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1 Title page

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4

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16

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18 discussion)

19 Abstract

20 Global warming threatens the productivity of forest plantations. We propose here
21 the integration of environmental information into a genomic evaluation scheme
22 using individual reaction norms, to enable the quantification of resilience in forest
23 tree improvement and conservation strategies in the coming decades. Random
24 regression models were used to fit wood ring series, reflecting the longitudinal
25 phenotypic plasticity of tree growth, according to various environmental gradients.
26 The prediction accuracy of the models was considered to select the most relevant
27 environmental gradient, namely a gradient derived from an ecophysiological model
28 and combining trunk water potential and temperature. Even if the individual
29 ranking was preserved over most of the environmental gradient, strong genotype
30 x environment interactions were detected in the extreme unfavorable part of the
31 gradient, which includes environmental conditions that are very likely to be more
32 frequent in the future. Combining genomic information and longitudinal data
33 allowed to predict the growth of individuals in environments where they have not
34 been observed. Phenotyping of 50% of the individuals in all the environments
35 studied allowed to predict the growth of the remaining 50% of individuals in all
36 these environments with an accuracy of 0.62. By adding observations in a reduced
37 number of environments for the individuals to be predicted, while decreasing the
38 number of phenotyped individuals across all environments, the prediction accuracy
39 reached 1.37, highlighting the importance of phenotypic data allocation. Genomic

40 reaction norms are useful for the characterization and prediction of the function of
41 genetic parameters and facilitate breeding in a climate change context.

42

43 Introduction

44 Forest trees are keystone species in forest ecosystems supporting biological
45 diversity and providing ecosystem services (Brockerhoff *et al.*, 2017). They also
46 produce wood, which will be a key material for meeting the challenges of the near
47 future, thanks to its multiple uses (construction, paper, furniture, energy,
48 chemistry) and its ability to sequester carbon for long periods of time
49 (Ramachandran Nair *et al.*, 2009; Domke *et al.*, 2020). In this context, forest
50 plantation has been expanding for several decades (FAO, 2010), with the aim of
51 concentrating timber production and relieve pressure on natural forest. However,
52 these benefits of forest plantation will require the adaptation of forest to a new,
53 more challenging climate (Allen *et al.*, 2010; Pawson *et al.*, 2013; Payn *et al.*, 2015)
54 One of the major levers for ensuring sustainable wood productivity for forest
55 plantations will be the deployment of trees capable of maintaining high growth
56 rates even in extreme environments. To meet this goal, the integration of
57 phenotypic plasticity, which is defined as the ability of an individual to produce
58 different phenotypes in different environmental conditions (Bradshaw, 1965), is
59 becoming a major issue in forest tree breeding programs (Ray *et al.*, 2022). An
60 individual is considered here as a unique genetic combination found in a single tree,
61 or in several vegetative copies genetically identical. The challenges posed by
62 climate change faced limited scope of traditional genetic analyses of forest trees
63 focusing principally on phenotypic plasticity between experimental sites (Baltunis
64 *et al.*, 2010; Correia *et al.*, 2010; Shalizi and Isik, 2019). Although these studies

65 highlight the existence of genotype x environment (GxE) interactions for conifer
66 trees, i.e. differential variations in plasticity between individuals, they often
67 consider a limited number of environments, selected so as to avoid high mortality
68 rates. They are, therefore, not designed to be representative of the full range of
69 environments of relevance in a context of rapid climate change. The cost and
70 difficulty of exposing the same individuals to different environmental conditions,
71 particularly for species difficult to propagate vegetatively, are major obstacles to
72 the systematic evaluation of across-site plasticity in the context of tree breeding.

73 Phenotypic plasticity can be effectively modeled by reaction norms if repeated
74 measurements across ages or clones are available, together with a relevant
75 descriptor of the environment in which the phenotype was expressed (Schlichting
76 and Pigliucci, 1998; Sanchez *et al.*, 2013). A reaction norm is a representation of
77 phenotypic values as a function of an environmental gradient. Various methods for
78 constructing reaction norms have been developed, but the random regression
79 model described by (Kirkpatrick and Heckman, 1989) is particularly relevant in
80 breeding contexts. Through the integration of genetic data, this model can
81 continuously estimate genetic parameters and breeding values according to the
82 gradient. The gradient most frequently chosen is time (age), and this approach is
83 frequently used in animal breeding (Jamrozik *et al.*, 1997; Schaeffer, 2004; Boligon
84 *et al.*, 2012) and more rarely in plant breeding contexts (Sun *et al.*, 2017; Campbell
85 *et al.*, 2018) including tree breeding (Apiolaza and Garrick, 2001; Wang *et al.*, 2009).
86 However, reaction norms can also be modelled along an environmental gradient
87 (Ravagnolo *et al.*, 2000, Zumbach *et al.*, 2008, Sanchez *et al.*, 2009). In forest

88 breeding, the more recent modelling of this type of reaction norm appears to be a
89 way to meet the challenges of rapid climate change in tree breeding (Marchal *et*
90 *al.*, 2019; Alves *et al.*, 2020).

91 Selection objectives for forest tree breeding have focused mainly on the final
92 volume of the tree trunk. Historical selection criteria evaluate the size of trees at
93 an advanced age (Mullin *et al.*, 2011; Pâques, 2013). . The continuous growth of
94 trees and their reactions to the different environments encountered over the years
95 are thus summarized by a very integrative measurement. It is not therefore possible
96 to trace back and identify the environmental factors contributing to the final
97 phenotype, as environment can be considered only in a global manner over the
98 whole period. However, yearly growth increments can be correlated with well-
99 characterized environments (Martinez-Meier *et al.*, 2008; Zas *et al.*, 2020). This can
100 be achieved with the use of wood ring series, which define the annual radial growth
101 of each individual in temperate climates. Indeed, the cambial activity of trees
102 depends strongly on environmental conditions, particularly temperature and water
103 availability (Schweingruber, 2007). The variability of annual ring width and wood
104 density characterizes the plastic response of trees to changing environmental
105 conditions. It has been shown to have genetic determinism (Sánchez-Vargas *et al.*,
106 2007; Dalla-Salda *et al.*, 2009) and could be used as a proxy for the potential
107 reaction of trees to changes in environmental conditions. The analysis of these
108 repeated phenotypes therefore provides an ideal longitudinal dataset for studying
109 phenotypic plasticity at individual level (Marchal *et al.*, 2019). Such analyses can be
110 explanatory in nature, seeking to identify the optimal combination of

111 environmental factors making a significant contribution to annual growth, but they
112 can also be predictive, with the development of functional models for inferring
113 growth in environments where individuals have not been observed.

114 The integration of molecular markers into genetic evaluations provides not only
115 more accurate estimates of genetic parameters, but also opportunities to
116 implement genomic selection (GS) approaches (R2D2 Consortium *et al.*, 2021). In
117 forest trees, such approaches pave the way for the early selection of important
118 traits, such as wood traits, that would otherwise be measured only after many years
119 of cumulative growth. GS is also particularly valuable in tree breeding, as it allows
120 the integration of traits that are costly and complex to measure (Grattapaglia and
121 Resende, 2011). In many species, the gains provided by the use of genomic data
122 have tended to eclipse the interest in longitudinal data (Oliveira *et al.*, 2019).
123 However, these two approaches are not antagonistic and their beneficial effects
124 can be combined (Rutkoski *et al.*, 2016; Sun *et al.*, 2017). Genomic reaction norms
125 based on environmental measurements are rarely used (Ly *et al.*, 2018), but are
126 potentially of great value in this context, as they allow prediction of growth in as
127 yet environments where individuals have not been observed, thus decreasing the
128 complex and costly evaluation procedures associated with experimentation and
129 phenotyping under different environmental conditions.

130 We propose here an integration of environmental information into genetic
131 evaluations, using reaction norms in the context of forest tree breeding. A random
132 regression model based on annual ring growth data for maritime pine (*Pinus*

133 *pinaster* Ait.) and including genomic data was used to fit individual-level reaction
134 norms. The genetic components of these norms were described and the
135 implications of their use in the context of breeding were further investigated with
136 respect to a classical analysis targeting final radial growth. Finally, we investigated
137 the model's ability to predict the growth of individuals in environments where they
138 have not been observed, considering realistic phenotyping conditions for the
139 maritime pine breeding program in a GS context. To our knowledge, this is the first
140 study in a tree breeding context to use a random regression model to combine
141 environmental gradient and genomic information.

142 Materials and Methods

143 Plant material

144 A maritime pine trial was established at two sites in 1997: Site 1 (Cestas, France:
145 Lat 44.74, Lng -0.68) and Site 2 (Escource, France: Lat 44.16, Lng -1.03). Soil
146 characterization revealed lower soil fertility (+16.8 g organic matter/kg of soil) and
147 a deeper water table (mean difference of +6 m) at Site2 than at Site1. Climatic
148 measurements showed that there was more rainfall at Site 2 (mean of +15% for
149 total annual rainfall), whereas temperatures were similar at the two sites
150 (supplementary Table S1). A total of 192 half-sib families obtained from crosses
151 between identified seed parents and two pollen mixtures of identified donors were
152 studied here. 171 families were planted on both sites with 35 individuals per family
153 in a complete block design with single-tree plots (1,250 trees/ha). 21 families were
154 planted on Site2 only, with the same design. Each site also includes 5 checklots
155 composed of individuals from improved and unimproved reference varieties.
156 Thinning operations were performed at both sites in 2012 and exclusively at Site 1
157 in 2017, when the trees were 16 and 21 years old, respectively. A subsample (POP)
158 of 25 half-sib families, with 13 individuals per family and per site, was selected as
159 representative of the variability of growth (total of 650 individuals). In the absence
160 of cloning in this maritime pine experimental context, the notions of “individual”,
161 “tree” and even “genotype” are considered equivalent in our study.

162 Genetic characterization of POP

163 Genomic DNA was extracted from needles collected from POP, to which we added
164 186 randomly selected duplicates for repeatability estimates. The concentration
165 and quality of DNA for each sample were determined with a NanoDrop
166 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). Genotyping
167 was performed by Thermo Fisher Scientific (Thermo Fisher Scientific, Santa Clara,
168 CA, USA) with the 4TREE Axiom single nucleotide polymorphism (SNP) multi-species
169 array (Guilbaud *et al.*, 2020). Of the 50,000 SNPs on this array, 13,407 have been
170 designed for maritime pine and are considered polymorphic for this species. The
171 preliminary filters recommended by Thermo Fisher Scientific were applied to the
172 genotyping results, at the sample (DishQC \geq 0.4, CallRate \geq 90) and SNP (CallRate \geq
173 95, fld-cutoff \geq 3.2, het-so-cutoff: \geq -0.1) levels. In addition, sequential filtering was
174 applied, with the removal, in the following order, of SNPs with less than 85%
175 repeatability, SNPs with more than 5% Mendelian segregation errors and SNPs with
176 a minor allele frequency (MAF) below 1%. A genomic relationship matrix (G) was
177 calculated with the VanRaden formula (VanRaden, 2008) using the AGHmatrix
178 package (Amadeu *et al.*, 2016) in R 4.2.2 environment (R Core Team, 2022):

$$179 \quad G = \frac{(M - P)(M - P)'}{2\sum p_i(1 - p_i)} \quad (1)$$

180 where the M matrix (n : number of individuals \times m : number of markers) contains
181 marker information coded as -1 for one of the homozygotes, 0 for heterozygotes
182 and 1 for the other homozygotes; and the P matrix ($n \times p$) contains allele

183 frequencies expressed as $2(p_i - 0.5)$, where p_i is the frequency of the second
184 allele at locus i for all individuals.

185 In addition, pedigree recovery was performed for each tree from POP, with a subset
186 of 161 SNPs used to infer the identities of the parents (25 seed parents and 85
187 pollen parents) and grandparents (69 initial progenitors from the base population
188 of the breeding program) (supplementary Method. S1). The most complete version
189 of the pedigree was used to compute an additive relationship matrix A for further
190 analyses.

191 Phenotypic data

192 Circumference measurements were performed at breast height (~1.30m from the
193 ground) on all the trees in the trial in 2004, 2008, 2012 and 2018, at the ages of 8,
194 12, 16 and 22 years, respectively. In addition, cores were removed from the trees
195 of POP in December 2019, at breast height, along the same north-south direction
196 for each tree. These cores were cut into 2-mm-thick radial strips for X-ray analysis
197 (Polge, 1966) to obtain wood density profiles (Fig. 1). The limits between the
198 different rings were identified with Windendro software (Guay *et al.*, 1992) and
199 validated by visual examination. The area of ring y ($RA_{raw,y}$) was calculated at
200 individual level as follows:

$$201 \quad RA_{raw,y} = \pi (L + l_y)^2 - \pi L^2 \quad (2)$$

202 where L is the sum of the ring widths from the pith to ring y (ring y excluded) and
203 l_y is the width of ring y . RA_{raw} values are a good proxy for biomass produced each

204 year independently of tree age, in contrast to ring widths which tend to decrease
205 progressively over the years due to radial growth of the tree.

206 We chose to study the 2005-2019 period (15 successive years) here because rings
207 for this period were available for at least 99% of POP and this period excludes the
208 juvenile phase of the trees (supplementary Table S2 and Fig. S1). Using the
209 circumference measurements, RA_{raw} values were spatially corrected for each site
210 with spline functions (via the *BreedR* R package; Muñoz and Sanchez, 2020; see
211 supplementary Method. S2) and named RA (adjusted ring area). A complete
212 phenotyping series for an individual is thus composed of 15 RA values.

213 [Characterization of the environment during ring growth](#)

214 The environmental conditions associated to each ring were characterized with two
215 classes of environmental indices, which depend on both year and site variables. The
216 first class focused on a purely climatic description, with two versions (DM and DM')
217 of the de Martonne aridity index (de Martonne, 1926), whereas the second
218 provided a finer description of the environmental conditions with two indices (GP
219 and GP'), extracted from an ecophysiological model combining climatic,
220 silvicultural and soil data (Moreaux *et al.*, 2020).

221 The de Martonne aridity index was calculated for each ring formed in year y at site
222 z with:

$$223 \quad DM_{y,z} = \frac{1}{8} \sum_{i=3}^{10} \frac{12P_{i,z}}{T_{i,z} + 10} \quad (3)$$

224 where $P_{i,z}$ is the amount of precipitation (in mm) and $T_{i,z}$ is the mean air
225 temperature (in °C), for month i in site z . Only the 8 months from March ($i = 3$) to

226 October ($i = 10$) were included here as we considered, as a first approximation,
227 that climatic conditions outside the growth period of maritime pine has no impact
228 on annual RA. In addition, we considered a modified version of the de Martonne
229 index (DM') based on a 30-day sliding window average (instead of calendar months)
230 and considering the impact of the climate of year ($y - 1$) on environmental
231 conditions in year y (inspired by Botzan *et al.*, 1998, supplementary Method. S3A).
232 The environmental indices of the second class derived from the ecophysiological
233 model GO+ 3.0 (Moreaux *et al.*, 2020) based on climatic data, silvicultural
234 parameters, soil water properties, soil fertility and reference values for maritime
235 pine growing in the Landes massif (supplementary Method. S3B). The growth
236 potential index (GP) was calculated for each ring, based on mean trunk water
237 potential and temperature estimated daily by the GO+ model (supplementary
238 Method. S3C). Similarly, to the de Martonne aridity indices, a second index GP' was
239 used to consider a sliding window of 10 days over the course of a year and to take
240 into account the impact of previous year.

241 Genetic analysis of radial growth with a random regression model 242 (RRM)

243 Unlike a standard analysis of radial tree growth based on final circumference
244 measurements (supplementary Method. S4), we proposed here to model individual
245 RA series for POP as a function of the environmental gradient, using an RRM
246 implemented in Wombat software (Meyer, 2007). The environmental gradients
247 associated with the four indices previously described were modeled independently

248 according to the RRM formulation. Regardless of the environmental index used, the
 249 joint analysis of the two sites and year series provided an overall environmental
 250 gradient of 30 levels (15 environmental levels per site). Legendre polynomials were
 251 used as the base functions (Kirkpatrick *et al.*, 1990) for the following RRM (Mrode
 252 and Thompson, 2005):

$$253 \quad RA_{ijs} = \sum_{k=0}^{k_m} \Phi_{ijk} m_{sk} + \sum_{k=0}^{k_\alpha} \Phi_{ijk} \alpha_{ik} + \sum_{k=0}^{k_p} \Phi_{ijk} p_{ik} + e_{ijs} \quad (4)$$

254

255 where RA_{ijs} is the ring area of individual i for environmental level j at site s ; m_{sk}
 256 is the k^{th} fixed regression coefficient used to model the average trajectory at site s ;
 257 α_{ik} and p_{ki} are the k^{th} random regression coefficients for the genetic additive and
 258 permanent environmental effects, respectively, of individual i , the latter effect
 259 representing the similarity between repeated records for the same individual of
 260 environmental and non-additive genetic origin; Φ_{ijk} is the k^{th} Legendre polynomial
 261 for the RA of individual i at environmental level j ; k_m, k_α, k_p are the order of
 262 polynomials for mean trajectory, genetic additive and permanent environmental
 263 effects, respectively; and r_{ijs} is a random residual. The goodness-of-fit of the
 264 models for the different orders of the polynomials used was assessed by comparing
 265 Aikake information criterion (AIC) and Bayesian information criterion (BIC)
 266 (supplementary Fig. S2). According to these criteria, the order 2 and order 3 models
 267 appeared to be the most relevant models for the environmental indices and
 268 relationship matrices considered in this study. Order 2 models have been preferred

269 since they allows a drastic reduction in computational demand with no loss or
270 marginal loss of goodness-of-fit compared to order 3 models. By setting $k_m = k_\alpha =$
271 $k_p = 2$, we fitted a baseline, a linear and a quadratic regression on RA.

272 The equivalent matrix notation for this model is (Mrode and Thompson, 2005):

$$273 \quad y = Xb + Zu + Qpe + e \quad (6)$$

274 where y is the vector of RA over the environmental levels; b is the vector of
275 solutions for site fixed effect; u and pe are the vectors of the individual genetic
276 additive and permanent environmental random regression coefficients,
277 respectively; e denotes the residuals. X , Z and Q are the corresponding incidence
278 matrices. For genomic-based RRM, it is assumed that $u \sim N(0, G \otimes \Omega)$,
279 $pe \sim N(0, I \otimes P)$, and $e \sim N(0, I \otimes D)$, where \otimes denotes the Kronecker product, G
280 the relationship matrix described above, Ω and P the covariance matrices for the
281 RR coefficients for the genetic additive and permanent environmental effects,
282 respectively, and D is a diagonal matrix of heterogeneous residuals for each
283 environmental level. For pedigree-based RRM, G is replaced by A .

284 With a second-order model ($k_m = k_\alpha = k_p = 2$), the RRM estimates three genetic
285 coefficients per individual. From these, individual GEBV were then obtained at all
286 environmental levels as a trajectory, following the formulation of (Mrode and
287 Thompson, 2005). GEBV estimated with an RRM integrating all available phenotypic
288 data and solved at each environmental level are denoted $GEBV_{ref}$.

289 Genomic selection

290 *Cross-validation (CV) scenarios*

291 The prediction accuracy of the RRM was assessed over two CV scenarios (Fig. 2).

292 First, the reference scenario, denoted **CV-A**, where the training set (T_{set}) included

293 the complete phenotyping series for 50% of the individuals (randomly selected

294 within sites and families), whilst the remaining 50% of individuals constituting the

295 validation set (V_{set}). Second, the CV-B scenario explored the possibility of retaining

296 the same amount of phenotypic information as for the CV-A (i.e. 50% of total

297 phenotypic data) but distributed differently over the individuals. Scenario **CV-B**

298 mimicked the use of a high-throughput phenotyping tool for quick estimation of

299 the last five RA which, in a context of global warming, would typically correspond

300 to unfavorable years. The T_{set} for **CV-B** included complete phenotypic series (i.e. 15

301 phenotypic records per individual) for 25% of individuals and only five phenotypic

302 records for the remaining individuals (75% of individuals). For each site, we kept

303 the same five environments for each repetition of the CV scenario. These 5

304 environments were chosen to represent the most unfavorable half of the gradient.

305 The Kennard-Stone algorithm (Kennard and Stone, 1969) was applied via the

306 *prospectr* R package (Stevens and Ramirez-Lopez, 2022) to maximize the Euclidean

307 distances between the GP' values and thus select 5 representative environments

308 from the 8 most unfavorable environments at each site.

309 The prediction accuracy (i.e. the correlation between the true and predicted

310 breeding values) of the RRM was estimated for each environmental level as the

311 Pearson correlation coefficient between predicted ($GEBV_{pred}$) and observed RA in
312 V_{set} divided by the square root of heritability. The overall prediction accuracy was
313 then obtained by averaging the prediction accuracies of each environment. For
314 each CV scenario, 10 independent repetitions of this process were performed. Such
315 performance estimator was used as a criterion for assessing modeling quality (Ly *et al.*,
316 2018; Arnal *et al.*, 2019; Momen *et al.*, 2019).

317 *Genetic gains*

318 The prediction accuracy of the RRM for genetic gains in our reference scenario CV-
319 A was assessed over each of the environmental levels. The assessment consisted of
320 calculating the differences in genetic gain between a selection based on $GEBV_{pred}$
321 obtained in V_{set} and the corresponding maximum that would have been obtained
322 with the same selection intensity based on $GEBV_{ref}$. For this, at each environmental
323 level, the top 5% of individuals selected according to $GEBV_{pred}$ were identified and
324 their corresponding $GEBV_{ref}$ (obtained with all the phenotypic information) used to
325 calculate the true genetic gain (GG_{true}) as the $GEBV_{ref}$ average of the selected
326 individuals. This amount was compared for the corresponding environmental level
327 to the maximum gain (GG_{max}), which was calculated as the $GEBV_{ref}$ average of the
328 top 5%. Finally, GG_{true} and GG_{max} were centered and reduced to ensure
329 comparability between environmental levels. Any difference between GG_{true} and
330 GG_{max} would indicate a decrease in the correlation between $GEBV_{pred}$ and $GEBV_{ref}$
331 for the selected percentage.

332 Results

333 Size and genetic characterization of POP

334 After phenotype curation (9 wood-density profiles were excluded as they were not
335 readable enough to allow ring limits to be positioned with confidence) and
336 genotyping quality control (13 individuals excluded), POP was finally composed of
337 628 trees (303 from Site 1 and 325 from Site 2).

338 Pedigree recovery on POP validated 93% of the pedigree seed parents (monoicous
339 individuals acting as mothers) and allowed the correction of 5%. The remaining 2%
340 of the pedigree seed parents was classified as unknown, as no candidate parent
341 could be validated. Pollen parents (acting as fathers) were successfully recovered
342 for 65% of the individuals. Note that the original design of the study was based on
343 crosses with a mixture of pollen donors, resulting in the fathers initially being
344 unknown in the pedigree. Finally, based on the curated pedigree, a status number
345 (N_S ; Lindgren *et al.*, 1996) of 21 was obtained for POP, suggesting a high level of
346 relatedness between the families studied.

347 The genotyping of POP resulted in the characterization of the 628 individuals over
348 3,832 SNPs, with a repeatability of 97% and a total missing data rate of 1%. Additive
349 genomic relationship coefficients (g_{xy}) estimated in G were consistent with the
350 pedigree-based additive relationship coefficients (a_{xy}) calculated in A (Fig. 3). The
351 a_{xy} values were discrete, whereas the g_{xy} values were normally distributed for
352 each level of relatedness. Note that, for most pedigree-based additive coefficient
353 levels, the normal distribution has a long upward-sloping tail (revealing some rare

354 cases of unrecorded relatedness), and a mean slightly below the theoretical value,
355 the latter being represented by the gray line in Fig. 3. This shift is expected insofar
356 as the standardization of G matrix with the observed allele frequencies sets its
357 average to 0.

358 Quality of model fit

359 The prediction accuracy (estimated with CV-A) was used as a criterion for assessing
360 the quality of RRM (Fig. 4). Mean prediction accuracies were moderate to high, with
361 correlation coefficients ranging from 0.19 to 0.25. Prediction accuracy was slightly
362 better (+0.04 better, on average) for genomic-based RRM than for pedigree-based
363 RRM, except for RRM based on the *DM* environmental index (equivalent mean
364 prediction accuracy of 0.21). The best prediction accuracies were obtained for
365 genomic-based RRM with the *DM'* (0.24) and *GP'* (0.25) indices. The optimization
366 of environmental indices improved slightly RRM prediction accuracy by 16% and 3%
367 relative to the initial *DM* and *GP* indices, respectively. Finally, the genomic-based
368 RRM using the *GP'* index was selected for the analyses described below, due to its
369 best prediction accuracy (0.25) for GS. Detailed prediction accuracies by
370 environment for this model are presented in the supplementary Fig. S3.

371 Narrow-sense heritability estimated for this model varied between 0.12 and 0.24
372 over the environmental gradient (supplementary Fig. S5). Despite the high level of
373 variation between similar environmental levels, heritability was significantly higher
374 for the most favorable environmental levels.

375 Individual reaction norms estimated by genomic-based RRM

376 Reordering longitudinal data by the annual environmental index, which
377 characterizes the conditions of ring formation, instead of the ordinal year greatly
378 modified the shape of the mean RA curve in a more easily interpretable way (Fig.
379 5). When expressed as a function of the environmental index GP' , RA increases
380 significantly. The lowest GP' values are associated with the most unfavorable
381 environmental conditions for growth, whereas the highest values are associated
382 with the most favorable conditions for growth. This pattern suggests plasticity at
383 the population level, but hides individual behaviors, which may deviate from this
384 central trajectory.

385 Random individual deviations from the mean trajectories due to additive genetic
386 effects are represented in Fig. 6 and were solved over the environmental gradient
387 of GP' ($GEBV_{ref}$). For most individuals, $GEBV_{ref}$ showed a dependence on GP' ,
388 highlighting the existence of plasticity for RA. These different behaviors can be
389 characterized simply by the slope of the trajectories, depicted in different colors
390 (Fig. 6). A majority of individual reaction norms were characterized by shallow
391 slopes and mean $GEBV_{ref}$ close to 0. This does not mean that the phenotypic
392 trajectory of these individuals is flat. Instead, it indicates that they have trajectories
393 indistinguishable from the mean trajectory and due to its additive genetic origin,
394 they would give only very limited extra plasticity to the offspring, unlike individuals
395 whose trajectories are significantly further from 0. The highest and lowest mean
396 $GEBV_{ref}$ were those obtained for individuals whose trajectory is colored dark blue

397 and dark red, respectively. These individuals also display reaction norms with the
398 strongest positive and negative slopes (for dark blue trajectories and dark red
399 trajectories, respectively), leading to a greater range of variation in individual
400 genetic values in favorable than in unfavorable environmental levels. The average
401 $GEBV_{ref}$ for each individual is, thus, strongly correlated with the regression
402 coefficients describing the slope of its trajectory (+0.52 and +0.95 with quadratic
403 and linear regression coefficients, respectively). There appear to be few
404 intersections between reaction norms, corresponding to changes in individual
405 ranks across environmental levels, over most of the environmental gradient. This is
406 confirmed by strong genetic correlations (>0.90) between environmental levels
407 (supplementary Fig. S4). However, large overlaps occur in the part of the gradient
408 corresponding to unfavorable environmental levels (Fig. 6). This results in lower
409 genetic correlations (between 0.83 and 0.90) between the two most unfavorable
410 environmental levels ($GP^i=52$ and $GP^i=53$) and more favorable environmental
411 levels (when $GP^i>65$). These lower values point out variations in how certain
412 individuals behave at these unfavorable environmental levels, which affects their
413 ranking.

414

415 Genomic selection scenarios and cross-validation

416 *Genetic gain over the environmental gradient*

417 The overall prediction accuracy of the genomic-based RRM (using the GP^i
418 environmental index) estimated with the CV-A scenario was 0.62 (Fig. 4). Breeding

419 efficiency, based on predicted values, was assessed by calculating genetic gains for
420 different environmental levels (Fig. 7). GG_{\max} increased until the environmental
421 value of 62, above which it reached a plateau with maximum value of 2.35. The
422 difference between GG_{\max} and GG_{true} was minimal (+0.64) for GP' environmental
423 level 62, increasing towards the two extreme environmental levels. Indeed, this
424 difference reached +0.73 and +0.72 respectively for the most unfavorable
425 environmental level 53 and for the most favorable environmental level 82. The
426 relatively moderate prediction accuracy of the RRM (0.62) necessarily led to a
427 significant loss of genetic gain (no overlap between GG_{\max} and GG_{true} boxplots).
428 Nevertheless, depending on the environmental level, GG_{true} accounted for 68% to
429 73% of GG_{\max} . GG_{true} was always significantly different from 0 ($p - \text{value}_{t-\text{tes}} <$
430 0.001), indicating a certain efficiency of selection based on predicted values, even
431 in the most extreme environmental levels.

432 *Prediction accuracy over the CV scenarios*

433 We considered an alternative cross-validation scenario (CV-B) (Fig. 2), to improve
434 selection efficiency while preserving phenotyping effort with respect to CV-A. As in
435 CV-A, 50% of the phenotypic data were used to constitute the T_{set} of the CV-B. The
436 key difference between the two is due to a better distribution of phenotypic effort,
437 both between individuals and between environments, in CV-B. This alternative
438 distribution had a considerable impact on improving the prediction accuracy of the
439 RRM, which increased from 0.62 for the CV-A to 1.37 for the CV-B (Fig. 8), with no
440 increase in phenotyping effort. It should be noted that the CV-A scenario is a major
441 challenge for RRM, as it imposes the prediction of entire trajectories for half of the

442 population. This challenge is relaxed in CV-B by including at least partial information
443 for all individuals.

444

445 Discussion

446 Deciding which genetic material should be planted now to form the forests of
447 tomorrow is becoming increasingly challenging due to the rapidity of climate
448 change (Thomas *et al.*, 2004; Wiens, 2016). Using longitudinal tree-ring data and
449 parallel environmental descriptors, we have successfully modeled genomic
450 individual reaction norms based on random regression. This first example for forest
451 trees provided consistent results for use in the maritime pine breeding program,
452 but may inspire other programs in perennial species.

453 Reaction norms in forest trees

454 Growth measurements at advanced age are generally used for the calculation of
455 breeding values. Such measurements constitute highly integrative phenotypes that
456 can be associated only with a global environmental site index. Using sites with
457 contrasting indices has been a classic strategy to establish comparative trials for
458 genetic x environment evaluation. In this sense, our two sites present strong
459 contrast in terms of fertility and water table depth at the scale of the Landes massif,
460 but even with their differences they are still part of the same breeding area (Jolivet
461 *et al.*, 2007). Wood cores give us access to phenotypic inter-annual variation and
462 can be used to generate longitudinal annual growth data that can be associated

463 with annual environmental variation. Our results showed indeed that the
464 environmental variation between years was much greater than the one between
465 sites (Fig. 5). Indeed Cir22 was associated with a mean environmental index GP' of
466 68.2 for site 1 and 72.1 for site 2, whereas analysis based on ring measurements
467 covered a larger index range (GP' from 52.6 to 81.9). This much greater annual
468 variation provides an opportunity to infer plasticity at individual level over a large
469 environmental gradient.

470 In addition to longitudinal data collection, which can be operationally costly, there
471 are other challenges that arise with these data. One is autocorrelation between
472 repeated measurements on the same individual in a time series. Another, not least,
473 is ontogenetic differences between phases of phenotype expression (Sanchez *et al.*,
474 2013). Finally, a third challenge is the choice of a relevant environmental descriptor.
475 Although we have not shown it for simplicity, we have performed a preliminary
476 RRM for RA with a one-year lag in the climatic index in order to match RA of year n
477 with the environmental index of year $n - 1$, and its results pointed to an absence of
478 autocorrelated effect. As for the ontogeny challenge, we have ignored in our
479 longitudinal data series the initial segments corresponding to the juvenile phase,
480 keeping only the remaining adult phase for which the RA trend was generally flat,
481 despite strong inter-annual oscillations (Fig. 5A).

482 The third challenge is probably the most difficult to address, the choice of a relevant
483 environmental index (Li *et al.*, 2017). This study was not designed to identify
484 precisely the environmental factors most relevant to tree growth, but we defined

485 two classes of biologically meaningful environmental indices that integrate the key
486 components of temperature and water (Begum *et al.*, 2013; Rathgeber *et al.*, 2016).
487 Both of them depend on the year and the site in which the ring was formed. The
488 first class (aridity indices) is easy to obtain, since it only considers the climatic data
489 (temperature and precipitation over the growing period) of the site and the year
490 associated with to the rings under study. On the other hand, the second class
491 (growth potential indices) requires more complex modeling, including for example
492 the characterization of the daily water status of the trunk. A major difference
493 between the two types of indices is the insensitivity of the former to the intra-
494 annual distribution of precipitation and temperatures. Thus, similar annual aridity
495 values (DM or DM') may reflect different climatic realities over the course of the
496 growing season, with temperatures and/or precipitation occurring at different
497 periods and leading to differences in growth. Conversely, by considering the daily
498 environmental status and tree physiology, the growth potential indices (GP and
499 GP') allowed a more detailed consideration of within-year environmental variation.

500 Finally, the prediction accuracy obtained with DM , DM' , GP , GP' (Fig. 4) confirms
501 the relevance of the proposed environmental indices, but also suggest that they
502 only partially capture the environmental factors influencing radial growth and the
503 differences between individuals' reactions. More specifically, the variability due to
504 site is not fully described by the index, given the remaining high significance value
505 of the corresponding fixed effect in RRM (Fig. 5B).

506 Modelling reaction norms with RRM

507 Unlike univariate single-point analyses, which are easy to implement but do not
508 integrate longitudinal phenotypic information, or multi-trait models, which can
509 integrate it but are computationally demanding, RRM provides genetic estimates
510 over the chosen continuous environmental gradient with reduced parametrization
511 (Sun *et al.*, 2017). The continuous trajectory of GEBV predicted by the RRM allows
512 a position to be considered at any environmental level within the range defined by
513 the two most extreme environments, whether it has actually been observed or not.
514 The RRM can model highly complex curves using orthogonal base functions such as
515 Legendre polynomials, which are widely used and described in the context of
516 breeding (Schaeffer, 2004; Campbell *et al.*, 2018; Marchal *et al.*, 2019). Despite
517 their great flexibility and computational advantages, Legendre polynomials may
518 present numerical problems (Runge's phenomenon) at the extremities for high-
519 order fits (de Boor, 1978; Meyer and Kirkpatrick, 2005). In this study, the
520 adjustment at the extremities of the environmental gradient was particularly
521 important as the unfavorable extreme conditions are likely to increase in frequency
522 in the future (Coumou and Rahmstorf, 2012; Spinoni *et al.*, 2018). The use of low-
523 order polynomials to model RA trajectories overcame this problem. The
524 consistency and quality of the norms fitted with Legendre polynomials were
525 verified by a comparison with norms fitted with B-spline functions, which are
526 considered a more robust alternative to high-order polynomials in terms of
527 extremum fitting, although less advantageous computationally (de Boor, 1978;
528 Meyer and Kirkpatrick, 2005) (Kendall correlations between $GEBV_{ref}$ estimated with

529 Legendre polynomials and those estimated with B-splines yielding coefficients of
530 up to 0.95 over the entire gradient for the final RRM).

531 [Exploration of individual genetic trajectories](#)

532 Random individual trajectories (Fig. 6) highlight the existence of plasticity for
533 genetic values that can be targeted by breeders. It is not easy to discriminate
534 between individual reaction norms that follow a trajectory close to the population
535 average, given their high frequency, the fact that they present shallow slopes and
536 mean $GEBV_{ref}$ close to 0. However, individuals with potentially good growth along
537 the entire gradient are much easier to discriminate from the rest, for which the
538 proposed clustering allows simple and efficient visualization (cluster E), useful for
539 selection purposes.

540 The distribution of individual GEBV varied between environmental levels, and those
541 more favorable levels enhanced the expression of differences between trajectories
542 relative to less favorable levels, which has already been observed in other biological
543 models (Arnold *et al.*, 2019). Individual ranking was globally preserved over the
544 trajectories for most of the gradient (van Eeuwijk *et al.*, 2016). This trend was
545 confirmed by strong genetic correlations (supplementary Fig. S4) between the
546 environmental levels. However, these genetic correlations were weaker with
547 unfavorable environments (environmental index below 65), in agreement with the
548 reranking of individuals observed for the individual trajectories at the most
549 unfavorable end (Fig. 6). This precise and localized GxE interaction in our gradient,
550 only possible thanks to the use of the RRM, should not be considered marginal or

551 potentially negligible considering that it affects only one segment of the gradient.
552 In fact, climate projections (supplementary Fig. S6) suggest that such unfavorable
553 environments are likely to become much more common in the future. Even if the
554 expected global level of aridity in 2075 remains close to current levels, according to
555 our de Martonne calculation, aridity in 2100 will be much stronger, with a higher
556 frequency of extreme events as predicted by other studies (Sillmann and Roeckner,
557 2008; Lehner *et al.*, 2017). Our 15-year study period was already affected by a high
558 global level of aridity and included extreme annual climates that may become
559 frequent in the future. The environmental gradient used for the inference of
560 reaction norms is therefore particularly relevant for identifying individuals with
561 better potential for growth in the unfavorable years to come.

562 When GxE interactions must be taken into account in selection decisions, a robust
563 strategy would involve prioritizing the best adapted individuals across the entire
564 environmental gradient (Li *et al.*, 2017), focusing on the notion of persistence. The
565 definition of persistence may vary according to species and breeding aims (Gengler,
566 1996; Rocha *et al.*, 2018), but it is generally defined as the capacity of a species to
567 maintain a stable or high level of growth or production over time or in the face of
568 different environmental conditions. For reactions norms, several ways of evaluating
569 persistence and integrating the slope of trajectories in an operational breeding
570 context have been proposed. For example, for feed conversion ratio in large white
571 pigs, (Huynh-Tran *et al.*, 2017) suggested combining the EBV estimated by the RRM
572 with the coefficients of eigenvectors estimated from the eigenvalue decomposition
573 of the covariance matrix of additive genetic effects. In their study, two summarized

574 breeding values for each individual were sufficient to describe most of the variation
575 in terms of mean genetic values (first dimension) and the slopes of EBV trajectories
576 (second dimension), and could be used directly in selection. In another example in
577 goat lactation, (Arnal *et al.*, 2019) considered “the cumulative deviation in genetic
578 contribution to yield relative to an average animal having the same (initial) yield”
579 for the calculation of persistence-related EBV. Finally, (Peixoto *et al.*, 2020)
580 suggested the ranking of cotton genotypes on the basis of area under the reaction
581 norm, the genotype with the highest norm being the most persistent. Another
582 interesting approach would involve calculating the final GEBV for each individual as
583 the mean of the GEBV for each environmental level weighted by the probability of
584 occurrence of the environmental level in the future. Such a strategy would make
585 use of the GxE interaction to maximize genetic gain for individuals performing in
586 environmental conditions close to those predicted for the near future, while
587 ensuring a certain level of resilience to environmental variation. Any of these
588 proposals could be applied to our data. A possible advantage of the latter strategy
589 could be to take more explicit account of future climate predictions, provided that
590 they have some control over uncertainty.

591 [Reaction norm in a GS context](#)

592 The use of genomic reaction norms to predict the growth of individuals in
593 environments where they have not been observed is a good example of the
594 potential benefits of GS approaches for traits that are complex to evaluate. Wood
595 density profiles provide highly informative longitudinal data on tree growth over

596 the years, but its acquisition via the coring process remains costly and time-
597 consuming at breeding-program scale. This limitation has motivated one of our
598 alternative cross-validation scenarios (CV-B), with a more homogeneous
599 distribution of phenotypic effort, resulting in a training population involving all
600 individuals, 25% of which contribute full time series and the remaining 75% only
601 partial 5-year series. Indeed, relative to our baseline scenario (CV-A), which aimed
602 to predict the full trajectories of 50% of individuals, the CV-B scenario achieved a
603 much higher level of prediction accuracy (1.37), demonstrating that the allocation
604 of phenotyping effort to constitute the training population is a key optimization to
605 consider. The scenario CV-B would reflect the use of a high-throughput
606 phenotyping tool usable on a large number of individuals at the cost of a smaller
607 number of rings scanned per tree, which is basically what a resistograph does
608 (Bouffier *et al.*, 2008). Resistograph measures the resistance of the wood to
609 penetration with a needle and can estimate RA efficiently for the rings closest to
610 the bark, i.e. the last five rings formed (personal communication). These
611 measurements provide only partial information about plasticity, but when applied
612 to the whole population, they have the advantage of providing information
613 complementary to that obtained by coring. Overall, less phenotyping effort is
614 required, but the benefits are substantial.

615 The genetic component of reaction norms, the one of greatest interest to breeders,
616 was estimated by integrating pedigree or genomic information in the RRM.
617 Genomic-based RRM had a significantly better prediction accuracy (with the GP'
618 index) than pedigree-based RRM, suggesting that refining the coefficients of

619 relationships between individuals through their molecular characterization with
620 SNPs results in the generation of more suitable models (Gamal El-Dien *et al.*, 2016;
621 Bouvet *et al.*, 2016). The pedigree information tended towards a systematic
622 overestimation of pairing coefficients relative to the genomic information (Fig. 3).
623 However, some rare pairs of individuals appeared to be much more related on the
624 basis of genomics than on the basis of pedigree, suggesting that, in some cases, the
625 pedigree may be incomplete, or may contain errors, despite the correction and
626 recovery steps (Tan *et al.*, 2017; Li *et al.*, 2019). The use of genomic data for
627 genomic evaluation is often proposed for forest trees (Grattapaglia and Resende,
628 2011; Lebedev *et al.*, 2020), but first GS studies for maritime pine (Isik *et al.*, 2016;
629 Bartholomé *et al.*, 2016) highlighted the difficulty of demonstrating a superiority of
630 genomic models over pedigree-based models. In this study, we provide some
631 arguments to go beyond these limitations in the application of the genomic
632 prediction model. The RRM takes greater advantage of genomic information to
633 predict individual trajectories than pedigree information. Indeed, in a context of
634 intense climate change, the importance of integrating environmental information
635 into genetic evaluation may fully justify the additional cost of genotyping (Isik,
636 2014).

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657 **Author contributions**

658 LB and LS: conceptualization, supervision and validation

659 VP, AB, LB and LS: methodology

660 VP: data curation, formal analysis, visualization, writing – original draft preparation

661 VP and AB: software

662 AB, LB and LS: writing – Review & Editing

663 **Competing Interests**

664 The authors declare no competing financial interests

665 **Data Archiving**

666 The data underlying this article are accessible via the private following link (Data

667 INRAE):

668 <https://entrepot.recherche.data.gouv.fr/privateurl.xhtml?token=15f2101e-ebb8->

669 [4b7c-838b-9703090cfec4](https://entrepot.recherche.data.gouv.fr/privateurl.xhtml?token=15f2101e-ebb8-4b7c-838b-9703090cfec4)

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672 The data will be made public and accessible to all once the article has been

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674 **Research Ethics Statement**

675 Not applicable.

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Figures

Figure 1: From wood increment core to wood density profile. *From the bottom to the top: the wood increment core picture from one tree, its corresponding radiography, its wood density profile (black line) from pith (position: 0mm) to the bark obtained after processing. Sudden and high drops in wood density mark the end of annual growth and were used to fit each ring limitations.*

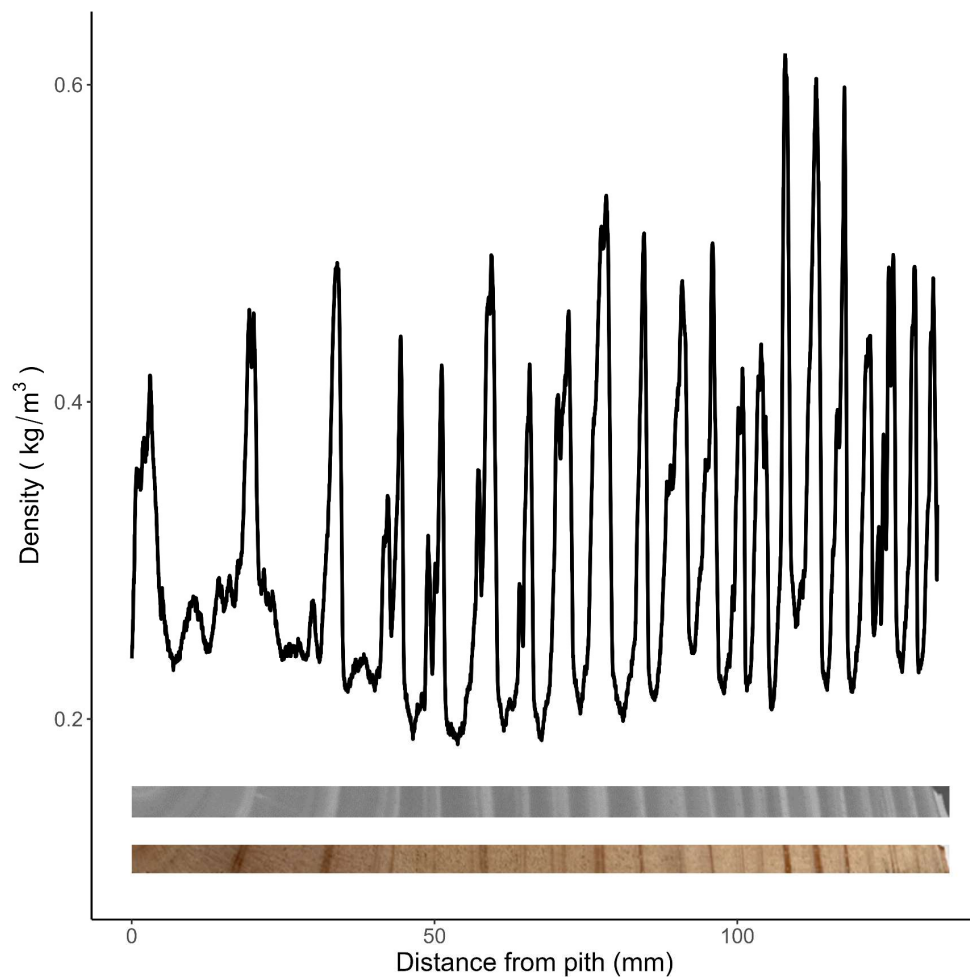


Figure 2: Cross-validation scenarios CV-A, CV-B, CV-C and CV-D performed with a RRM according to the GP' index. All scenarios include the same amount of phenotypic information in the training set (i.e. 50% of total phenotypic data); only the distribution of this information across individuals and environmental levels differ. All families contributed equally to the training set.

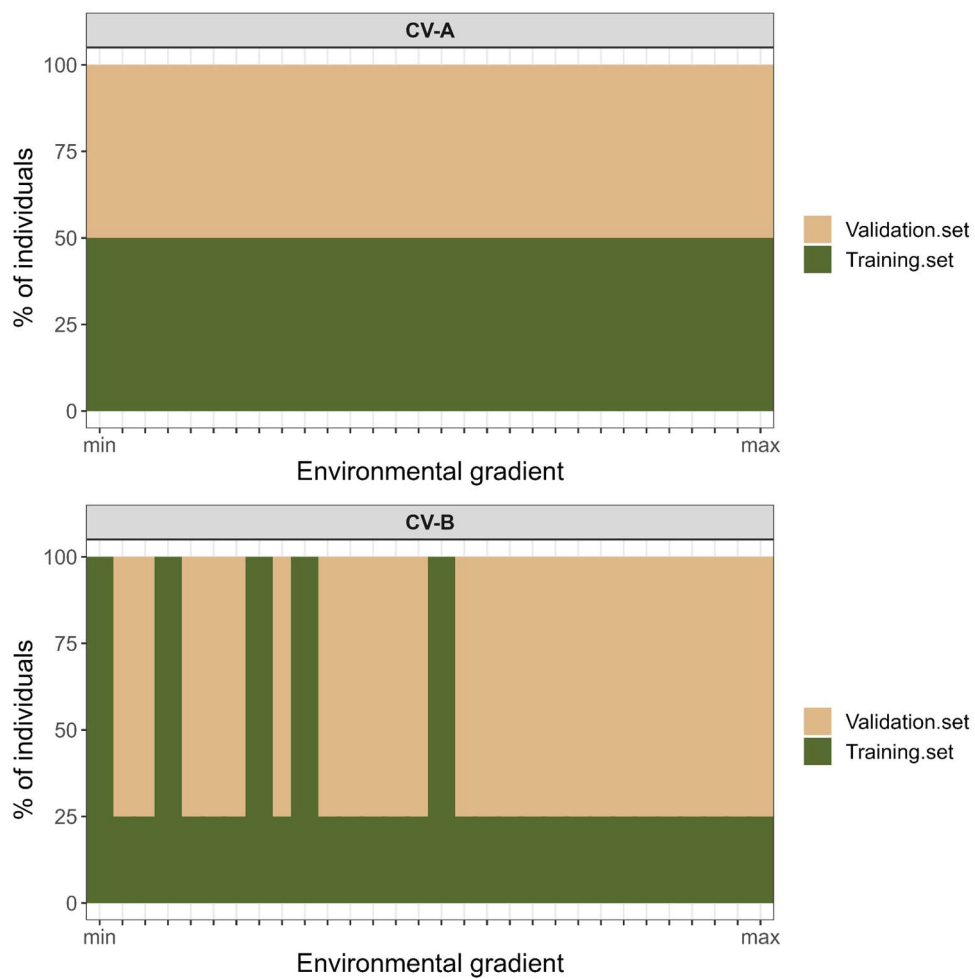


Figure 3: Comparison between pedigree-based additive relationship coefficients derived from pedigree and additive genomic relationship coefficients derived from SNP markers for individuals of POP. For each value of the discrete scale taken by the pedigree-based additive relationship coefficients, the corresponding violin plot represents the continuous distribution of additive genomic relationship coefficients. Numbers below each violin plot denote the number of relationship included in the corresponding violin plot. Grey line is the bisector passing through the origin of the graph. The two highest pedigree-based additive relationship coefficients (1.25 and 1.5) are unique and so represented by single points instead of violin plots.

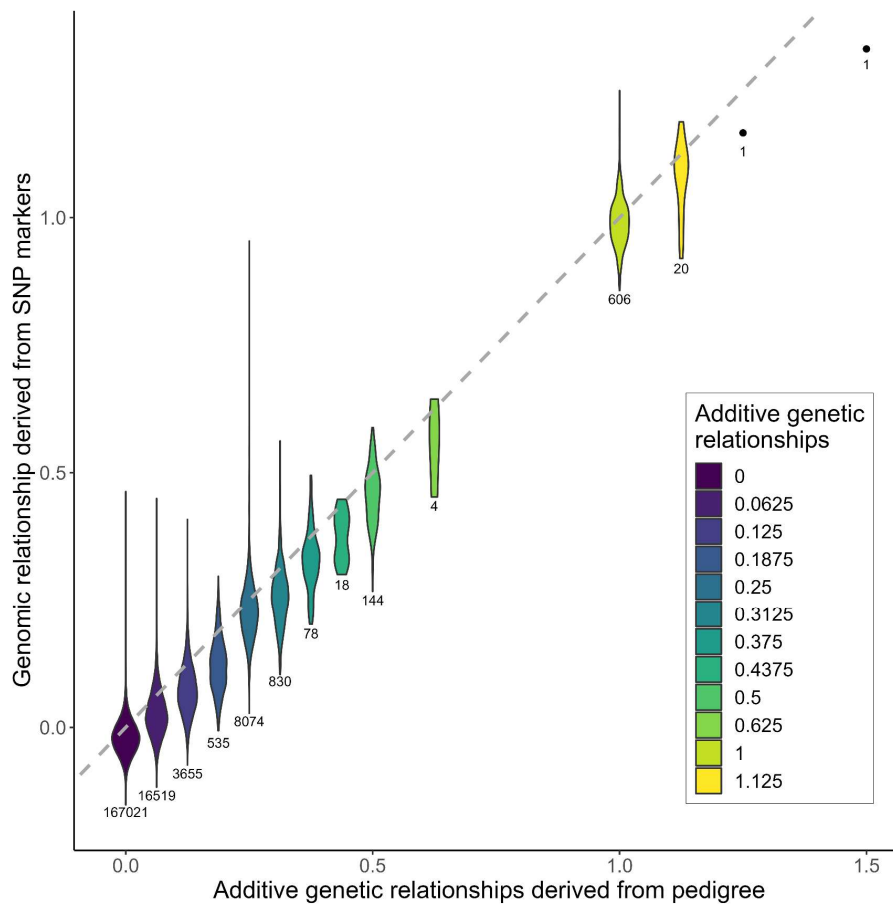


Figure 4: Prediction accuracy of the RRM according to the environmental gradient and the genetic information used. *Boxplots indicates the Pearson correlation coefficient between observed and predicted RA values over the whole environmental gradient for 10 repetitions of the CV-A scenario. Boxplots are blue when the RRM implemented integrated pedigree-based additive relationship coefficients while boxplots are yellow when the RRM implemented integrated additive genomic relationship coefficients. For each kind of genetic information, RRM were run independently with each of the four environmental gradients, respectively derived from DM, DM', GP and GP' indices.*

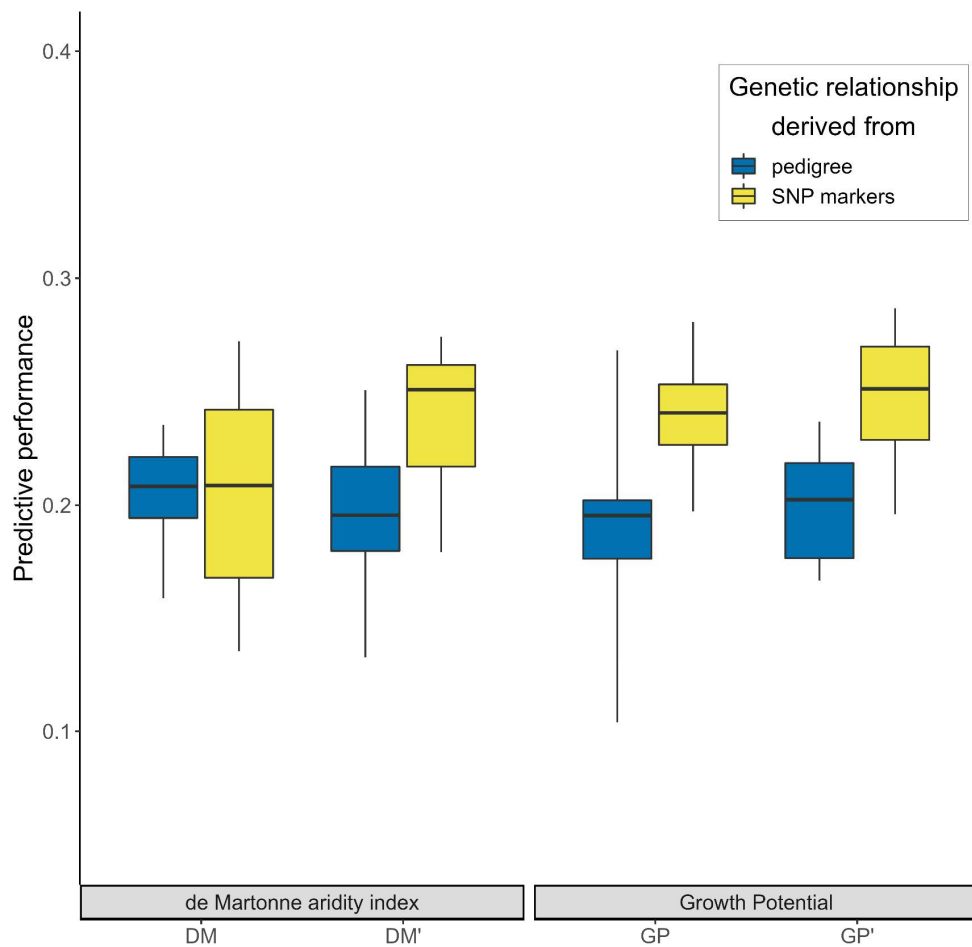


Figure 5: Evolution of mean RA according to the years for each site (Fig. 5A) or according to the GP' index (Fig. 5B). Figure 5.A presents mean phenotypic trajectories of RA and Figure 5.B presents mean trajectories adjusted by the RRM for each site. Both trajectories are the result of the same model. The significance of the slope parameter for each trajectory in Figure 5.B was assessed with Student's t-test (p -value <0.01)

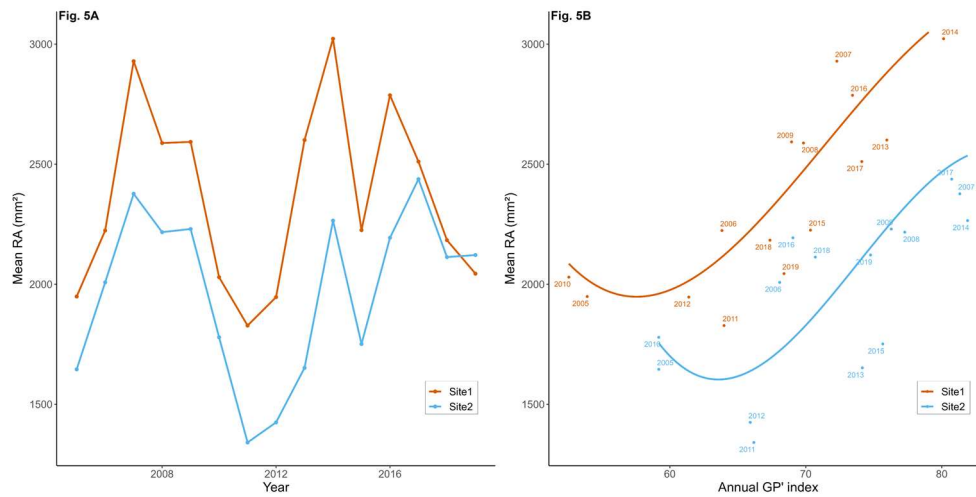


Figure 6: Individual trajectories of $GEBV_{ref}$ associated to RA according to the GP' index. Trajectories correspond to the genetic component of the reaction norms estimated by the genomic based RRM. Trajectories were colored according to their slope (quadratic regression coefficient), from the steepest negative slopes (dark red) to the steepest positive slopes (dark blue).

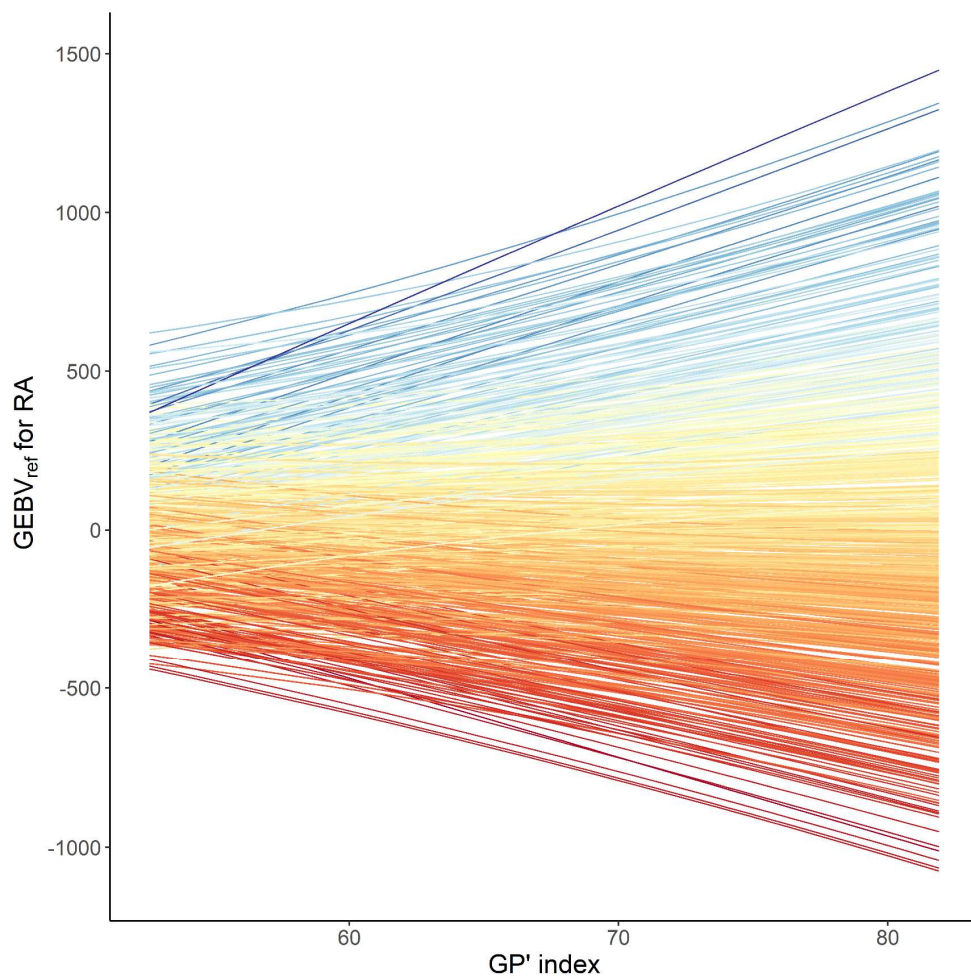


Figure 7: Maximum genetic gain (GG_{max}) and true genetic gain (GG_{true}) according to GP' index. *The RRM was used with complete phenotypic information for all individuals to estimate $GEBV_{ref}$ over the gradient; and then independently repeated 10 times with the scenario CV-A to predict $GEBV_{pred}$ for individuals in the validation set. GG_{max} was calculated as the mean of the top 5% of $GEBV_{ref}$ and for each iteration GG_{true} was calculated as the mean of $GEBV_{ref}$ associated to the top 5% individuals selected based on $GEBV_{pred}$ for each GP' values. GG_{max} and GG_{true} are centered and reduced.*

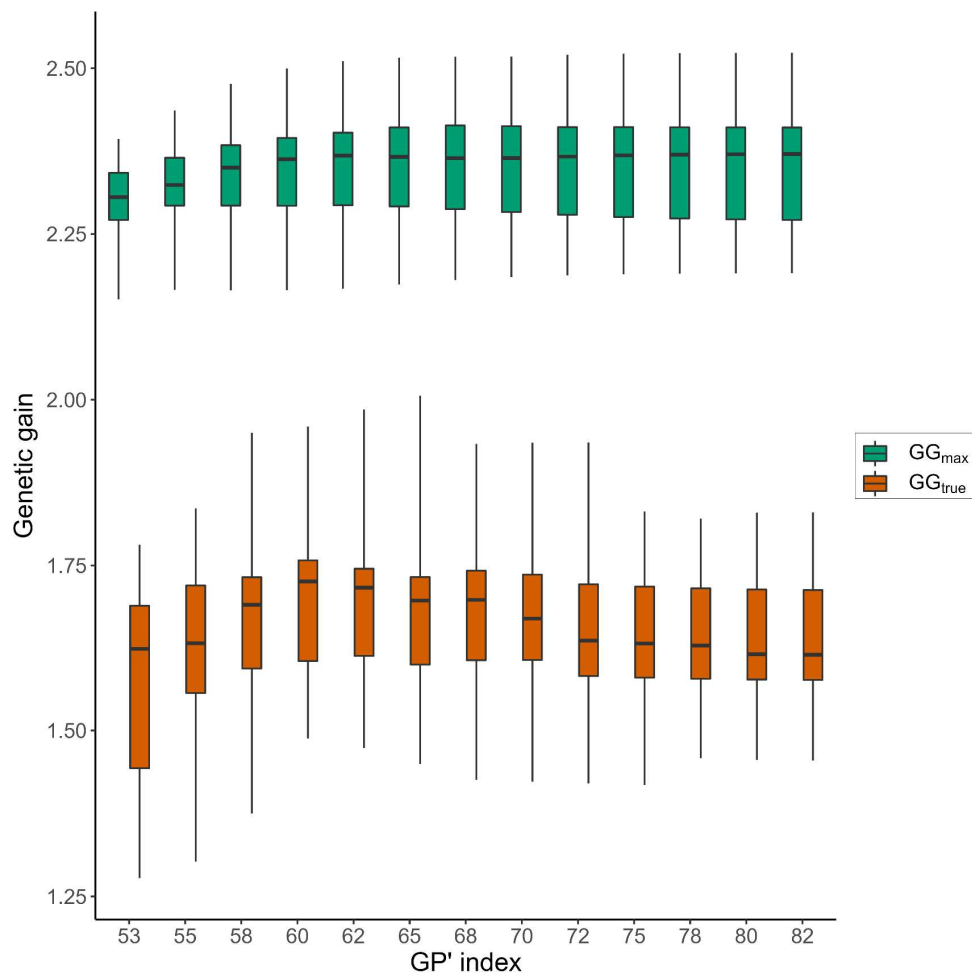


Figure 8: Prediction accuracy of the RRM according to the CV-A and CV-B scenarios. *Boxplots indicates the Pearson correlation coefficient between observed and predicted RA values over the whole environmental gradient for 10 independent repetitions of the CV scenario. The significance between prediction accuracies was assessed by a Student's t-test (****: p-value <1e10-4)*

