJS' Ph.D. studentship), Dave 695 Sim, Joan Beaton, Sheila Reid, and Ben Moore (James Hutton Institute) for making the seed 696 collections and experimental assistance, NERC, the Forestry Commission, the Scottish 697 Government's Rural and Environment Science and Analytical Services Division (RESAS). 698 and EU-funded Network of Excellence EVOLTREE for support, UK Meteorological Office 699 for the climate data, and anonymous reviewers for constructive comments on the manuscript, 700 701 702 Notes on contributors 703 Matti Salmela is an evolutionary biologist interested in local adaptation, its quantitative 704 705 genetic basis, and maintenance of adaptive genetic variation in natural populations. 706 Stephen Cavers is a senior scientist at CEH, studying genetic diversity, gene flow, and 707 adaptation in plants, in both tropical and temperate ecosystems. 708 709 Joan Cottrell is head of the molecular team at Forest Research. Her research interests include 710 711 the assessment of genetic diversity, adaptation, and gene flow in native British trees and their 712 associated woodland dwelling flora and fauna. She is particularly interested in translating 713 research results into informed conservation advice on policy and management of our 714 woodland resource. 715 716 Glenn Iason is an ecologist with interests in herbivore nutritional ecology and ecosystem 717 management. He is also studying the extent of genetic and environmental variation in plant 718 secondary metabolites and how these determine plant-herbivore interactions, associated

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725	
726	References
727	
728	Aitken SN, Hannerz M. 2001. Genecology and gene resource management strategies for
729	conifer cold hardiness. In: Conifer Cold Hardiness. Dordrecht (Netherlands): Kluwer
730	Academic Publishers. p. 23-53.
731	
732	Aitken SN, Yeaman S, Holliday JA, Wang T, Curtis-McLane S. 2008. Adaptation, migration
733	or extirpation: climate change outcomes for tree populations. Evolutionary Applications 1:95-
734	111.
735	
736	Alberto F, Bouffier L, Louvet J-M, Lamy J-B, Delzon S, Kremer A. 2011. Adaptive responses
737	for seed and leaf phenology in natural populations of sessile oak along an altitudinal gradient.
738	Journal of Evolutionary Biology 24:1442-1454.
739	
740	Barton NH, Keightley PD. 2002. Understanding quantitative genetic variation. Nature
741	Reviews Genetics 3:11-21.
742	
743	Benito Garzón M, Alía R, Robson TM, Zavala MA. 2011. Intra-specific variability and
744	plasticity influence potential tree species distributions under climate change. Global Ecology
745	and Biogeography 20:766-778.
746	
747	Beuker E. 1994. Adaptation to climatic changes of the timing of bud burst in populations of

750	Billington HL, Pelham J. 1991. Genetic variation in the date of budburst in Scottish birch
751	populations: implications for climate change. Functional Ecology 5:403-409.
752	
753	Bull JJ. 1987. Evolution of phenotypic variance. Evolution 41:303-315.
754	
755	Burczyk J, DiFazio SP, Adams WT. 2004. Gene flow in forest trees: how far do genes really
756	travel? Forest Genetics 11:179-192.
757	
758	Campbell, R. K. (1979) Genecology of Douglas-fir in a watershed in the Oregon Cascades.
759	Ecology 60, 1036-1050.
760	NO
761	Chmielewski F-M, Rötzer T. 2001. Response of tree phenology to climate change across
762	Europe. Agricultural and Forest Meteorology 108:101-112.
763	
764	Chmura DJ, Anderson PD, Howe GT, Harrington CA, Halofsky JE, Peterson DL, Shaw DC,
765	St. Clair JB. 2011. Forest responses to climate change in the northwestern United States:
766	ecophysiological foundations for adaptive management. Forest Ecology and Management
767	261:1121-1142.
768	G
769	Critchfield WB, Little E. 1966. Geographic distribution of the pines of the world. U.S.
770	Department of Agriculture Forest Service Miscellaneous Publication, 991.
771	
772	Davis MB, Shaw RG. 2001. Range shifts and adaptive responses to Quaternary climate
773	change. Science 292:673-679.
774	

Pinus sylvestris L. and Picea abies (L.) Karst. Tree Physiology 14:961-970.

- Downloaded by [CEH Edinburgh] at 03:30 16 April 2013
- Elith J, Leathwick JR. 2009. Species distribution models: ecological explanation and
- prediction across space and time. Annual Review of Ecology, Evolution, and Systematics 40:

777 677-697.

- 778
- Ellner S, Hairston NG. Jr. 1994. Role of overlapping generations in maintaining genetic
- variation in a fluctuating environment. American Naturalist 143:403-417.
- 781
- 782 Eriksson G, Andersson S, Eiche V, Ifver J, Persson A. 1980. Severity index and transfer effects
- on survival and volume production of *Pinus sylvestris* in Northern Sweden. Studia Forestalia
 Suecica 156:1-32.

785

- 786 Falconer DS, Mackay TFC. 1996. Introduction to Quantitative Genetics. Second edition.
- 787 Essex (UK): Pearson Education Limited.

788

- 789 Fish T, Wilson R, Edwards C, Mills C, Crone A, Kirchefer AJ, Linderholm AW, Loader NJ,
- 790 Woodley E. 2010. Exploring for senescence signals in native Scots pine (Pinus sylvestris L.)
- in the Scottish Highlands. Forest Ecology and Management 260:321-330.

792

- 793 Garcia-Gonzalez F, Simmons LW, Tomkins JL, Kotiaho JS, Evans JP. 2012. Comparing
- rounding the calculation and use of coefficients of additive
- 795 genetic variation. Evolution 66:2341-2349.

- Harrison SJ. 1997. Changes in the Scottish climate. Botanical Journal of Scotland 49:287-300.
- 799 Hedrick PW. 1986. Genetic polymorphism in heterogeneous environments a decade later.
- 800 Annual Review of Ecology and Systematics 17:535-566.
- 801

802 Hendry AP, Kinnison MT, Heino M, Day T, Smith TB, Fitt G, Bergstrom CT, Oakeshott J,

803 Jørgensen PS, Zalucki MP et al. 2011. Evolutionary principles and their practical application.

804 Evolutionary Applications 4:159-183.

805

806 Hilli A, Hokkanen T, Hyvönen J, Sutinen M-L. 2008. Long-term variation in Scots pine seed

crop size and quality in northern Finland. Scandinavian Journal of Forest Research 23:395-403.

809

810 Hoffmann AA, Merilä J. 1999. Heritable variation and evolution under favourable and

811 unfavourable conditions. Trends in Ecology and Evolution 14:96-101.

812

813 Hoffmann AA, Sgrò CM. 2011. Climate change and evolutionary adaptation. Nature 470:

814 479-485.

815

816 Houle D.1992. Comparing evolvability and variability of quantitative traits. Genetics 130:817 195-204.

818

819 Howe GT, Aitken SN, Neale DB, Jermstad KD, Wheeler NC, Chen THH. 2003. From

820 genotype to phenotype: unraveling the complexities of cold adaptation in forest trees.

821 Canadian Journal of Botany 81:1247-1266.

822

823 Hughes MK, Schweingruber FH, Cartwright D, Kelly PM. 1984. July-August temperature at

824 Edinburgh between 1721 and 1975 from tree-ring density and width data. Nature 308:341-

826

825

344.

827 Johnsen Ø, Dæhlen OG, Østreng G, Skrøppa T. 2005. Daylength and temperature during seed

828 production interactively affect adaptive performance of *Picea abies* progenies. New

829 Phytologist 168:589-596.

850	
831 832	Kawecki TJ, Ebert D. 2004. Conceptual issues in local adaptation. Ecology Letters 7:1225- 1241.
833	
834	Kinloch BB. Westfall RD. Forrest GI. 1986. Caledonian Scots pine - origins and genetic
835	structure. New Phytologist 104:703-729.
836	
837	Kramer AT, Havens K. 2009. Plant conservation genetics in a changing world. Trends in Plant
838	Science 14:599-607.
839	
840	Leinonen I. 1996. Dependence of dormancy release on temperature in different origins of
841	Pinus sylvestris and Betula pendula seedlings. Scandinavian Journal of Forest Research 11:
842	122-128.
843	
844	Linhart YB, Grant MC. 1996. Evolutionary significance of local genetic differentiation in
845 846	plants. Annual Review of Ecology and Systematics 27:237-277.
847	Luterbacher J, Dietrich D, Xoplaki E, Grosjean M, Wanner H. 2004. European seasonal and
848	annual temperature variability, trends, and extremes since 1500. Science 303:1499-1503.
849	
850	Mason WL, Hampson A, Edwards C. 2004. Managing the Pinewoods of Scotland. Edinburgh:
851	Forestry Commission.
852	
853	Merilä J, Sheldon BC. 1999. Genetic architecture of fitness and nonfitness traits: empirical
854	patterns and development of ideas. Heredity 83:103-109.
855	
856	Mikola J. 1982. Bud-set phenology as an indicator of climatic adaptation of Scots pine in

857 Finland. Silva Fennica 16:178-184.

858

859 Milner JM, Elston DA, Albon SD. 1999. Estimating the contributions of population density

- 860 and climatic fluctuations to interannual variation in survival of Soay sheep. Journal of Animal
- 861 Ecology 68:1235-1247.

862

- 863 Mimura M, Aitken SN. 2010. Local adaptation at the range peripheries of Sitka spruce
- 864 Journal of Evolution Biology 23:249-258.

865

- 866 Notivol E, García-Gil MR, Alía R, Savolainen O. 2007. Genetic variation of growth rhythm in
- the limits of a latitudinal cline in Scots pine. Canadian Journal of Forest Research 37:540-551.

868

869 Oleksyn J, Tjoelker MG, Reich PB. 1998. Adaptation to changing environment in Scots pine
870 populations across a latitudinal gradient. Silva Fennica 32:129-140.

871

Parmesan C. 2006. Ecological and evolutionary responses to recent climate change. Annual
Review of Ecology, Evolution, and Systematics 37:637-669.

874

- 875 Perry M, Hollis D. 2005. The generation of monthly gridded datasets for a range of climatic
- variables over the UK. International Journal of Climatology 25:1041-1054.

877

Persson B, Ståhl EG. 1990. Survival and yield of *Pinus sylvestris* L. as related to provenance
transfer and spacing at high altitudes in northern Sweden. Scandinavian Journal of Forest
Research 5:381-395.

881

Petit RJ, Hampe A. 2006. Some evolutionary consequences of being a tree. Annual Review of
Ecology, Evolution, and Systematics 37:187-214.

884	
885	Ray D. 2008. Forestry Commission Research Note 101: Impacts of climate change on forestry
886	in Scotland - a synopsis of spatial modelling research. Forestry Commission Scotland.
887	
888	Rehfeldt GE, Tchebakova NM, Parfenova YI, Wykoff WR., Kuzmina NA, Milyutin LI. 2002.
889	Intraspecific responses to climate in <i>Pinus sylvestris</i> . Global Change Biology 8:912-929.
890	
891	Reich PB, Oleksyn J. 2008. Climate warming will reduce growth and survival of Scots pine
892	except in the far north. Ecology Letters 11:588-597.
893	
894	Rousi M, Heinonen J. 2007. Temperature sum accumulation effects on within-population
895	variation and long-term trends in date of bud burst of European white birch (Betula pendula).
896	Tree Physiology 27:1019-1025.
897	
898	Ruotsalainen S, Nikkanen T, Haapanen M. 1995. Effect of seed-maturing conditions on the
899	growth and hardiness of one-year old Pinus sylvestris seedlings. Forest Genetics 2:189-198.
900	
901	Sagnard F, Barberot C, Fady B. 2002. Structure of genetic diversity in Abies alba Mill. from
902	southwestern Alps: multivariate analysis of adaptive and non-adaptive traits for conservation
903	in France. Forest Ecology and Management 157:175-189.
904	
905	Salmela MJ, Cavers S, Cottrell JE, Iason GR, Ennos RA. 2011. Seasonal patterns of
906	photochemical capacity and spring phenology reveal genetic differentiation among native
907	Scots pine (Pinus sylvestris L.) populations in Scotland. Forest Ecology and Management
908	262:1020-1029.
909	$\overline{\mathbf{v}}$
910	Salmela MJ, Cavers S, Cottrell JE, Wachowiak W, Iason GR, Ennos RA. 2010. Understanding
911	the evolution of native pinewoods in Scotland will benefit their future management and

Downloaded by [CEH Edinburgh] at 03:30 16 April 2013

912 conservation. Forestry 83:535-545.

913

- 914 Savolainen O, Bokma F, García-Gil R, Komulainen P, Repo T. 2004. Genetic variation in
- 915 cessation of growth and frost hardiness and consequences for adaptation of Pinus sylvestris to
- 916 climatic changes. Forest Ecology and Management 197:79-89.
- 917
- 918 Savolainen O, Kujala ST, Sokol C, Pyhäjärvi T, Avia K, Knürr T, Kärkkäinen K, Hicks S
- 919 2011. Adaptive potential of northermost tree populations to climate change with emphasis on
- 920 Scots pine (Pinus sylvestris L.). Journal of Heredity 102:526-536.

921

- 922 Savolainen O, Pyhäjärvi T, Knürr T. 2007. Gene flow and local adaptation in trees. Annual
- 923 Review of Ecology, Evolution, and Systematics 38:595-619

924

Smouse PE, Sork VL. 2004. Measuring pollen flow in forest trees: an exposition of alternative
approaches. Forest Ecology and Management 197:21-38.

927

Steiner KC. 1979. Patterns of variation in bud-burst timing among populations in several *Pinus* species. Silvae Genetica 28:185-194.

930

- 931 Stenseth NC, Mysterud A, Ottersen G, Hurrell JW, Chan K-S, Lima M. 2002. Ecological
- 932 effects of climate fluctuations. Science 297:1292-1296.

933

Valladares F, Gianoli E, Gómez JM. 2007. Ecological limits to plant phenotypic plasticity.
New Phytologist 176:749-763.

936

937 Vasseur DA, Yodzis P. 2004. The color of environmental noise. Ecology 85:1146-1152.

939	Wachowiak W, Salmela MJ, Ennos RA, Iason G, Cavers S. 2010. High genetic diversity at the
940	extreme range edge: nucleotide variation at nuclear loci in Scots pine (Pinus sylvestris L.) in
941	Scotland. Heredity 106:775-787.
942	
943	Westfall RD, Millar CI. 2004. Genetic consequences of forest population dynamics influenced
944	by historic climatic variability in the western USA. Forest Ecology and Management
945	197:159-170.
946	
947	White TL, Adams WT, Neale DB. 2007. Forest Genetics. Wallingford (UK): CABI
948	Publishing.
949	
950	Willi Y, Van Buskirk J, Hoffman AA. 2006. Limits to adaptive potential of small populations.
951	Annual Review of Ecology, Evolution, and Systematics 37:433-438.
952	
953	Yeaman S, Jarvis A. 2006. Regional heterogeneity and gene flow maintain variance in a
954	quantitative trait within populations of lodgepole pine. Proceedings of the Royal Society B:
955	Biological Sciences 273:1587-1593.
956	
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968 Table 1 Populations of *Pinus sylvestris* in Scotland included in the study, their latitude (Lat.),

969 longitude (Long.), altitudinal range sampled (Alt.), core pinewood area according to Mason et

970 al. (2004), and mean (1961-2000) calculated climate features: growing season length (GSL;

971 days), growing degree days (GDD: day degrees), and February and July mean temperatures

972 (FMT and JMT).

Population	Lat.	Long.	Alt. (m)	Area (ha)	GSL	GDD	FMT (°C)	JMT (°C)
Abernethy (AB)	57.21	3.61	311-370	2452	211	990	1.15	12.73
Allt Cul (AC)	57.04	3.35	435-512	13	145	513	-1.01	10.41
Amat (AM)	57.87	4.60	39-201	181	214	892	1.22	12.29
Ballochbuie (BB)	56.98	3.30	421-531	775	116	446	-1.69	9.46
Beinn Eighe BE)	57.63	5.40	17-91	182	283	1329	3.68	14.16
Black Wood of Rannoch (BW)	56.68	4.37	250-321	1011	254	1138	2.12	13.55
Coille Coire Chuilc (CCC)	56.42	4.71	222-311	67	226	928	1.64	12.32
Conaglen (CG)	56.79	5.33	89-193	189	246	887	2.20	11.73
Crannach (CR)	56.58	4.68	258-338	70	231	1019	1.81	12.62
Glen Affric (GA)	57.26	4.92	205-293	1532	210	769	0.88	11.62
Glen Cannich (GC)	57.35	4.95	182-381	301	212	778	0.96	11.71
Glen Derry (GD)	57.03	3.58	426-493	235	168	593	-0.46	11.34
Glen Einig (GE)	57.96	4.76	45-92	27	242	1089	2.19	13.15
Glen Loy (GL)	56.91	5.13	136-219	74	191	541	0.49	9.80
Glen Tanar (GT)	57.02	2.86	289-422	1564	235	1105	2.21	13.63
Loch Clair (LC)	57.56	5.36	98-166	126	277	1253	3.44	13.68
Meggernie (MG)	56.58	4.35	254-385	277	223	916	1.07	12.04
Rhidorroch (RD)	57.89	4.98	138-220	103	221	840	1.51	11.62
Rothiemurchus (RM)	57.15	3.77	295-329	1744	224	1087	1.39	13.15
Shieldaig (SD)	57.50	5.63	44-132	103	273	1093	3.21	12.83
Strath Oykel (SO)	57.98	4.61	35-160	14	257	1276	2.69	14.05

- 981 Table 2 Variation in timing of bud flush. Results of the nested ANOVA testing the effects of
- 982 population (fixed factor), families within populations (random factor), and blocks (random
- 983 factor) in the a) Edinburgh and b) Aberdeen trials.

- 984
- a) Edinburgh trial

a) Edinburgh trial					×
Source of variation	df	MS	<i>F</i> -ratio	P-value	Variance component
Population	20	1153.35	1.70	0.058	2.98
Families within populations	63	679.52	13.50	<0.001	15.39
Block	39	176.66	3.51	<0.001	1.50
Residual	3120	50.35			50.4
b) Aberdeen trial					CC.
Source of variation	df	MS	<i>F</i> -ratio	P-value	Variance component
Population	20	183.26	2.18	<0.01	1.45
Families within populations	188	83.96	2.01	<0.001	5.88
Block	39	221.37	5.29	<0.001	5.09
Residual	1216	41.87			42.26

- 986 Table 3 The amount of within-population variation in timing of bud flush. Standard deviations
- 987 (SD_{POP}) , variance components due to families (V_{AF}) , residual variation (V_{WF}) , and blocks
- (V_{Block}) , and the range among family means in each population within the two trials. 988

	Deputation			Edinburgh					Aberdeen		
	Population	SD _{POP}	V _{AF}	Mean range	$V_{\rm WF}$	V_{Block}	SD_{POP}	V_{AF}	Mean range	$V_{\rm WF}$	$V_{\rm Block}$
	Abernethy (AB)	9.53	43.95	15.63	70.12	0.00	8.48	0.00	14.09	76.92	0.00
	Allt Cul (AC)	7.76	8.90	6.20	57.18	0.00	8.98	2.39	12.19	62.75	16.32
	Amat (AM)	7.32	15.41	8.90	40.15	1.80	8.68	3.43	12.09	70.03	2.11
	Ballochbuie (BB)	8.86	27.93	12.38	62.28	0.00	8.30	0.00	10.30	52.46	22.99
	Beinn Eighe (BE)	7.62	9.56	6.72	48.00	2.84	6.29	0.00	6.09	34.88	6.64
	Black Wood of Rannoch (BW)	8.00	11.79	8.24	55.55	0.00	7.92	0.00	9.62	27.07	35.22
	Coille Coire Chuilc (CCC)	7.25	6.26	5.73	42.20	5.65	7.92	13.16	12.15	43.73	6.08
	Conaglen (CG)	7.97	4.56	5.69	54.22	5.81	4.29	0.01	7.08	20.36	0.00
	Crannach (CR)	9.19	55.18	16.34	51.34	0.80	5.52	11.40	10.02	23.81	0.00
	Glen Affric (GA)	6.79	5.14	4.78	43.25	0.00	5.29	0.00	3.15	36.67	0.00
	Glen Cannich (GC)	8.17	23.58	12.02	45.50	3.70	6.18	12.00	14.13	24.92	3.02
	Glen Derry (GD)	7.95	30.24	11.85	39.48	1.07	11.10	29.05	24.03	94.57	5.24
	Glen Einig (GE)	7.93	2.95	5.06	66.41	0.00	5.81	3.79	8.79	19.06	10.44
	Glen Loy (GL)	6.78	7.31	6.75	37.61	2.75	6.71	7.03	10.19	41.43	0.00
	Glen Tanar (GT)	6.25	8.09	6.22	32.43	0.44	8.17	11.87	16.91	53.83	2.84
	Loch Clair (LC)	8.51	0.00	1.98	64.00	9.16	7.76	0.00	11.99	76.21	0.00
	Meggernie (MG)	7.10	0.72	3.48	47.31	2.61	8.51	4.76	13.92	71.13	0.00
	Rhidorroch (RD)	8.10	8.00	6.53	58.24	0.41	5.42	2.88	12.46	26.72	0.06
	Rothiemurchus (RM)	9.02	22.73	10.55	61.92	2.35	6.17	11.53	12.84	18.53	6.97
	Shieldaig (SD)	8.49	26.01	10.84	53.52	0.00	5.42	0.00	6.28	37.54	0.00
89	Strath Oykel (SO)	7.27	14.68	6.94	42.00	0.00	5.55	0.00	6.61	27.00	4.22
				5							

992 Figure 1 Map of the sampled native Pinus sylvestris populations and locations of the two trial

- 993 sites. Climatic features of the sites are given in Table 1. Inset: full distribution of *P. sylvestris*,
- 994 with study area highlighted by box.
- 995
- 996 Figure 2 Relationship between mean growing degree days (GDD) at origin and population
- 997 means of timing of bud flush in the two trials. In the Edinburgh trial: $\beta_0 = 8.14$, $\beta_1 = 0.0066$,
- 998 $P < 0.01, R^2 = 35\%$; in the Aberdeen trial: $\beta_0 = 15.70, \beta_1 = 0.0040, P < 0.001, R^2 = 32\%$. Error bars
- 999 indicate standard errors of the means.

1000

- 1001 Figure 3 Relationship between altitude at origin and the amount of variation (SD_{POP}) in timing
- 1002 of bud flush within populations in the Aberdeen trial ($\beta_0=5.14$, $\beta_1=0.0080$, P<0.01, $R^2=35\%$).

1003

Figure 4 Among-year variation in the Scottish climate. a) Temporal variation in growing 1004 season length (GSL) and growing degree days (GDD); b) temporal variation in February 1005 temperature (FT) and February North Atlantic Oscillation (NAO) indices; c) relationship 1006 1007 between the altitudes of the 21 native pinewood sites and among-year variability of winter 1008 and summer temperatures (combined MAD of FT and JT; d) temporal variation in GSL and 1009 GDD (coefficients of variation, CV) plotted against the altitudes of the 21 sampled sites. The 1010 climate data used cover the period 1961-2000. In a) and b), annual means were calculated 1011 over the 5 km \times 5 km grids within which the 21 pinewood sites are located. 1012

1013 1014 Figure 1. 1015

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