- 1 Range-wide variation in local adaptation and phenotypic plasticity
- of fitness-related traits in Fagus sylvatica and their implications
- 3 under climate change
- 4 Homero Gárate-Escamilla¹, Arndt Hampe¹, Natalia Vizcaíno-Palomar¹, T. Matthew Robson² &
- 5 Marta Benito Garzón^{1*}.

- ¹BIOGECO INRA UMR 1202 University of Bordeaux, Pessac, 33400, France
- 8 ² Organismal and Evolutionary Biology (OEB), Viikki Plant Science Centre (ViPS), Faculty of
- 9 Biological and Environmental Sciences, University of Helsinki, P.O. Box 65, Finland 00014.
- *Corresponding author: marta.benito-garzon@inra.fr

11 ACKNOWLEDGEMENTS

- 12 This study was funded by the "Investments for the Future" program IdEx Bordeaux (ANR-10-
- 13 IDEX-03-02). HGE was funded by the CONACYT- Mexico and by the Institute of innovation
- and technology transfer of Nuevo Leon (I2T2), Mexico. TMR was funded by the Academy of
- Finland (decision 304519). We thank the scientific network INRA-ACCAF.

16 **BIOSKETCH**

20

- 17 The authors' research is focused on plant ecology and global change. The authors use
- multidisciplinary approaches including ecophysiology, genetics and ecological modelling to
- understand complex processes in ecology at large geographical scales.

22 Running title: range-wide multi-trait variation 23 **Keywords**: phenotypic variation, species distribution models, beech, acclimation, trait co-24 variation, common gardens 25 26 **ABSTRACT** 27 **Aim:** To better understand and more realistically predict future species distribution ranges, it is 28 29 critical to account for local adaptation and phenotypic plasticity in populations' responses to 30 climate. This is challenging because local adaptation and phenotypic plasticity are traitdependent and traits co-vary along climatic gradients, with differential consequences for fitness. 31 Our aim is to quantify local adaptation and phenotypic plasticity of vertical and radial growth, 32 33 leaf flushing and survival across Fagus sylvatica range and to estimate each trait contribution to 34 explain the species occurrence. 35 **Location:** Europe 36 **Time period:** 1995 – 2014; 2070 37 **Major taxa studied:** Fagus sylvatica L. **Methods:** We used vertical and radial growth, flushing phenology and mortality of *Fagus* 38 39 sylvatica L. recorded in BeechCOSTe52 (>150,000 trees). Firstly, we performed linear mixed-40 effect models that related trait variation and co-variation to local adaptation (related to the 41 planted populations' climatic origin) and phenotypic plasticity (accounting for the climate of the 42 plantation), and we made spatial predictions under current and RCP 8.5 climates. Secondly, we 43 combined spatial trait predictions in a linear model to explain the occurrence of the species.

Results: The contribution of plasticity to intra-specific trait variation is always higher than that of local adaptation, suggesting that the species is less sensitive to climate change than expected; different traits constrain beech's distribution in different parts of its range: the northernmost edge is mainly delimited by flushing phenology (mostly driven by photoperiod and temperature), the southern edge by mortality (mainly driven by intolerance to drought), and the eastern edge is characterised by decreasing radial growth (mainly shaped by precipitation-related variables in our model); considering trait co-variation improved single-trait predictions. Main conclusions: Population responses to climate across large geographical gradients are traitdependent, indicating that multi-trait combinations are needed to understand species' sensitivity to climate change and its variation across distribution ranges.

1. INTRODUCTION

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

Climate change is having a major impact on the structure, composition and distribution of forests worldwide (Trumbore, Brando, & Hartmann, 2015). Accordingly, numerous models have projected significant range shifts of forest tree species towards higher latitudes and elevations (Urban et al., 2016). However, to date, the two most important processes in the response of tree populations to a rapidly changing climate, local adaptation and phenotypic plasticity (Aitken, Yeaman, Holliday, Wang, & Curtis-Mclane, 2008; Savolainen, Pyhäjärvi, & Knürr, 2007), are not systematically considered by species distribution models (but see Duputié, Rutschmann, Ronce, & Chuine, 2015; Richardson, Chaney, Shaw, & Still, 2017; Valladares et al., 2014). Phenotypic plasticity enables a given genotype to express different phenotypes in response to changing environments, while local adaptation produces new genotypes with a greater ability to cope with the new environment. The two mechanisms are ubiquitous in natural populations, although their respective importance is considered to vary extensively through time and across species ranges (Des Roches et al., 2018; Reich et al., 2016). To better understand and more realistically predict future species distribution ranges, it is therefore critical to identify and quantify the respective importance of local adaptation and phenotypic plasticity in the response of local populations to a changing climate.

From an ecological perspective, fitness can be associated with several phenotypic traits which directly affect survival and reproduction, creating a fitness landscape (Laughlin, 2018) that allows them to be used to bound species ranges (Benito-Garzón, Ruiz-Benito, & Zavala, 2013; Stahl, Reu, & Wirth, 2014). Fitness-related traits vary across large geographical gradients, mainly depending on how natural selection drove differences among populations in the past. For instance, tree height is generally greatest at the core of a species range and decreases towards its

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

margins (Pedlar & McKenney, 2017). Climate-driven mortality commonly increases towards the driest part of a species range, which is related to drought-induced stress conditions (Benito Garzón et al., 2018). The onset of flushing phenology tends to be delayed towards high latitudes (Duputié et al., 2015) as a consequence of genetic adaptation to late frost and fluctuating photoperiod (Way & Montgomery, 2015). Moreover, traits tend to co-vary across climatic gradients (Laughlin & Messier, 2015). A conspicuous example is the demographic compensation found between survival and growth near range margins (Benito-Garzón et al., 2013; Doak & Morris, 2010; Peterson, Doak, & Morris, 2018). New climatic conditions can result in maladaptation of some populations, which may change intra-specific patterns of trait variation and co-variation across geographical gradients, and eventually, species ranges. For example, increasing temperatures at high-latitude or high-elevation range margins are likely to produce higher growth rates, but they can also induce higher mortality owing to late frosts (Delpierre, Guillemot, Dufrêne, Cecchini, & Nicolas, 2017; Vitasse, Lenz, Hoch, & Korner, 2014). Hence, species ranges are likely to be delimited by the interaction of multiple traits and their responses across environmental gradients (Benito-Garzón et al., 2013; Enquist et al., 2015; Stahl et al., 2014).

Common gardens or provenance tests provide us with the necessary experiments to quantify phenotypic plasticity and local adaptation of fitness-related traits in response to climate (Mátyás, 1999). Models based on reaction norms of phenotypic traits using measurements recorded in common gardens show that: (i) geographic variation in populations' responses to climate is more strongly based on phenotypic plasticity than on local (Benito Garzón, Robson & Hampe, *Under Review*); (ii) phenotypic variation can strongly differ among traits, in particular for tree survival, growth, and flushing phenology - traits that are directly related to fitness and

typically measured in common gardens (Benito Garzón, Alía, Robson, & Zavala, 2011; Duputié et al., 2015; Richardson et al., 2017; Valladares et al., 2014); (iii) as a consequence, predictions of future species ranges are likely to be strongly influenced by the combined response of different fitness-related traits to climate (Laughlin, 2018), but this structured combination of intra-specific multi-trait variation defining species ranges has not been explored with empirical data.

Fagus sylvatica L. (European beech, henceforth "beech") is a widely distributed deciduous broadleaved temperate tree. In some parts of its range, beech has a late flushing strategy to avoid late frosts, which has a generally detrimental effect on tree growth (Delpierre et al., 2017; Gömöry & Paule, 2011; Robson, Rasztovits, Aphalo, Alia, & Aranda, 2013). Beech is currently expanding at its northern distribution edge, whereas it experiences drought-induced radial growth decline and increasing mortality at its southern edge (Farahat & Linderholm, 2018; Stojnic et al., 2018). The extent to which this pattern will continue in the future depends on how the combination of several fitness-related traits will influence the species' response to new climates.

Here, we propose a new modeling approach that quantifies local adaptation and phenotypic plasticity of four major phenotypic traits related to fitness (vertical and radial growth, survival, and flushing phenology) and their interactions, to delimit species ranges under current and future climates. To this purpose, we use the phenotypic measurements recorded in the BeechCOSTe52 database (Robson, Benito Garzón, & BeechCOSTe52 database consortium, 2018), the largest network of common gardens for forest trees in Europe, covering virtually the entire distribution range of the species. Our specific objectives are: (i) to quantify range-wide patterns of phenotypic plasticity and local adaptation in growth, survival and flushing phenology;

(ii) to identify interactions among the different traits and the extent of their geographical variation in local adaptation and phenotypic plasticity; (iii) to discuss how these fitness-related traits delimit species ranges, and (iv) to better understand species ranges under new climate scenarios and the role of trait variation in shaping the future species range.

2. MATERIAL AND METHODS

We calibrated two types of linear mixed-effect models using a combination of trait measurements from common gardens where seeds coming from provenances from different origins have been planted (provenances) and of environmental variables that we obtained for these common gardens and provenances. The first model type (one-trait models) used single traits as response variables and environmental data as explanatory variables. The second model type (two-trait models) added a second trait as co-variate, which allowed the interaction of both traits to be accounted for in the model. Finally, to quantitatively estimate the contribution of each trait to explain beech range, we performed a binomial model using the occurrence of the species as response variable (presence/absence) and the spatial predictions of all traits as explanatory variables.

2.1. Trait measurements

We analyzed total tree height (vertical growth), diameter at breast height (DBH; radial growth), survival and flushing phenology measured on a total of 153,711 individual beech trees that originated from seeds collected from 205 populations (hereafter referred to as "provenances") across Europe and planted at 38 common gardens (hereafter "trials") (Table 1; Supporting

Information Figure S1.1, Appendix S1). Briefly, the seeds were germinated in greenhouses and planted in the trials at an age of two years. Plantations were carried out in two consecutive campaigns, the first campaign (comprising 14 trials) in 1995 and the second one (comprising 24 trials) in 1998 (see more details in Robson et al., 2018). Survival was recorded as individual tree survival. Leaf flushing was transformed from observational stage score data (qualitative measurements that slightly differ among trials) to Julian days by adjusting flushing stages for each tree with the Weibull function by trial (Robson et al., 2011, 2013).

2.2. Environmental data

We used the EuMedClim database that gathers climatic information from 1901 to 2014 gridded at 1km (Fréjaville & Benito Garzón, 2018). The climate of the provenances was averaged for the period from 1901 to 1990, with the rationale that the seeds planted in the common gardens stemmed from trees growing during that period (Leites, Robinson, Rehfeldt, Marshall, & Crookston, 2012). To characterize the climate of the common gardens, we calculated average values for the period between the date of planting (either 1995 or 1998) and the year of measurement of each trait for 21 climate variables (Supporting Information Table S1.1, Appendix S1). In addition, we used the latitude and longitude of the provenance and of the trial as proxies for the photoperiod and continentality, respectively (used in our flushing phenology models).

Phenotypic predictions under future climates were performed using the representative concentration pathway (RCP 8.5) in GISS-E2-R from WorldClim (http://www.worldclim.org/cmip5 30s) for 2070. We deliberately chose only this pessimistic

scenario because for long-lived organisms such as forest trees it makes little difference whether the projected situation will be reached in 2070 or some decades later.

2.3. Statistical analysis

2.3.1. Environmental variable selection

To avoid co-linearity and reduce the number of environmental variables to use in models, we performed two principal component analyses (PCA), one for the climate variables related to the provenance site and one for the climate variables related to the trial site. For tree height, DBH and survival, we considered 21 variables for the provenance and 21 variables for the trial (Supporting Information Figure S1.2, Appendix S1); whereas for leaf flushing, we only included the temperature-related variables as well as latitude and longitude (a total of 20 variables), because leaf flushing is known to be mainly driven by them (Basler & Körner, 2014).

The retained variables after the PCA screening were combined in models containing one variable to characterize the climate of the provenance and one variable to characterize the climate of the trial (Supporting Information Table S1.2, Appendix S1). We selected the combination of environmental variables for each trait that minimize the AIC (Supporting Information Table S1.2, Appendix S1).

2.3.2. One-trait and two-trait mixed-effect models

We used linear mixed-effect models to analyze the response of individual traits (one-trait models) and the co-variation between two traits (two-trait models) to climate. We included the climate at the provenance and the trial site as previously selected (Supplementary Table 1), the

age of trees, and for the leaf flushing model also latitude and longitude as fixed effects. The trial, blocks nested within the trial and trees nested within block and trial, were included as random effects to control for differences among sites and for repeated measurements of the same trees. The random effect of the provenance was also included in the model. The common form of the one-trait model was:

$$log(TR_{ijk}) = \alpha_0 + \alpha_1(Age_{ik}) + \alpha_2(CP_{ij}) + \alpha_3(CT_{ik}) + \alpha_4(CP_{ij}^2) + \alpha_5(CT_{ik}^2)$$
$$+ \alpha_6(Age_{ik} \times CP_{ij}) + \alpha_7(Age_{ik} \times CT_{ik}) + \alpha_8(CP_{ij} \times CT_{ik}) + \beta + \varepsilon$$

208 (Equation 1)

Where TR = trait response of the i^{th} individual of the j^{th} provenance in the k^{th} trial; Age = tree age of the i^{th} individual in the k^{th} trial; CP = climate at the provenance site of the i^{th} individual of the j^{th} provenance; CT = climate at the trial site of the i^{th} individual in the k^{th} trial; β = random effects and ε = residuals. In addition, the model included the following interaction terms: Age and CP, Age and CT, and CP and CT.

We analyzed trait co-variation across the species range by adding two specific traits of interest in the same model. The common form of the two-trait model was:

$$log(TR_{ijk}) = \alpha_0 + \alpha_1(Age_{ik}) + \alpha_2(Cov_{ij}) + \alpha_3(CP_{ij}) + \alpha_4(CT_{ik}) + \alpha_5(Cov_{ik} \times CP_{ij}) +$$

$$\alpha_6(Cov_{ik} \times CT_{ik}) + \alpha_7(Cov_{ik} \times Age_{ik}) + \alpha_8(Age_{ik} \times CP_{ij}) + \alpha_9(Age_{ik} \times CT_{ik}) +$$

$$\alpha_{10}(CP_{ij} \times CT_{ik}) + \beta + \varepsilon$$
219
(Equation 2)

Where TR = trait response of the i^{th} individual of the j^{th} provenance in the k^{th} trial; Age = tree age of the i^{th} individual in the k^{th} trial; Cov = trait co-variate of the i^{th} individual in the k^{th} trial; CP =

climate at the provenance site of the i^{th} individual of the j^{th} provenance; CT = climate at the trial site of the i^{th} individual in the k^{th} trial; β = random effects and ε = residuals. In addition, the model included the following interaction terms: Cov and CP, Cov and CT, Cov and Age, Age and CP, Age and CT, and CP and CT.

The one-trait and two-trait models for vertical and radial growth and leaf flushing were fitted with the 'Imer' function, while the one-trait model for survival was fitted with the 'glmer' function to accommodate logistic regressions (binomial family) in the analysis. We implemented a stepwise-model procedure with four main steps to choose the best supported model (Akaike, 1992): (i) we fitted saturated models that included all the variables in the fixed part of the model; (ii) we chose the optimal random component of the model by comparing the battery of models using restricted maximum likelihood (REML), and selected the best model using the Akaike information criterion (AIC) with criteria delta <2 (Mazerolle, 2006); (iii) we compared the battery of models using maximum likelihood (ML) and selected the optimal fixed component using the AIC criterion; (iv) we combined the best optimal random and fixed component previously selected and adjusted them using REML to obtain the best performing model. All model fits were done using the package 'Ime4' (Bates et al., 2018).

For the best supported models, we visually analyzed the interactions of vertical growth, radial growth, survival and leaf flushing with the environment (one-trait models) and between traits (between the response and co-variate variable, i.e. the two-trait models). To do so, tree age was fixed to 12 years for the radial and vertical growth and leaf flushing models and to 6 years for the survival model. Mathematical interactions in one-trait models (CP x CT in equation 1) represent the differences in trait values that can be attributed to the provenance (interpretable as local adaptation) and those that can be attributed to the trial (interpretable as phenotypic

plasticity). Mathematical interactions in two-trait models (Cov x CT in equation 2) represent the differences in trait values that can be attributed to a second trait that co-varies across the species range with the first trait, mediated by the climate of the trial (representing phenotypic plasticity). Unfortunately, survival could not be included in the two-trait models because there were insufficient measurements shared with other traits in the same trials.

We estimated the percentage of the variance explained by the model attributed to the fixed effects alone (marginal R^2) and attributed to the fixed and random effects together (conditional R^2). We measured the generalization capacity (Pearson correlation) of the model using cross-validation (64% of the data used for calibration and the remaining 34% for validation).

2.3.3. Spatial predictions

We made spatial predictions for each trait across the species range for current and future climatic conditions using the 'raster' package (Hijmans et al., 2017). For the prediction of current and future trait variation, the climate variable for provenance was represented by the average climate over the period from 1901 to 1990. The climate of the trial was set as the average climate from 2000 to 2014, for current trait predictions, and to 2070 for future predictions. For two-trait models, the predicted values of the co-variate (DBH and leaf flushing) in the present were used to estimate the predictions of vertical growth in the future. We calculated the spatial difference between the future and the current conditions (future values minus current values) to illustrate the amount of change that traits can accommodate. All spatial predictions of traits were delimited within the distribution range of the species (EUFORGEN, 2009).

2.3.4. Quantification of the trait contribution to delimit the range of beech

We regressed the occurrence (presence/absence) of the species (EUFORGEN, 2009) against the trait values obtained by the one-trait models using the 'glm' function to accommodate logistic regressions (binomial family). The equation takes the form:

$$(RV) = \alpha_0 + \alpha_1(Vg) + \alpha_2(Rg) + \alpha_3(S) + \alpha_4(Lf) + \alpha_5(Vg \times S) + \alpha_6(Rg \times S) + \alpha_7(Lf \times S)$$
$$+ \alpha_8(Vg \times Rg) + \alpha_9(Vg \times Lf) + \alpha_{10}(Rg \times Lf) + \varepsilon$$

270 (Equation 3)

Where RV = presence/absence of beech; Vg = vertical growth; Rg = radial growth; S = survival;
 Lf = leaf flushing; ε = residuals. In addition, the model included all possible pairwise linear

interactions of the included traits. The total deviance explained by the model was calculated

using the function 'Dsquared' of the package 'modEvA' (Barbosa, Brown, & Real, 2014). Then,

we performed an analysis of variance (ANOVA) of the model to obtain trait and trait interaction

deviances to estimate the percentage of the variance attributable to each trait.

277 All the models where performed with the R statistical framework version 3.2.0 (R Development

Core Team, 2015).

267

268

269

273

274

275

276

278

279

280

281

282

283

284

285

3. RESULTS

3.1. Environmental variables selection

The two PCA performed (provenance PCA and trial PCA) revealed two groups of variables, one related with temperature and another more related with precipitation (Supporting Information Figure S1.2, Appendix S1). The two most important axes of the provenance PCA explained

53.52 and 24.03% of the total variance, and those of the trial PCA explained 38.93 and 24.19% (Supporting Information Figure S1.2, Appendix S1). We to retain the following variables for tree growth and survival: BIO1, BIO5, BIO6, BIO12, BIO13, BIO14, Pet Mean and Pet Max. For the leaf flushing models, we retained BIO1, BIO5, BIO6, MTdjf, MTmam, MTjja, Mtson, and Mtdjfmam in addition to latitude and longitude.

3.2. One-trait and two-trait models

According to the best supported models (Table 2), the most important variable related to the climate at the provenance for vertical growth, radial growth and survival was maximal potential evapotranspiration (Pet Max). The most important variables related to climate at the trials were precipitation of the wettest month (BIO13) for vertical growth, annual precipitation (BIO12) for radial growth, and precipitation of the driest month (BIO14) for survival. In the case of leaf flushing, the mean temperature of December, January and February (MTdjf) was the most important climate variable for both the provenance and the trial site. The latitude of the provenance and the trial and the longitude of the trial were also significant in the leaf flushing model (see Supporting Information Table S1.3, Appendix S1 for detailed statistics on the models). We observed significant interactions between the climate of the trial and that of the provenance in all models (Table 2; Supporting Information Table S1.3, Appendix S1).

The capacity for generalization from the models (Pearson correlation coefficients) was high: between 0.53 for radial growth and 0.73 for leaf flushing. The marginal R^2 ranged from 18% for the survival model to 57% for the vertical growth model, while the conditional R^2 ranged from

40% for the survival model to 98% for the radial growth model (Supporting Information Table S1.3, Appendix S1).

The significance of the fixed and random effects in the one-trait models was positively affected (i.e., estimates were higher) by the addition of a second trait (Supporting Information Table S1.4, Appendix S1). Furthermore, the co-variates and their interactions with the climate variables of the trials were also significant in the two-trait models (Supporting Information Table S1.4, Appendix S1). The capacity to generalize from the two-trait models was high: 0.76 for the vertical growth-radial growth model and 0.77 for the vertical growth-leaf flushing model (Supporting Information Table S1.4, Appendix S1). The marginal R^2 was 62% in the vertical growth-radial growth model and 47% in the vertical growth-leaf flushing model, while the conditional R^2 was 95% in the vertical growth-radial growth model and 99% in the vertical growth-leaf flushing model (Supporting Information Table S1.4, Appendix S1).

3.3. Spatial patterns of phenotypic trait variation from one-trait models

Spatial predictions showed differences in phenotypic trait variation among traits (Figure 1, maps) and the interaction graphs permitted the way that plasticity and local adaptation shape these differences to be visualized (Figure 1, interaction graphs).

Vertical growth reached its maximum value at intermediate values of precipitation of the wettest month in the trials (Figure 1a, interaction graph). These largest trees were predicted to occur mostly over the northern and western part of the species range (Figure 1a, map). A signal of local adaptation was detected in our models and is shown by the interaction graph, where each

line represents the response of provenances to high, intermediate and low levels of maximal potential evapotranspiration.

Predicted radial growth across the species range presented a similar pattern to that of vertical growth, with the lowest values in marginal populations, particularly at the southern margin (Figure 1b, map). High annual precipitation coincided with high growth rates (Figure 1b map), with a moderate signal of local adaptation in the form of some variation among provenances (Figure 1b, interaction graph).

The lowest survival rates were predicted towards the east and at some isolated points in the southernmost part of the range (Figure 1c, map). Survival increased towards those trials where precipitation is high in the driest month, with weak local adaptation indicated by very small –though statistically significant– differences among provenances (Figure 1c, interaction graph).

Earlier flushing was predicted towards the south-eastern part of the range (Fig 1d, map), with notable local adaptation indicated by large differences among provenances depending on the latitude of origin (Figure 1d, interaction graph). Differences in flushing date among provenances were particularly large in trials where the winter temperature is low (Figure 1d, interaction graph).

3.3. Patterns of phenotypic trait variation from two-trait models

Overall, models with a second trait as co-variate produced different results to those considering a single trait only. Predicted vertical growth was larger across the range when either radial growth (Figure 2a) or leaf flushing (Figure 2b) was included as a co-variate. Vertical growth increased

with radial growth and precipitation (Figure 2a) and decreased in those regions where leaf flushing was predicted to be late in the year (which corresponded mainly to the northern part of the range) (Figure 2b).

3.4. Spatial predictions of traits under climate change considering one- and two-trait

models

Trait projections for 2070 showed an overall increase in tree growth, particularly for radial growth (Figure 3a, b), but following similar spatial patterns to those predicted under current conditions (Figure 1a, b). Tree survival was predicted to strongly decrease (with respect to that predicted under current conditions, Figure 1c) in the east and throughout the range periphery, while survival rates remained higher in the central part (Figure 3c). Leaf flushing showed similar patterns to those predicted under current conditions (Figure 1d) but with an overall advance in flushing dates (Figure 3d).

The prediction of vertical growth, considering radial growth as a covariate, showed an overall increase across the distribution range (Figure 3e) with respect to the model projection of vertical growth under future conditions (Figure 3a). This model showed differences in vertical growth increase compared to the same model applied to current conditions (Figure 2a; Supporting Information Figure S1.3e, Appendix S1). Predictions considering leaf flushing as a co-variate tended to constrain vertical growth throughout the range (Figure 3f) compared with the same model in current conditions (Figure 2b).

3.5. Total trait contribution to explain species ranges

All traits and their interactions significantly contributed to explain species occurrence (Table 3). The model explained 31% of the total deviance, with vertical growth accounting for 37%, radial growth for 33%, survival for 19%, and leaf flushing for 1%. The interaction between vertical growth and survival contributed with 3% to the total deviance, that between radial growth and leaf flushing with 2% and the remaining interactions with 1% or less (Table 3).

4. DISCUSSION

4.1. Contribution of phenotypic plasticity and local adaptation to range-wide variation in

beech growth, survival and leaf flushing

Altogether, our results underpin that range-wide variation in fitness-related traits of beech is driven markedly more by phenotypic plasticity than by local adaptation (Supporting Information Table S1.3, Appendix S1). as happens in other species (Benito Garzón *et al.*, *Under review*), and they imply that beech possesses a noteworthy capacity to respond to rapid climate change through acclimation. Although a short-term response through acclimation can be considered as positive for beech to keep pace with climate change, our results point out that the plastic component of tree growth and survival is mostly related to precipitation (Table 2), which follows highly unpredictable patterns (Pflug et al., 2018), making it difficult to evaluate whether acclimation will be enough for beeches to survive (but see our predictions for 2070 under RCP 8.5. showing an increase of mortality in young trees at the margins of the species ranges – Figure 3c). Local adaptation in tree growth (vertical and radial) and survival are driven by adaptation to

maximal potential evapotranspiration (Table 2), suggesting that populations are responding to selection factors related to drought (Volaire, 2018). This is in agreement with the general consensus that beech is a drought-sensitive species (Aranda et al., 2015), although there is ongoing debate over the extent of resistance that beech has to drought (Pflug et al., 2018).

The plastic response of leaf flushing to climate was mainly driven by winter temperatures (Table 2). There is a general consensus that these will increase globally in the future (Vautard et al., 2014), and, accordingly, our projection for 2070 anticipates an advance in flushing through most of the range (Figure 1d, 3d and S3d). However, leaf flushing can be constrained by local adaptation to photoperiod (Gauzere et al., 2017; Way & Montgomery, 2015). The fact that phenotypic plasticity and local adaptation in leaf flushing are driven by different environmental parameters implies that these two processes would interact in the long-term. For instance, phenotypic plasticity concerning winter temperatures might enhance local adaptation towards new photoperiodical cues (i.e., shorter spring days), but the evolutionary time scale of local adaptation makes this interaction very unlikely in the short-term.

4.2 Trait relationships across the species range

Trait inter-dependence varied to some extent along geographical gradients as the two-trait models had higher predictive power and explained more variance than those based on a single trait (Supporting Information Table S1.3 and Si1.4, Appendix S1). The tight albeit not perfect positive interaction between tree vertical and radial growth (Figure 2a, interaction graph) is unsurprising because of allometric relationships between these two variables, particularly in a common-garden plantation that avoids competition among trees.

The biological basis of the observed co-variation between vertical growth and leaf flushing is less obvious. One possible explanation is that vertical growth is greatly restricted by late flushing in northern beech populations (Kollas, Körner, & Randin, 2014). This would also explain our observation that the one-trait model predicts taller trees to occur in the North, whereas the two-trait model predicts just the opposite. Interestingly, the two-trait model thus implies that strong local adaptation of leaf flushing to photoperiod tends to constrain phenotypic plasticity for vertical growth in northern beech populations (Way & Montgomery, 2015).

4.3. Are spatial patterns of growth, survival and leaf flushing delimiting the range of

beech?

Beech populations from certain eastern and southern parts of the distribution range seem most sensitive to perturbation, as suggested by the lowest values for all traits considered (Figure 1). Our analysis of species occurrence as a function of spatial trait values also suggests that a combination of these traits contributes differently to the delimitation of the species range (Table 3), in particular: (i) mortality delimits certain parts of the southern and eastern range of beech, reflecting the climatic marginality of the species in these areas, and meaning that these populations are most threatened and making eastwards expansion of beech difficult (although more studies on regeneration are needed to confirm this result); this is the case for many species whose highest mortality is in the driest part of their range (Anderegg et al., 2015; Benito-Garzón et al., 2013; Camarero, Gazol, Sancho-Benages, & Sangüesa-Barreda, 2015); (ii) the smallest girths are predicted in the southern part of the distribution and the eastern part of the range, suggesting that radial growth is mostly restricted by drought (interaction graph and map, Figure 1b), as has already been pointed out (Farahat & Linderholm, 2018); (iii) with very little variation

across climatic gradients, vertical growth alone will not delimit beech range. This is not the case for other tree species, for which tree height is clearly delimiting species range (Chakraborty, Schueler, Lexer, & Wang, 2018), highlighting the fact that no single best trait delimits tree species ranges; (iv) projections of trees growing in southern and south-eastern regions that flush early also have higher mortality and lower growth predictions than elsewhere within the species range. However, when tree height and leaf flushing are pooled together in the two-trait model, this leads to an decrease in vertical growth in the North; (v) it seems that in beech, and likely in other species with local adaptation to photoperiod, phenology could restrict the northern expansion of ranges (Duputié et al., 2015; Saltré, Duputié, Gaucherel, & Chuine, 2015), although the link between phenology, survival and fitness is still unclear, and more experiments are needed to better understand the interaction between photoperiod and phenology.

4.4. Implications of using trait approaches based on phenotypic variation to forecast beech

sensitivity to climate change

Overall, spatial patterns of vertical and radial growth, survival and leaf flushing predicted for the future (Figure 3), are relatively similar to those predicted by the models under current conditions (Figure 1 & 2), which is likely due to the high plasticity of these traits that allows populations to respond to short-term changes in the environment. Our results, based on the study of phenotypic variation, predict species persistence in the future (through high trait values) rather than extinction and migration northwards as predicted by species distribution models based on the occurrence of the species (Kramer et al., 2010; Stojnic et al., 2018).

Nevertheless, the direct comparison of our trait predictions for current and future conditions allows us to detect some differences in their spatial patterns and total trait values (Supporting Information Figure S1.3, Appendix S1), and gives us a better understanding of the temporal dynamics of traits and their relative importance for beech persistence in the future. For instance, our models of leaf flushing predict reduced geographical variability in phenology in the future (from day 94 to 160 -Figure 1d- and from day 94 to 147 – Figure 3d-), as has been reported worldwide (Ma, Huang, Hänninen, & Berninger, 2018). This is mostly explained by larger advances in the phenology of populations at colder sites than those at warmer sites, likely as a consequence of the larger increases in winter temperatures that happen in the North (Kjellström et al., 2018). Survival of young trees is predicted to decrease at the margins of the distribution, but less markedly than is predicted by species distribution models (Kramer et al., 2010; Stojnic et al., 2018).

Including more than one trait related to growth likely reflects a conserved allometric relationship between vertical and radial growth in the future (Figure 3e), but this may be a direct consequence of the lack of competition among trees in our experimental design. Including phenology in two-trait models seems to be detrimental for vertical growth, at least for northern populations where growth is likely constrained by phenology (Figure 3f). However, our trait covariation approaches are limited to vertical growth as response variables, limiting our understanding of the interplay that other traits can have across species range in the future.

4.5. Limitations, perspectives and future research

Although this study relied on the largest network of common gardens for a forest tree in Europe, the resulting inferences suffer from a number of limitations. Our models are based in a limited set of ages (from 2 to 15 years old). However, the expression of phenotypic plasticity changes with age (Mitchell & Bakker, 2014), which can restrict the broad scope of our results to those ages that we considered. This limitation is particularly pronounced for the case of survival (age range 2 to 6 years), for which data only reflect early recruit survival. Other relevant proxies for tree fitness as fecundity and reproduction have not been considered in our approach. In beech, climate warming tends to increase seed production in northern populations (Drobyshev et al., 2010) and to cause a decline in seedling density in southern ones (Barbeta, Peñuelas, Ogaya, & Jump, 2011), which would be expected to continue under climate change.

Our predictions should help to shape future controlled experiments on those populations most sensitive to climate (in the South – East of the range), and others designed to test those trait relationships that are still unclear (phenology – growth – mortality) at the northernmost distribution edge.

REFERENCES

Aitken, S. N., Yeaman, S., Holliday, J. A., Wang, T., & Curtis-Mclane, S. (2008). Adaptation , migration

or extirpation : climate change outcomes for tree populations. *Evolutionary Applications*, (1), 95–

111. https://doi.org/10.1111/j.1752-4571.2007.00013.x

Akaike, H. (1992). Data analysis by statistical models. *No To Hattatsu*, 24, 127–133.

Anderegg, W. R. L., Flint, A., Huang, C., Flint, L., Berry, J. A., Davis, F. W., ... Field, C. B. (2015). Tree

mortality predicted from drought-induced vascular damage. *Nature Geoscience*, 8, 367–371.

505

506

507

508

509

510

511

512

513

514

515

516

517

518

519

520

521

522

523

524

525

526

https://doi.org/10.1038/NGEO2400 Aranda, I., Cano, F. J., Gascó, A., Cochard, H., Nardini, A., Mancha, J. A., ... Sánchez-Gómez, D. (2015). Variation in photosynthetic performance and hydraulic architecture across European beech (Fagus sylvatica L.) populations supports the case for local adaptation to water stress. Tree Physiology, 35(1), 34–46. https://doi.org/10.1093/treephys/tpu101 Barbeta, A., Peñuelas, J., Ogaya, R., & Jump, A. S. (2011). Reduced tree health and seedling production in fragmented Fagus sylvatica forest patches in the Montseny Mountains (NE Spain). Forest Ecology and Management, 261(11), 2029–2037. https://doi.org/10.1016/j.foreco.2011.02.029 Barbosa, A. M., Brown, J. A., & Real, R. (2014). modEvA – an R package for model evaluation and analysis. R package, version 0.1. Available at: http://modeva.r-forge.r-project.org/. Basler, D., & Körner, C. (2014). Photoperiod and temperature responses of bud swelling and bud burst in four temperate forest tree species. Tree Physiology, 34(4), 377–388. https://doi.org/10.1093/treephys/tpu021 Bates, D., Maechler, M., Bolker, B., Walker, S., Bojesen, R. H., Singmann, H., ... Green, P. (2018). lme4: Linear mixed-effects models using Eigen and S4. R package version 1.1-18-1. Available at: http://CRAN.R-project. org/package=lme4. Benito-Garzón, M., Ruiz-Benito, P., & Zavala, M. A. (2013). Interspecific differences in tree growth and mortality responses to environmental drivers determine potential species distributional limits in Iberian forests. Global Ecology and Biogeography, 22(10), 1141–1151. https://doi.org/10.1111/geb.12075 Benito Garzón, M., Alía, R., Robson, T. M., & Zavala, M. a. (2011). Intra-specific variability and plasticity influence potential tree species distributions under climate change. Global Ecology and Biogeography, 20, 766–778. https://doi.org/10.1111/j.1466-8238.2010.00646.x

528

529

530

531

532

533

534

535

536

537

538

539

540

541

542

543

544

545

546

547

548

549

Benito Garzón, M., Gonzalez Munoz, N., Wigneron, J.-P., Moisy, C., Fernandez-Manjarres, J., & Delzon, S. (2018). The legacy of water deficit on populations having experienced negative hydraulic safety margin. Global Ecology & Biogeography, 27, 346–356. https://doi.org/10.1111/geb.12701 Camarero, J. J., Gazol, A., Sancho-Benages, S., & Sangüesa-Barreda, G. (2015). Know your limits □? Climate extremes impact the range of Scots pine in unexpected places. Annals of Botany, 116, 917-927. https://doi.org/10.1093/aob/mcv124 Chakraborty, D., Schueler, S., Lexer, M. J., & Wang, T. (2018). Genetic trials improve the transfer of Douglas \square fir distribution models across continents. *Ecography*, 41, 1–14. https://doi.org/10.1111/oik.02629 Delpierre, N., Guillemot, J., Dufrêne, E., Cecchini, S., & Nicolas, M. (2017). Tree phenological ranks repeat from year to year and correlate with growth in temperate deciduous forests. Agricultural and Forest Meteorology, 234–235, 1–10. https://doi.org/10.1016/j.agrformet.2016.12.008 Des Roches, S., Post, D. M., Turley, N. E., Bailey, J. K., Hendry, A. P., Kinnison, M. T., ... Palkovacs, E. P. (2018). The ecological importance of intraspecific variation. *Nature Ecology & Evolution*, 2(1), 57–64. https://doi.org/10.1038/s41559-017-0402-5 Doak, D. F., & Morris, W. F. (2010). Demographic compensation and tipping points in climate-induced range shifts. *Nature*, 467(7318), 959–962. https://doi.org/10.1038/nature09439 Drobyshev, I., Övergaard, R., Saygin, I., Niklasson, M., Hickler, T., Karlsson, M., & Sykes, M. T. (2010). Masting behaviour and dendrochronology of European beech (Fagus sylvatica L.) in southern Sweden. Forest Ecology and Management, 259(11), 2160–2171. https://doi.org/10.1016/j.foreco.2010.01.037 Duputié, A., Rutschmann, A., Ronce, O., & Chuine, I. (2015). Phenological plasticity will not help all species adapt to climate change. Global Change Biology, 21(8), 3062–3073.

551

552

553

554

555

556

557

558

559

560

561

562

563

564

565

566

567

568

569

570

571

572

https://doi.org/10.1111/gcb.12914 Enquist, B. J., Norberg, J., Bonser, S. P., Violle, C., Webb, C. T., Henderson, A., ... Savage, V. M. (2015). Scaling from Traits to Ecosystems: Developing a General Trait Driver Theory via Integrating Trait-Based and Metabolic Scaling Theories. Advances in Ecological Research, 52, 249-318. https://doi.org/10.1016/bs.aecr.2015.02.001 EUFORGEN. (2009). Distribution map of Beech (Fagus sylvatica). Available at: www.euforgen.org. Farahat, E., & Linderholm, H. W. (2018). Growth-climate relationship of European beech at its northern distribution limit. European Journal of Forest Research, 137(5), 1–11. https://doi.org/10.1007/s10342-018-1129-9 Fréjaville, T., & Benito Garzón, M. (2018). The EuMedClim Database □: Yearly Climate Data (1901 – 2014) of 1 km Resolution Grids for Europe and the Mediterranean Basin. Frontiers in Ecology and Evolution, 6, 1–5. https://doi.org/10.3389/fevo.2018.00031 Gauzere, J., Delzon, S., Davi, H., Bonhomme, M., Garcia de Cortazar-Atauri, I., & Chuine, I. (2017). Integrating interactive effects of chilling and photoperiod in phenological process-based models. A case study with two European tree species: Fagus sylvatica and Quercus petraea. Agricultural and Forest Meteorology, 244–245, 9–20. https://doi.org/10.1016/j.agrformet.2017.05.011 Gömöry, D., & Paule, L. (2011). Trade-off between height growth and spring flushing in common beech (Fagus sylvatica L.). Annals of Forest Science, 68(5), 975–984. https://doi.org/10.1007/s13595-011-0103-1 Hijmans, R. J., Van Etten, J., Cheng, J., Mattiuzzi, M., Sumner, M., Greenberg, J. A., ... Ghosh, A. (2017). Package 'raster': Geographic Data Analysis and Modeling. Available at: https://cran.rproject.org/web/packages/raster/raster.pdf. Kjellström, E., Nikulin, G., Strandberg, G., Christensen, O. B., Jacob, D., Keuler, K., ... Vautard, R.

574

575

576

577

578

579

580

581

582

583

584

585

586

587

588

589

590

591

592

593

594

595

(2018). European climate change at global mean temperature increases of 1.5 and 2 degrees C above pre-industrial conditions as simulated by the EURO-CORDEX regional climate models. Earth System Dynamics, 9(2), 459–478. https://doi.org/https://doi.org/10.3929/ethz-a-010180262 Rights Kollas, C., Körner, C., & Randin, C. F. (2014). Spring frost and growing season length co-control the cold range limits of broad-leaved trees. Journal of Biogeography, 41(4), 773–783. https://doi.org/10.1111/jbi.12238 Kramer, K., Degen, B., Buschbom, J., Hickler, T., Thuiller, W., Sykes, M. T., & de Winter, W. (2010). Modelling exploration of the future of European beech (Fagus sylvatica L.) under climate change-Range, abundance, genetic diversity and adaptive response. Forest Ecology and Management, 259(11), 2213–2222. https://doi.org/10.1016/j.foreco.2009.12.023 Laughlin, D. C. (2018). Rugged fitness landscapes and Darwinian demons in trait-based ecology. New Phytologist, 217, 501–503. https://doi.org/10.1111/nph.14908 Laughlin, D. C., & Messier, J. (2015). Fitness of multidimensional phenotypes in dynamic adaptive landscapes. Trends in Ecology & Evolution, 30(8), 487–496. https://doi.org/10.1016/j.tree.2015.06.003 Leites, L. P., Robinson, A. P., Rehfeldt, G. E., Marshall, J. D., & Crookston, N. L. (2012). Height-growth response to changes in climate differ among populations of interior Douglas-fir: a novel analysis of provenance-test data. Ecological Applications, 22(1), 154-165. https://doi.org/10.1890/11-0150.1 Ma, Q., Huang, J., Hänninen, H., & Berninger, F. (2018). Agricultural and Forest Meteorology Reduced geographical variability in spring phenology of temperate trees with recent warming. Agricultural and Forest Meteorology, 256–257, 526–533. https://doi.org/10.1016/j.agrformet.2018.04.012 Mátyás, C. (1999). Forest genetics and sustainability. Dordrecht, Boston: Kluwer Academic Publishers. https://doi.org/DOI: 10.1007/978-94-017-1576-8

597

598

599

600

601

602

603

604

605

606

607

608

609

610

611

612

613

614

615

616

617

618

Mazerolle, M. J. (2006). Improving data analysis in herpetology: Using Akaike's information criterion (AIC) to assess the strength of biological hypotheses. Amphibia Reptilia, 27(2), 169–180. https://doi.org/10.1016/j.jclepro.2013.10.062 Mitchell, R. M., & Bakker, J. D. (2014). Intraspecific Trait Variation Driven by Plasticity and Ontogeny in Hypochaeris radicata. *PLoS ONE*, 9(10). https://doi.org/10.1371/journal.pone.0109870 Pedlar, J. H., & McKenney, D. W. (2017). Assessing the anticipated growth response of northern conifer populations to a warming climate. Scientific Reports, 7, 1–10. https://doi.org/10.1038/srep43881 Peterson, M. L., Doak, D. F., & Morris, W. F. (2018). Both life-history plasticity and local adaptation will shape range-wide responses to climate warming in the tundra plant Silene acaulis. Global Change Biology, 24(4), 1614–1625. https://doi.org/10.1111/gcb.13990 Pflug, E. E., Buchmann, N., Siegwolf, R. T. W., Schaub, M., Rigling, A., & Arend, M. (2018). Resilient Leaf Physiological Response of European Beech (Fagus sylvatica L.) to Summer Drought and Drought Release. Frontiers in Plant Science, 9, 187. https://doi.org/10.3389/fpls.2018.00187 R Development Core Team, R. (2015). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. Avaliable at: http://www.Rproject.org. Reich, P. B., Sendall, K. M., Stefanski, A., Wei, X., Rich, R. L., & Montgomery, R. A. (2016). Boreal and temperate trees show strong acclimation of respiration to warming. *Nature*, 531, 633–636. https://doi.org/10.1038/nature17142 Richardson, B. A., Chaney, L., Shaw, N. L., & Still, S. M. (2017). Will phenotypic plasticity affecting flowering phenology keep pace with climate change? Global Change Biology, 23(6), 2499–2508. https://doi.org/10.1111/gcb.13532 Robson, M., Alia, R., Bozic, G., Clark, J., Forsteuter, M., Gomory, D., ... von Wühlisch, G. (2011). The timing of leaf flush in European beech (Fagus sylvatica L.) saplings. Genetic Resources of European

620

621

622

623

624

625

626

627

628

629

630

631

632

633

634

635

636

637

638

639

640

641

Beech (Fagus Sylvatica L.) for Sustainable Forestry: Proceedings of the COST E52 Final Meeting . *SERIE FORESTAL*, 22, 61–80. Robson, M., Benito Garzón, M., & BeechCOSTe52 database consortium. (2018). Data Descriptor: Phenotypic trait variation measured on European genetic trials of Fagus sylvatica L. Scientific Data, 5, 1–7. https://doi.org/10.1038/sdata.2018.149 Robson, M., Rasztovits, E., Aphalo, P. J., Alia, R., & Aranda, I. (2013). Flushing phenology and fitness of European beech (Fagus sylvatica L.) provenances from a trial in La Rioja, Spain, segregate according to their climate of origin. Agricultural and Forest Meteorology, 180, 76–85. https://doi.org/10.1016/j.agrformet.2013.05.008 Saltré, F., Duputié, A., Gaucherel, C., & Chuine, I. (2015). How climate, migration ability and habitat fragmentation affect the projected future distribution of European beech. Global Change Biology, 21(2), 897–910. https://doi.org/10.1111/gcb.12771 Savolainen, O., Pyhäjärvi, T., & Knürr, T. (2007). Gene Flow and Local Adaptation in Trees, Annual *Review of Ecology, Evolution, and Systematics*, 38(1), 595–619. https://doi.org/10.1146/annurev.ecolsys.38.091206.095646 Stahl, U., Reu, B., & Wirth, C. (2014). Predicting species' range limits from functional traits for the tree flora of North America. Proceedings of the National Academy of Sciences, 111(38), 13739–13744. https://doi.org/10.1073/pnas.1300673111 Stojnic, S., Suchocka, M., Benito-Garzon, M., Torres-Ruiz, J., Cochard, H., Bolte, A., ... Delzon, S. (2018). Variation in xylem vulnerability to embolism in European beech from geographically marginal populations Variation in xylem vulnerability to embolism in European beech from geographically marginal populations. *Tree Physiology*, 38, 173–185. https://doi.org/10.1093/treephys/tpx128

643

644

645

646

647

648

649

650

651

652

653

654

655

656

657

658

659

660

661

662

663

Trumbore, S., Brando, P., & Hartmann, H. (2015). Forest health and global change. Science, 349(6250), 814–818. https://doi.org/10.1126/science.aac6759 Urban, M. C., Bocedi, G., Hendry, A. P., Mihoub, J. B., Pe'er, G., Singer, A., ... Travis, J. M. J. (2016). Improving the forecast for biodiversity under climate change. *Science*, 353(6304). https://doi.org/10.1126/science.aad8466 Valladares, F., Matesanz, S., Guilhaumon, F., Araújo, M. B., Balaguer, L., Benito-Garzón, M., ... Zavala, M. A. (2014). The effects of phenotypic plasticity and local adaptation on forecasts of species range shifts under climate change. Ecology Letters, 17(11), 1351–1364. https://doi.org/10.1111/ele.12348 Vautard, R., Gobiet, A., Sobolowski, S., Kjellström, E., Stegehuis, A., Watkiss, P., ... Jacob, D. (2014). The European climate under a 2 ° C global warming. *Environmental Research Letters*, 9, 1–11. https://doi.org/10.1088/1748-9326/9/3/034006 Vitasse, Y., Lenz, A., Hoch, G., & Korner, C. (2014). Earlier leaf-out rather than difference in freezing resistance puts juvenile trees at greater risk of damage than adult trees. Journal of Ecology, 102, 981–988. https://doi.org/10.1111/1365-2745.12251 Volaire, F. (2018). A unified framework of plant adaptive strategies to drought: crossing scales and disciplines. Global Change Biology (in Press). https://doi.org/10.1111/ijlh.12426 Way, D. A., & Montgomery, R. A. (2015). Photoperiod constraints on tree phenology, performance and migration in a warming world. Plant, Cell & Environment, 38(9), 1725–1736. https://doi.org/10.1111/pce.12431

DATA ACCESSIBILITY All phenotypic data used in this study are available at https://zenodo.org/record/1240931#.XBuSa81CeUk (Robson et al., 2018). All the maps generated in this study are available from the authors.

Tables & Figures

Table 1. The extent of data from the BeechCOSTe52 database used for modelling. Measurements: total number of measurements; Trees: total number of individual trees; Trials: total number of trials; Provenances: total number of provenances, Age: the age at which the trees were measured. Columns indicate sample sizes for the traits: tree height (Height), diameter at breast height (DBH), survival and leaf flushing that were used in the one-trait models, as well as for the combined height-DBH, and height-leaf flushing records that were used in the two-trait models.

Ī						
	Height	DBH	Survival	Leaf flushing	H-DBH	H-Leaf flushing
Measurements	203 105	34 237	41 309	7 863	34 237	12 087
Trees	108 415	31 339	37 433	7 863	31 339	10 634
Trials	36	19	7	7	19	6
Provenances	205	186	114	62	186	150
Age	2 to 15	8 to 15	2 to 6	12	8 to 15	6, 9, 11, 12, 15

Table 2. Summary of the variables included in the final best-supported models (one- and two-trait) for each trait analyzed. Environmental variables selected for the provenances and the trials for the one-trait models of height, DBH, survival and flushing, and for the two-trait models of height-DBH and height-leaf flushing. H: height; DBH: diameter at breast height; Lf: leaf flushing; Pet Max: maximal monthly potential evapotranspiration; BIO12: annual precipitation; BIO13: precipitation of wettest month; BIO14: precipitation of driest month; MTdjf: mean temperature of December, January and February; Co-variate: trait covariate.

		one-trait models					two-trait models	
		Height	DBH	Survival	Leaf flushing	H-DBH	H-Lf	
ables	Environment of the provenance	Pet Max	Pet Max	Pet Max	MTdjf Latitude	Pet Max	Pet Max	
	Environment of the trial	BIO13	BIO12	BIO14	MTdjf Latitude Longitude	BIO13	BIO13	
	Co-variate					DBH	Lf	

Table 3. Summary statistics for a generalized linear model (binomial family) of beech occurrence (presence/absence) as a function of trait spatial predictions and their interactions. Estimate: coefficient of the regression shown on a logarithmic scale; SE: standard error of fixed variables; *t*: Wald statistical test that measures the point estimate divided by the estimate of its standard error, assuming a Gaussian distribution of observations; *p*: p-value; DE: deviance explained; Vg: vertical growth; Rg: radial growth; S: survival; Lf: leaf flushing.

	Estimate	SE	t	p	DE
(Intercept)	-5.84	1.15e-02	-509.03	2.00E-16	
Vg	5.45	1.64e-02	332.93	2.00E-16	0.37
Rg	0.51	7.93e-03	64.67	2.00E-16	0.33
S	2.11	3.75e-03	562.83	2.00E-16	0.19
Lf	3.12	1.48e-02	210.94	2.00E-16	0.01
Vg x S	0.10	4.30e-03	21.08	2.00E-16	0.03
Rg x S	-0.60	2.04e-03	-295.94	2.00E-16	0.01
S x Lf	-1.40	4.02e-03	-348.1	2.00E-16	0.01
Vg x Rg	-1.11	4.62e-03	-240.58	2.00E-16	0.01
Vg x Lf	-7.81	2.15e-02	-363.18	2.00E-16	0.01
Rg x Lf	3.43	1.09e-02	313.89	2.00E-16	0.02
Model total deviance					0.31

Fig 1. Spatial projections for (a) vertical growth (cm), (b) radial growth (mm), (c) survival (probability) and (d) leaf flushing (Julian days) generated using one-trait models (maps on the left), and corresponding graphs of interactions between the best environmental predictor variable across the trials divided according to environment at the provenance for each of the four traits (graphs on the right). Black, dark grey, and light grey lines represent high, medium and low values of the climatic variable of the provenances (as opposed to those of the trial, indicated on the x-axis). The vertical lines represent the confidence intervals. The maps display the trait projection for contemporary climate (inferred from 2000-2014 meteorological data) across the current species range. The color gradient depicts the clinal variation from low (red) to high (green) values of each trait. The values of the different traits are represented in the following way: vertical growth (cm), radial growth (mm), probability of survival (0 =dead, 1=alive) and leaf flushing (Julian days). Pet max prov: maximal monthly potential evapotranspiration at the provenance; Latitude prov: latitude of the provenance.

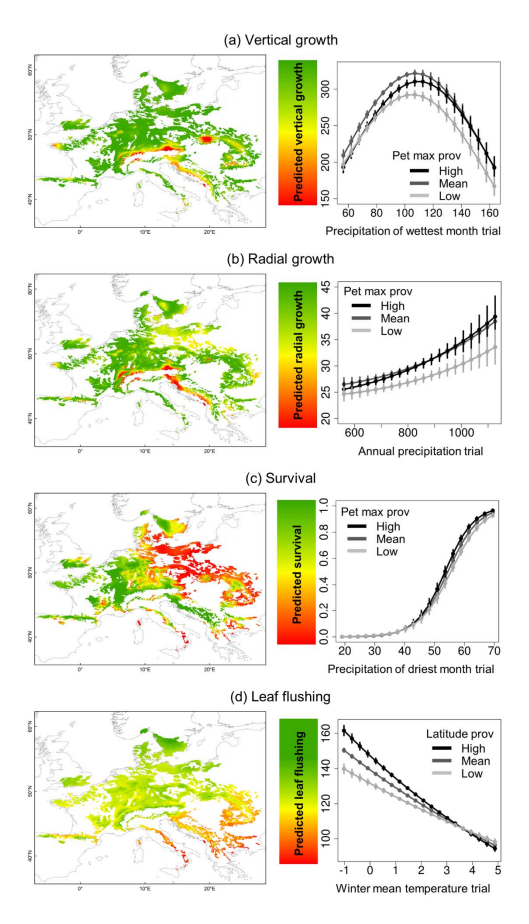
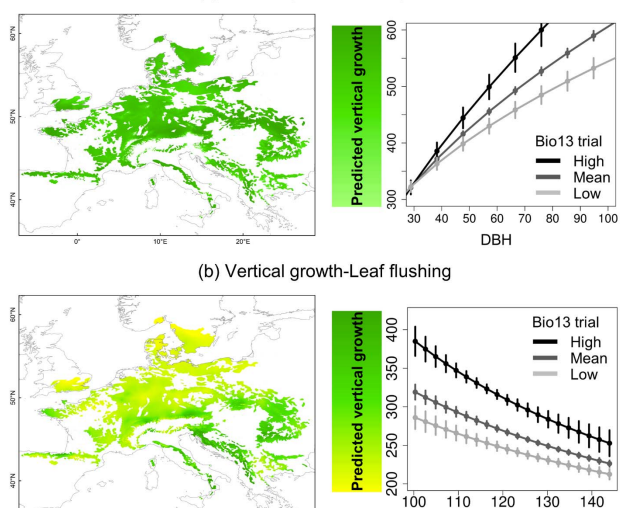


Fig 2. Spatial projections of vertical growth (cm) for (a) Vertical-radial growth model and (b) vertical growth-leaf flushing models (maps on the left), and the corresponding graphs of covariation between vertical growth and the covariate: (a) DBH (mm) and (b) leaf flushing (Julian days). Black, dark grey, and light grey lines represent high, medium and low values of the precipitation of the wettest month of the trial (BIO13). The vertical lines represent the confidence intervals. The maps display the trait projection for contemporary climate (inferred from 2000-2014 meteorological data) across the current species range. The color gradient depicts the clinal variation in vertical growth from 200 cm (yellow) to 600 cm (green).

(a) Vertical growth-radial growth



Leaf flushing

Figure 3. Spatial predictions for 2070 (RCP 8.5) across the species range for one-trait models:

(a) vertical growth (cm); (b) radial growth (mm); (c) probability of survival (0=dead; 1=alive);

(d) leaf flushing (Julian days); and for two-trait models: (e) vertical growth (cm; co-variate radial growth) and (f) vertical growth (cm; co-variate leaf flushing). The color gradients depict the

clinal variation from low (red) to high (green) values.

