1 Title: Responses of competitive understorey species to spatial environmental gradients

2 inaccurately explain temporal changes

3	Authors: Emiel De Lombaerde ^{a,*} , Kris Verheyen ^a , Michael P. Perring ^{a,b} , Markus Bernhardt-							
4	Römermann ^c , Hans Van Calster ^d , Jörg Brunet ^e , Markéta Chudomelová ^{f,g} , Guillaume Decocq ^h , Martin							
5	Diekmann ⁱ , Tomasz Durak ^j , Radim Hédl ^f , Thilo Heinken ^k , Patrick Hommel ^I , Bogdan Jaroszewicz ^m ,							
6	Martin Kopecký ^{n,o} , Jonathan Lenoir ^h , Martin Macek ⁿ , František Máliš ^{p,q} , Fraser J.G. Mitchell ^r , Tobias							
7	Naaf ^s , Miles Newman ^r , Petr Petřík ⁿ , Kamila Reczyńska ^t , Wolfgang Schmidt ^u , Krzysztof Świerkosz ^v ,							
8	Ondřej Vild ^f , Monika Wulf ^s , Lander Baeten ^a							
9	*: Corresponding author. Tel.: +32 9 246 90 30							
10	Email address: emiel.delombaerde@ugent.be (E. De Lombaerde)							
11	^a : Forest & Nature Lab, Campus Gontrode, Faculty of Bioscience Engineering, Ghent University,							
12	Geraardsbergsesteenweg 267, 9090 Melle-Gontrode, BELGIUM (mailing address for							
13	correspondence)							
14	^b : School of Biological Sciences, The University of Western Australia, 35 Stirling Highway, Crawley							
15	WA 6009 AUSTRALIA							
16	^c : Institute of Ecology and Evolution, Friedrich Schiller University, Jena, Dornburger Str. 159, 07743							
17	Jena, GERMANY							
18	^d : Research Institute for Nature and Forest, Kliniekstraat 25, 1070 Brussel, BELGIUM							
19	^e : Southern Swedish Forest Research Centre, Swedish University of Agricultural Sciences, Box 49,							
20	230 53 Alnarp, SWEDEN							
21	^f : Department of Vegetation Ecology, Institute of Botany, The Czech Academy of Sciences, Lidická							
22	25/27, CZ-657 20 Brno, CZECH REPUBLIC							

^g: Department of Botany and Zoology, Faculty of Sciences, Masaryk University, Kotlářská 2, CZ-611

- 24 37 Brno, CZECH REPUBLIC
- 25 ^h: Unité de recherche "Ecologie et Dynamique des Systèmes Anthropisés" (EDYSAN, FRE 3498 CNRS-
- 26 UPJV), Université de Picardie Jules Verne, 1 rue des Louvels, F-80037 Amiens Cedex 1, FRANCE
- ¹: Vegetation Ecology and Conservation Biology, Institute of Ecology, FB 2, University of Bremen,
- 28 Leobener Str. 5, DE-28359 Bremen, GERMANY
- 29 ^j: Department of Ecology, University of Rzeszów, ul. Rejtana 16C, PL-35- 959 Rzeszów, POLAND
- 30 ^k: General Botany, Institute of Biochemistry and Biology, University of Potsdam, Maulbeerallee 3,
- 31 DE-14469 Potsdam, GERMANY
- 32 ¹: Wageningen Environmental Research (Alterra), P.O. Box 47, 6700 AA Wageningen, THE NETHERLANDS
- 33 ^m: Białowieża Geobotanical Station, Faculty of Biology, University of Warsaw, ul. Sportowa 19, 17-
- 34 230 Białowieża, POLAND
- 35 ⁿ: Department of GIS and Remote Sensing, Institute of Botany, The Czech Academy of Sciences,
- 36 Zámek 1, CZ-252 43, Průhonice, CZECH REPUBLIC
- ^o: Department of Forest Ecology, Faculty of Forestry and Wood Sciences, Czech University of Life
 Sciences Prague, Kamýcká 129, CZ-165 00 Prague 6 Suchdol, CZECH REPUBLIC
- ⁹: Technical University in Zvolen, Faculty of Forestry, T. G. Masaryka 24, 960 53 Zvolen, SLOVAKIA
- 40 ^q: National Forest Centre, T. G. Masaryka 22, 960 92 Zvolen, SLOVAKIA
- 41 ^r: Botany Department and Trinity Centre for Biodiversity Research, School of Natural Sciences,
- 42 Trinity College, the University of Dublin, College Green, Dublin 2, IRELAND
- 43 ^s: Leibniz Centre for Agricultural Landscape Research (ZALF), Eberswalder Straße 84, 15374
- 44 Müncheberg, GERMANY

^t: Department of Botany, Faculty of Biological Sciences, University of Wrocław, Kanonia 6/8, PL-50 328 Wrocław, POLAND

⁴⁷ ^u: Department Silviculture and Forest Ecology of the Temperate Zones, Georg-August-University
48 Göttingen, Büsgenweg 1, D-37077 Göttingen, GERMANY

49 ^v: Museum of Natural History, University of Wrocław, Sienkiewicza 21, PL-50-335 Wroclaw, POLAND

50 Abstract

51 Understorey plant communities play a key role in the functioning of forest ecosystems. Under favourable environmental conditions, competitive understorey species may develop high 52 53 abundances and influence important ecosystem processes such as tree regeneration. Thus, 54 understanding and predicting the response of competitive understorey species as a function of 55 changing environmental conditions is important for forest managers. In the absence of sufficient 56 temporal data to quantify actual vegetation changes, space-for-time (SFT) substitution is often used, i.e. studies that use environmental gradients across space to infer vegetation responses to 57 environmental change over time. Here we assess the validity of such SFT approaches and analysed 58 59 36 resurvey studies from ancient forests with low levels of recent disturbances across temperate 60 Europe to assess how six competitive understorey plant species respond to gradients of overstorey 61 cover, soil conditions, atmospheric N deposition and climatic conditions over space and time. The combination of historical and contemporary surveys allows (i) to test if observed contemporary 62 63 patterns across space are consistent at the time of the historical survey, and, crucially, (ii) to assess 64 whether changes in abundance over time given recorded environmental change match expectations 65 from patterns recorded along environmental gradients in space. We found consistent spatial relationships at the two periods: local variation in soil variables and overstorey cover were the best 66 predictors of individual species' cover while interregional variation in coarse-scale variables, i.e. N 67 68 deposition and climate, was less important. However, we found that our SFT approach could not

69 accurately explain the large variation in abundance changes over time. We thus recommend to be

70 cautious when using SFT substitution to infer species responses to temporal changes.

Keywords: temperate forest; herb layer; tree regeneration; global change; nitrogen deposition;
 canopy; spatiotemporal resurvey data; cover abundance; chronosequence; forestREplot

73 Introduction

74 The importance of understorey plant communities and their key role in the functioning of forest ecosystems are increasingly recognized (Gilliam, 2007; Nilsson & Wardle, 2005; Thrippleton, 75 76 Bugmann, Kramer-priewasser, & Snell, 2016). One important influence of the understorey is its 77 effect on tree regeneration; each tree in the overstorey has recruited in and passed through this 78 forest layer as a seedling. Through the initial competitive interactions with the regeneration of 79 overstorey tree species, the understorey community acts as a filter and may have long-term impacts 80 on forest overstorey structure and composition (George & Bazzaz, 1999; Royo & Carson, 2006). Opportunistic, fast-growing understorey plant species develop high abundances when resource 81 82 availability is high, leading to reduced seedling growth and survival, and even complete failure of 83 the tree regeneration (Balandier, Collet, Miller, Reynolds, & Zedaker, 2006; George & Bazzaz, 1999; 84 Royo & Carson, 2006). Thus, it is important for forest managers to understand which (combinations 85 of) environmental factors mainly drive the abundance response of these competitive species.

Understorey species' distribution and abundance are first of all determined by the local-scale environment. The overstorey community can determine the composition and abundance of understorey plants by controlling resources and conditions on the forest floor (Gilliam, 2007; Härdtle, Oheimb, & Westphal, 2003; Li et al., 2012; Nieto-lugilde et al., 2015). Overstorey opening results in increased light availability at the forest floor, but can also improve nutrient and water availability and temperature conditions for understorey plants (Barbier, Gosselin, & Balandier, 2008; Wagner, Fischer, & Huth, 2011). This may lead to a shift in species composition, with a higher cover

93 of light-demanding, competitive species (Degen, Devillez, & Jacquemart, 2005; Kelemen, Mihók, 94 Gálhidy, & Standovár, 2012; Naaf & Wulf, 2007). Understorey species composition and abundance 95 also depend strongly on local soil conditions such as moisture, pH or nutrient availability (Marage & 96 Gégout, 2009; Van Couwenberghe, Collet, Lacombe, & Gégout, 2011; Wagner et al., 2011). In 97 addition to the local site conditions, environmental drivers that vary over broad gradients such as 98 climate and atmospheric nitrogen (N) deposition may be important as well. Coudun and Gégout 99 (2007) found that mean annual temperature predicted the cover of the competitive dominant 100 species Vaccinium myrtillus, in addition to soil acidity and nutrient levels. An experiment by De 101 Frenne et al. (2015) reported that tall, competitive plants, increased in response to elevated 102 temperature, especially under high light availability. Studies on atmospheric N depositions have also 103 documented increasing dominance of fast growing, nitrophytes at the expense of oligotrophic and 104 stress tolerant species (Bobbink et al., 2010; Dirnböck et al., 2014; Gilliam et al., 2016). Integrating 105 all these variables acting at the local and regional scale may thus be very important in predicting 106 understorey species cover along environmental gradients.

107 Environmental conditions in forests are changing over time due to global change and management 108 interventions. Understanding how these environmental changes are causing shifts in the 109 abundances of the species that hamper tree regeneration, preferably requires temporal data 110 (Verheyen et al., 2017). Repeated observations of species abundance are, however, often not 111 available because people have not had the (financial) means or foresight to establish permanent 112 plots or precisely georeference long-term data for particular species or vegetation. Space-for-time 113 (SFT) substitution, which can be broadly defined as using (contemporary) spatial data to infer 114 changes over time, can therefore provide a very useful alternative (Pickett, 1989). In the case of 115 forest understorey vegetation, vegetation inventories covering broad spatial gradients of climate 116 and deposition could be used to understand how competitive understorey species will potentially 117 respond to changing environmental conditions over time (Hedwall & Brunet, 2016). However,

opinions on whether the use of SFT substitution is valid differ and the assumption that drivers of
spatial gradients also drive temporal changes requires validation (Banet & Trexler, 2013; Blois,
Williams, Fitzpatrick, Jackson, & Ferrier, 2013; Johnson & Miyanishi, 2008).

121 In this study, we performed a literature search to identify which understorey plant species are the 122 most important and frequent competitors of tree seedlings in European temperate forests. We then 123 used the data from 36 vegetation resurvey studies from forests across temperate Europe to assess 124 how these competitive understorey species respond to broad gradients of overstorey cover, soil 125 conditions, atmospheric N deposition and climatic conditions over space and time. Each study 126 provides data from historical vegetation records and contemporary resurveys of those records after 127 at least two decades. Our general aim was to assess whether widely available contemporary 128 inventory data can be used to infer likely responses to changing conditions over time. Our main 129 research questions were: (i) which species are considered as most important competitors of tree 130 regeneration in temperate European forests?; (ii) which environmental conditions determine the 131 abundance of these species along spatial gradients and is the relative importance of these drivers 132 consistent across the contemporary and historical vegetation data? and (iii) do species abundance 133 changes over contemporary spatial environmental gradients allow predicting how environmental change over time cause species to actually change their abundances? 134

135 Materials and methods

136 Study species

We performed a formal literature search to identify which understorey species are the most important and frequent competitors of tree seedlings in European deciduous temperate forests (research question 1). We searched for peer-reviewed publications using the bibliographic database of the ISI Web of Knowledge in March 2016 over all available years (1955 to 2016). We used the following search string: Forest* AND tree* AND (seedling* OR sapling* OR regeneration) AND

142 ((understor* OR "ground layer" OR "herb* layer" OR "ground vegetation" OR "ground flora") AND 143 compet*) OR (vegetation AND management) OR (weed*) OR (neighbo*r* AND compet*). Returns 144 from this search were further inspected, and studies that met the following criteria were retained: 145 (i) experimental studies, field experiments, observational field studies or reviews, (ii) studies from 146 European temperate forests and (iii) a forest understorey species is reported to have had a negative 147 impact on the performance of tree seedlings (e.g. their emergence, survival or growth). Studies were 148 initially filtered by title and then by reading the abstracts to retain those studies with possible 149 relevance to the research question. We then examined relevant studies individually and we 150 searched the reference lists for additional publications. Of the 147 studies that we closely examined, 151 57 publications matched our criteria. A search using similar combinations of search terms using 152 Google Scholar, reviewing only the first two pages (sorted by relevance), did not yield additional 153 publications. For each understorey species, we counted the number of publications where it was 154 identified to have a negative effect on tree regeneration (Table 1and Appendix A: Table 1). Although we cannot be certain that we found all available studies, we are confident that our set is a 155 representative selection of species that are considered most important competitors of tree 156 157 regeneration in European temperate forests. Our list of study species was further restricted by 158 including only those that were mentioned to have a negative effect on tree regeneration in at least 159 five publications. Furthermore, species had to be present in more than 100 plots and 10 data sets 160 for both contemporary and historical surveys, that is, retaining species with sufficient data points 161 and spatial coverage. Finally, we excluded species that showed a low cover across almost all plots (percentage cover lower than 15% in over 90% of the plots), because at low cover values they 162 163 probably have a negligible negative effect on regeneration. Ultimately, six species matched these 164 criteria: Deschampsia flexuosa, Molinia caerulea, Pteridium aquilinum, Rubus fruticosus agg., Rubus idaeus and Vaccinium myrtillus (Table 1, bold species). 165

166 Data sets

167 We used the data from 36 independent vegetation resurvey data sets in semi-natural temperate 168 forests across Europe: from Switzerland to southern Sweden (south-north) and from the Ireland to 169 Poland (west-east) (Fig. 1 and Appendix A: Table 2). The data sets included in our analyses cover long/broad gradients of overstorey cover, soil conditions, atmospheric N deposition and climatic 170 171 conditions. Each data set is composed of multiple non-overlapping (in space) permanent or quasipermanent plots recorded at two time points. The historical surveys were carried out between 1935 172 173 and 1994 and the resurveys between 1987 and 2014. Time intervals between the two surveys 174 ranged between 17 and 75 years (38 years on average). The vast majority of plots in these data sets are described as ancient forest sites (sensu Peterken 1996) in which no forest stand replacement 175 176 had taken place between the surveys (e.g. no clear cutting and replanting with conifers). However, 177 management system changes could have taken place without abrupt changes in tree species 178 composition (e.g. gradual transformation from former coppicing to mature forest). Generally, forest management remained stable or became less intensive between the surveys (Bernhardt-179 Römermann et al., 2015). All data sets are included in the forestREplot network 180 (<u>www.forestreplot.ugent.be</u>), a global database combining biodiversity resurveys across temperate 181 182 forests to advance global change research (Verheyen et al., 2017). For further details, see Appendix 183 A (Table 2).

The data sets distinguish between three vegetation layers: the understorey layer (< 0.5-1 m plant height, incl. woody saplings/seedlings), shrub layer (woody plants of minimum height 0.5 to 1 m and maximum height 5 to 14 m) and tree layers. We used cover estimates of each species in each layer as a measure of abundance. Because species cover was recorded in different ways across data sets, cover data were harmonized by converting the different cover recording scales to mid-point percentages of their cover class.

For each species, two subsets of plots were selected: one with the contemporary plots where the species is present (i.e. cover > 0%) and one with the historical plots (Table 1). We also selected species-specific subsets of plot-pairs in which the study species is present at both survey times. The response variable used in our models is the cover percentage of the subject species per plot where it is present.

195 Explanatory variables

196 We used coarse-scale variables expressing gradients in climate and atmospheric N deposition to 197 explain variation in cover among data sets. We used mean annual temperature (MAT; °C) and mean 198 annual precipitation (MAP; mm) to characterize the climatic conditions. Climate data were derived 199 from the Climatic Research Unit at a spatial resolution of 0.5 ° covering monthly means for the 200 period 1901–2013 (Harris, Jones, Osborn, & Lister, 2014). For each data set, we calculated MAT and 201 MAP by averaging annual values for the 10 years preceding the historical surveys and the 202 contemporary resurveys. Nitrogen deposition rate ("Ndep"; kg N/ha/year) was quantified using the 203 EMEP database at a 50-km spatial resolution. We calculated a mean N deposition rate for the period 204 equal to the intercensus interval preceding both the historical survey and resurvey for each data set 205 in a similar way as Verheyen et al. (2012) and Bernhardt-Römermann et al. (2015) using the 206 correction factors provided by Duprè et al. (2010). De Schrijver et al. (2011) showed that the 207 modelled EMEP data and locally observed N deposition data are strongly correlated.

To explain variation in species cover between plots within data sets, we derived plot-level variables related to light availability at the forest floor and soil properties for the two surveys separately. As a measure for light availability at the forest floor, we calculated the total cover of tree and shrub layer species (overstorey cover; "OS") based on species-specific cover values using the approach developed by Fischer (2015). This approach takes into account the overlap between the layers by subtracting the product of the cover values from their sum. As proxies of the prevailing plot-specific

soil properties, we calculated cover-weighted mean Ellenberg indicator values using the individual species' indicator values for soil fertility (EIV_N), soil reaction (EIV_R) and soil moisture (EIV_F) (Diekmann, 2003; Ellenberg, Weber, Düll, Wirth, & Werner, 2001). The study species were excluded from the EIV calculations to avoid circularity. In this study, we used the product EIV_N x EIV_R, which is known to be a good proxy for the turnover rates of organic matter and soil nutrient availability (humus quality; "Hms") (Godefroid, Massant, & Koedam, 2005; Rogister, 1978).

220 Data analysis

221 To quantify which environmental variables determine the dominance of our six study species 222 (research question 2), we related the cover abundance of each species to the plot-level and coarse-223 scale environmental variables using multilevel models to account for the hierarchical structure of 224 the data. First, only the abundance variation along spatial gradients in the contemporary data sets 225 was modelled; these models are henceforth called "spatial models". Models were fitted with the 226 Imer function in the Ime4 package in R 3.4.1 (Bates, Maechler, Bolker, & Walker, 2015; R Core Team, 227 2017). To linearize the relationship between response and explanatory variables and stabilize 228 residual variance, we used a natural logarithmic transformation on the species cover data (Gelman 229 & Hill, 2007). All explanatory variables, measured at different scales, were standardized prior to 230 analysis which results in the estimation of regression coefficients that are comparable in magnitude 231 (Schielzeth, 2010). The parameter estimates of regression coefficients express how the log-cover 232 values change for a one standard deviation change in the explanatory variables. To detect possible 233 multicollinearity between the different explanatory variables, variance inflation factors (VIF) were 234 calculated according to Zuur et al. (2009). These VIF values were low (< 3), indicating low collinearity.

To obtain the most parsimonious model for each species, we started from the "beyond optimal model" (Zuur et al., 2009). This model contained all explanatory variables (Hms, EIV_F, OS, MAT, MAP and Ndep) as fixed effects and a random effect term for 'data set' (Spatial model):

239

Spatial model

In this model, y is the percentage cover of one of the study species at the contemporary survey, x240 241 is one of the six explanatory variables and the random part denotes the effect of 'data set' and 242 residual error. Backward elimination of explanatory variables was done using maximum likelihood-243 fitted models at a 5% level of significance. This procedure was automatized using the StepLmer 244 function (R package ImerTest) with F-tests for parameter estimates calculated using the "Kenward-245 Roger" approximation. For *M. caerulea*, a quadratic term for EIV_F was added after observing a bell 246 shaped pattern in the model residuals. For each species, the most parsimonious model was refitted 247 with restricted maximum likelihood. The goodness of fit for these models were estimated by calculating pseudo R² values following the method of Nakagawa and Schielzeth (2013). R²_{marginal} 248 249 expresses variance explained by fixed effects and R²_{conditional} expresses variance explained by both 250 fixed and random effects.

251 To determine whether the relative importance of the explanatory variables of species abundance 252 was consistent across the contemporary and historical vegetation data, we applied the same set of 253 explanatory variables identified in the most parsimonious spatial models above to the historical 254 survey data. This means we modelled the historical survey cover data of each species in response 255 to the historical survey values of only those explanatory variables that were retained in the models 256 for the contemporary data. By comparing parameter estimates and goodness-of-fit estimates for 257 the models using the historical and resurvey data, we explored to what degree explanatory variables of present-day patterns in cover abundance, especially those retained in the model selection, are 258 also relevant to explain patterns in the old data. 259

Finally, we test if species abundance changes over contemporary spatial environmental gradients allow predicting how environmental change over time cause species to change their abundances

262 (research question 3; see Appendix B for detailed information on our approach). First, we modelled observed changes in species abundance over time, by fitting regression models for each species 263 using the subset of plots in which a species was present at both survey times for each species 264 (Temporal model). For each plot-pair, temporal change was characterized as the natural logarithm 265 of the ratio between the contemporary resurvey and the historical survey. We did this for the 266 response variable (percentage cover), and for all the explanatory variables used in the spatial 267 268 models. Similar as for the spatial models, we used a multilevel modelling approach with random effects for 'data set'. 269

270
$$\ln\left(\frac{y_r}{y_i}\right) = \beta_0^T + \beta_1^T \ln\left(\frac{x_r}{x_i}\right) + \dots + random part$$

271

282

Temporal model

In this model, y_r and y_i are the percentage cover of one of the study species at the contemporary 272 and historical survey resp., x_r and x_i are one of the six explanatory variables at the contemporary 273 and historical survey resp. and random part denotes the effect of 'data set'. The intercept β_0^T 274 275 catches the average temporal change in abundance not accounted for by the used explanatory 276 variables. Second, these temporal models then allowed making predictions of abundance change 277 over time in response to a particular change in environmental conditions. For each predictor variable separately, we predicted the change in species cover (as log ratio) for a realistic change in 278 the predictor (also as log ratio): here we used the observed mean change in the predictor between 279 the two survey times, $\overline{r_{\chi}} = \frac{\overline{x_r - x_l}}{x_r}$ (Table 2). 280

281
$$\widehat{y^T} = \widehat{\ln\left(\frac{y_r}{y_\iota}\right)} = \beta_0^T + \beta_1^T \ln\left(\frac{x_r}{x_\iota}\right) = \beta_0^T + \beta_1^T \ln(1 + \overline{r_x})$$

Temporal prediction

Additionally, predictions were made for the mean change in abundance in the absence of environmental change,

$$285 \qquad \widehat{y^T} = \beta_0^T$$

286

'No change' prediction

Third, we made similar predictions of temporal change in abundance in response to the same environmental change in each predictor $\overline{r_x}$, but based on the species' abundance patterns along spatial gradients, that is, using the contemporary spatial models:

$$290 \quad \widehat{y^{SFT}} = \beta_1 \ln(1 + \overline{r_x})$$

291

Space-for-time prediction

292 In this way, we were able to compare the predicted change in the species' abundances for the actual 293 temporal vs. the spatial model, based on the same change $\overline{r_x}$ in an environmental predictor x. See 294 Appendix B for a more detailed explanation on the predictions using our SFT approach and how 295 these are comparable to the predictions using the temporal models. For the temporal models, 95% 296 confidence intervals were calculated using direct model output, i.e. the mean and standard error of 297 each parameter. For the predictions based on the spatial models, 95% confidence intervals were 298 calculated following an informal Bayesian approach (Gelman & Hill, 2007). For each prediction, we 299 drew 1000 random samples from a normal distribution for the mean and standard error of each 300 model parameter. For each of these samples we were able to calculate the log ratios and compute 301 the confidence intervals around the predictions.

302 Results

303 Spatial models

304 Results of the most parsimonious spatial models using resurvey data (x-axis in Fig. 2) show that local variation in the proxies for soil conditions (moisture and humus quality) and overstorey cover to a 305 306 lesser extent were the best predictors of individual species' cover. Even though covering a broad 307 range in climatic conditions, MAT and MAP were not even retained in the spatial models. Only the 308 interregional variation in N deposition rate had a significant but weak negative impact on the cover 309 of *R. idaeus*. We found that humus quality had a significant and negative effect on all species except 310 for *R. fruticosus* agg. and *R. idaeus*. Soil moisture (EIV_F) had a varying effect on species cover; we 311 observed a negative effect on D. flexuosa, whereas the cover of M. caerulea had a bell-shaped, quadratic response for EIV_F with a maximum between mean Ellenberg values of 6 and 7. Overstorey 312 313 cover was found to have a negative effect on the cover abundance of D. flexuosa, M. caerulea, R. 314 fruticosus agg. and R. idaeus, although the magnitude of the effect was relatively small. A more 315 detailed summary of parameter estimates, significance tests and goodness-of-fit estimations for 316 each species model can be found in Appendix A (Tables 3-8). Overall, the contemporary spatial models fitted the data well, indicated by the $R^{2}_{conditional}$ values ranging from 0.24 up to 0.55 (Fig. 2; 317 Appendix A: Tables 3-8). On average 32% of the total variability was explained by the random 'data 318 set' effect, reflected by the differences between corresponding R²_{marginal} and R²_{conditional} estimations. 319 320 For *R. fruticosus* agg. and *R. idaeus*, only 3% and 4% of variability respectively was explained by the 321 fixed effects (R²_{marginal}=0.03 and 0.04, respectively; Appendix A: Tables 3-8).

Relations between the species' cover abundance and the explanatory variables that were found to be significant in the spatial models using resurvey data were mainly consistent when linking the historical survey abundance data to the historical survey predictor values (Fig. 2; Appendix A: Fig. 1). For all retained explanatory variables, except for overstorey cover and EIV_F for *D. flexuosa* and humus quality for *P. aquilinum*, the confidence intervals of the parameter estimates overlapped with the 1:1 line, indicating that the effects of the predictors were consistent in the spatial models for both surveys. For *D. flexuosa* these effects differed in both magnitude and direction, whereas for *P. aquilinum* only marginally in magnitude. Both $R^2_{marginal}$ and $R^2_{conditional}$ estimates were similar

between the models fitted with resurvey and historical data (Fig. 2; Appendix A: Tables 3-8).

331 Temporal vs. space-for-time approach

332 The models fitted using contemporary spatial data could not accurately explain variation in 333 abundance response of our six study species to changes in the different explanatory variables over 334 time. None of the mean changes in explanatory variables (Table 2), for both temporal models or SFT 335 approach, lead to a significant change in cover, except for the SFT prediction for a mean change in 336 N dep rate for *R. idaeus* (Fig. 3). This shows that even though the effect of a certain predictor variable 337 may be large, a realistic change over time may not cause a significant change in plot-scale species' 338 cover abundance, as was the case here. Similar results were found when making predictions for 339 more extreme change in the explanatory variables, i.e. predictions for the 20% percentile and 80% 340 percentile of the observed change in the plot-pairs between historical and contemporary resurveys 341 (Appendix A: Fig. 2). The SFT approach cannot account for the changes in species' cover over time in absence of environmental change, i.e. β_0^T in the temporal models ('No change'; Fig. 3). The 342 confidence intervals of many predictions using our SFT approach did not overlap with the mean 343 344 prediction using the temporal models. Additionally, uncertainty of the predictions using temporal data was always substantially larger than the predictions using our SFT approach (except for N 345 346 deposition rate). For these reasons, predictions made using our SFT approach could not match the 347 predictions based on actual temporal data.

348 Discussion

Being able to predict how competitive understorey species respond to different environmental drivers can be of key importance in attaining tree regeneration success. In this study, we first identified which understorey species are most important competitors of tree regeneration in European temperate forests. Using a large set of observational data along contemporary spatial

353 gradients in environmental conditions, we then showed that local-scale variables related to light 354 and soil conditions were most important in predicting the abundance of competitive species' 355 abundance. Variation in coarse-scale variables such as climate and nitrogen deposition were less 356 important. Finally, we showed, however, that these contemporary spatial relations could not 357 accurately explain the large variation in species' abundance response over time.

358 Previous studies that analysed the response of individual understorey species cover to overstorey 359 openness observed strong effects (Gaudio, Balandier, & Marquier, 2008; Ricard & Messier, 1996; 360 Van Couwenberghe et al., 2011). In our study, we found, however, that overstorey cover had only a 361 relatively small and negative effect or no significant effect on cover abundance. This might be 362 because D. flexuosa, P. aquilinum, R. fruticosus and V. myrtillus have been shown to be able to 363 tolerate shaded conditions; they may, even under lower light levels, be able to maintain productivity and, especially in the case of R. fruticosus agg., remain dominant in the understorey (Balandier et 364 365 al., 2013; den Ouden, 2000; Van Couwenberghe et al., 2011). Furthermore, the vast majority of plots 366 used in this study are described as ancient forest sites (sensu Peterken, 1996) in which no forest 367 stand replacement had taken place between the surveys (e.g. no large cuttings). Studies using 368 spatial data linking species' abundance to light availability on the forest floor that also include more intensely managed and disturbed forests (e.g. Van Couwenberghe et al., 2011) may therefore show 369 370 stronger effects to overstorey openness.

In contrast to overstorey cover, the variables used as proxies for soil nutrient availability (humus quality) and soil moisture (EIV_F) showed stronger effects on cover abundance. All species except for *R. fruticosus* agg. and *R. idaeus* developed higher cover on sites with low humus quality, which indicates their association as acidophytes with oligotrophic site conditions. This is largely in agreement with previous studies (Coudun & Gégout, 2007; Taylor, Rowland, & Jones, 2001; Van Couwenberghe et al., 2011). It should be noted that *R. fruticosus* agg. is a polymorphic grouping of

377 numerous apomictic microspecies that are phylogenetically very close to each other and difficult to 378 differentiate. It prefers to grow on acid soils, but can grow on a wide variety of soil types (Ellenberg 379 et al., 2001). This can explain why proxies related to soil conditions did not have a significant effect 380 on the cover of *R. fruticosus* agg. The EIV for soil moisture had a negative effect on the abundance 381 of D. flexuosa and we found a bell-shaped, quadratic relation between cover of M. caerulea and the EIV for soil moisture with a maximum at high soil moisture content. This concurs with previous 382 383 studies which indicated that abundance of *M. caerulea* is primarily determined by soil water 384 saturation, soil aeration, and nutrient availability (Ellenberg et al., 2001; Taylor et al., 2001).

385 This data set, covering a large geographical range, gave us a unique opportunity to test the effect of 386 coarse-scale environmental variables on the cover of understorey species. In our results, neither 387 variables related to climate (MAT and MAP) or atmospheric N deposition rate were important in explaining variation in the cover of the study species. Species may simply be indifferent to variation 388 389 in temperature or precipitation, e.g. D. flexuosa, R. idaeus and M. caerulea (Ellenberg et al., 2001). 390 Previous studies have shown that forests with dense overstoreys can potentially buffer the effects 391 of N deposition as well as macroclimate warming on understorey plant communities (De Frenne et 392 al., 2013; Hedwall, Skoglund, & Linder, 2015; Verheyen et al., 2012). This can help to explain why 393 these variables did not have a significant effect in our study. Spatio-temporal resolution of the data 394 on the broad-scale drivers used in this study is relatively coarse and may fail to capture the variation 395 in cover abundance on the local scale. A more detailed characterisation of the environment and 396 measurements on a finer (micro)climatic scale could better explain variation in cover (Lenoir et al., 2013). 397

398 Other (a)biotic factors not included in this study may improve the amount of explained variability 399 for the understorey species' cover. Past land-use can possibly have a strong, underestimated effect 400 on the composition and abundance of temperate forest understoreys (Dupouey, Dambrine, Laffite,

401 & Moares, 2002; Perring et al., 2016; Randin, Jaccard, Vittoz, Yoccoz, & Guisan, 2009). This includes 402 both real changes in land use (e.g. forests on agricultural land) as well as historical transitions in 403 forest management (e.g. coppice to high forest). As the vast majority of the plots included in our 404 study were located in ancient forests, only the latter could be an important factor in this study. Past 405 forest management may have an influence on the composition of understories and abundance of species at present (den Ouden, 2000; Kopecký, Hédl, & Szabó, 2013). Also using data from 406 407 forestREplot, Perring et al. (2018) have shown that understorey community trajectories were clearly 408 influenced by interactions between management legacies from over 200 years ago and 409 environmental change. Yet, detailed data on management history is often lacking and hard to come 410 by. Large herbivores also have a large impact on the abundance of understorey species (Kirby & 411 Thomas, 2000; Rooney, 2001; Vild et al., 2017). Reductions in the cover of species such as Rubus 412 spp. are a common result in grazed woods (Kirby & Thomas, 2000), e.g. under selective browsing by 413 roe deer (Capreolus capreolus; Boulanger et al., 2017; Moser, Schütz, & Hindenlang, 2006). Including 414 detailed information on large herbivores may thus improve models predicting abundance cover. Abundance of these competitive species can also be influenced by the presence of other 415 416 competitors. Hester et al. (1991) found that abundance of V. myrtillus decreased where D. flexuosa 417 was present. Deschampsia flexuosa, in turn, was outcompeted by Agrostis capillaris, however less 418 successfully under shaded conditions. Due to this competitive hierarchy (cf. Boulanger et al., 2017), as a result of asymmetric competition, cover of the 'weaker' competitor may decrease even though 419 420 conditions are favourable. The presence of particular understorey plant species may therefore be 421 important in determining cover abundance response of understorey species to environmental 422 change.

There is no scientific consensus in ecological research on whether or not space-for-time (SFT) substitution is a valid method in predicting change over time. Conclusions from previous studies range from strong support (Banet & Trexler, 2013; Blois et al., 2013; Rolo, Olivier, Guldemond, & 426 van Aarde, 2016; Walker, Wardle, Bardgett, & Clarkson, 2010) to strong rejection (França et al., 2016; Isaac, Girardello, Brereton, & Roy, 2011; Johnson & Miyanishi, 2008). In our study, we show 427 428 that managers and researchers have to be cautious when using spatial data to infer abundance 429 changes of forest understorey species over time. The spatial models fitted using the historical plot 430 data showed a similar relationship between the species cover and the explanatory variables as the 431 models fitted using the resurvey data. This suggests that the contemporary spatial relations we 432 found are consistent at other moments in time and can be used to predict cover abundance over 433 spatial gradients at different points in time. However, by comparing the real observed temporal 434 changes in species abundance in response to environmental change with the predicted changes in 435 abundance based on a SFT, we found that the latter approach could not accurately predict how 436 environmental change over time may cause species to change their abundances. This is mainly due 437 to the fact that understorey species changed considerably in abundance even if changes in the 438 predictors (included in our study) were fairly small. An explanation for these discrepancies is that 439 understorey plant species may not be in equilibrium with the current environment. Observed cover 440 abundance may be lagging behind present environmental conditions and instead reflect past habitat 441 conditions (Bertrand et al., 2011; Dahlgren, Eriksson, Bolmgren, Strindell, & Ehrlen, 2006). This may 442 be due to buffering effect of the overstorey or due to the fact that plants are able to persist under 443 unfavourable conditions (Bertrand et al., 2016) and their lifespan can be as long as several decades 444 (Ehrlén & Lehtilä, 2002). Lauenroth and Sala (1992) showed that the main reason their SFT approach 445 did not match temporal models was due to a temporal lag in the time required for the studied vegetation to capitalize on the amount of precipitation at a given time. Such a time-lagged response 446 447 may also be reflected in the Ellenberg indicator values. The use of direct measurements such as soil 448 pH or soil moisture content (cf. Raduła, Szymura, & Szymura, 2018) instead of indirect values may therefore improve our models. Another reason SFT substitution fails is hysteresis: the rate of 449 450 changes in species' cover-abundance is not the same when the environment shifts from shade to

451 light (fast changes) or when it shifts from light to shade (slow changes). For example, R. fruticosus agg. can rapidly establish and spread in a clearing, but it will take several decades after canopy 452 453 closure before it significantly declines. Additionally, sources of unwanted variability or bias in the 454 used resurvey data may also arise because of relocation errors due to the use of quasi-permanent 455 plots, observer biases and differences in recording seasons (Kapfer et al., 2017; Milberg, Bergstedt, 456 Fridman, Odell, & Westerberg, 2008). These re-sampling errors may add a random error to the 457 temporal change in vegetation, observer-related differences in composition (identification bias) and 458 quantitative properties (abundance bias) among vegetation samples and may result in over- or under-estimation of species abundance. Furthermore, plant species may shift in their responses, 459 460 either across geographical gradients (Diekmann & Lawesson, 1999; Wasof et al., 2013) or shift their 461 niches over time (Pearman, Guisan, Broennimann, & Randin, 2008). This variation may result in 462 inaccuracy of the temporal models and in the mismatch between our SFT approach and temporal models. 463

464 We investigated whether space-for-time substitution allows predicting how species that are 465 considered important competitors for tree regeneration will respond to changing environmental 466 conditions. However, the used data set restricted us from directly modelling tree regeneration in 467 relation to these understorey species and environmental factors. In future research, given adequate 468 data on tree seedlings (e.g. cover or counts) is available, these relationships could be modelled 469 directly, for instance with structural equation modelling (SEM; Grace, Anderson, Olff, & Scheiner, 470 2010). Furthermore, the data used in this study mainly covered ancient forests with low levels of 471 disturbance. Repeating our analyses on an extended data set that also includes more open-canopy 472 forests could give further insights into how understorey species and tree regeneration change 473 abundance under different environmental contexts. Due to the large scale of the data, we were also 474 restricted to using indirect or coarse variables to characterise environmental conditions. In spite of 475 these shortcomings related to data availability, phytosociological vegetation (re)survey data of this

476 sort represents a valuable source of information to improve our understanding of how vegetation changes in relation to environmental gradients over space and time. Quantifying how ecosystems 477 and communities vary along environmental gradients using observational data is a relevant method 478 479 that complements the knowledge gained from experiments and modelling studies (Verheyen et al., 480 2017). We showed in our analyses that spatial (re)survey data can prove valuable in determining 481 which environmental variables affect competitive understorey species cover. Our results suggest, 482 however, that forest ecologists and managers should be cautious when using inventory data across 483 large spatial gradients to predict the impacts of global change. More studies, similar to ours, that 484 simultaneously examine variation in community composition over space, time, and along 485 environmental gradients may clarify under what circumstances using space-for-time substitution, 486 as a tool in ecology, is valid.

487 Acknowledgement

The research of EDL, KV and MPP is supported by the ERC Consolidator Grant 614839 – PASTFORWARD. The research of RH, MM, MK, MC has received funding from long-term research development project no. RVO 67985939 and the Grant Agency of the Czech Republic project 17-092835. FM was funded by Slovak Research and Development Agency, projects APVV-15-0270 and APVV-15-0176. We thank Gian-Reto Walther and Thomas Dirnböck for providing data. We kindly thank Jörg Ewald and an anonymous reviewer for reviewing this document, which has greatly improved through their comments.

495 Appendix A. Supplementary data

496 Appendix B. Detailed information on the temporal vs. space-for-time approach

497 Supplementary data associated with this article can be found, in the online version, at XXXXX.

498 References

Balandier, P., Collet, C., Miller, J. H., Reynolds, P. E., & Zedaker, S. M. (2006). Designing forest
vegetation management strategies based on the mechanisms and dynamics of crop tree
competition by neighbouring vegetation. *Forestry*, *79*(1), 3–27.
http://doi.org/10.1093/forestry/cpi056

Balandier, P., Marquier, A., Casella, E., Kiewitt, A., Coll, L., Wehrlen, L., & Harmer, R. (2013).
Architecture, cover and light interception by bramble (*Rubus fruticosus*): A common
understorey weed in temperate forests. *Forestry*, *86*(1), 39–46.
http://doi.org/10.1093/forestry/cps066

507 Banet, A. I., & Trexler, J. C. (2013). Space-for-time substitution works in Everglades ecological 508 forecasting models. *PLoS ONE*, *8*(11), 1–10. http://doi.org/10.1371/journal.pone.0081025

509 Barbier, S., Gosselin, F., & Balandier, P. (2008). Influence of tree species on understory vegetation

510 diversity and mechanisms involved-A critical review for temperate and boreal forests. *Forest*

511 *Ecology and Management*, 254(1), 1–15. http://doi.org/10.1016/j.foreco.2007.09.038

Bates, D., Maechler, M., Bolker, B. M., & Walker, S. (2015). Fitting Linear Mixed-Effects Models Using
Ime4. *Journal Of Statistical Software*, 67(1), 1–48. http://doi.org/10.18637/jss.v067.i01

514 Bernhardt-Römermann, M., Baeten, L., Craven, D., De Frenne, P., Hédl, R., Lenoir, J., ... Verheyen, K.

515 (2015). Drivers of temporal changes in temperate forest plant diversity vary across spatial

516 scales. *Global Change Biology*, *21*(10), 3726–3737. http://doi.org/10.1111/gcb.12993

517 Bertrand, R., Lenoir, J., Piedallu, C., Riofrio-Dillon, G., de Ruffray, P., Vidal, C., ... Gégout, J.-C. (2011).

518 Changes in plant community composition lag behind climate warming in lowland forests.

519 *Nature*, *479*, 517–20. http://doi.org/doi:10.1038/nature10548

520 Bertrand, R., Riofrío-Dillon, G., Lenoir, J., Drapier, J., De Ruffray, P., Gégout, J. C., & Loreau, M.

521 (2016). Ecological constraints increase the climatic debt in forests. *Nature Communications*, 7.

522 http://doi.org/10.1038/ncomms12643

- 523 Blois, J. L. J. L., Williams, J. W. J. W., Fitzpatrick, M. C. M. C., Jackson, S. T., & Ferrier, S. (2013). Space 524 can substitute for time in predicting climate-change effects on biodiversity. Proceedings of the National Academy of Sciences of the United States of America, 110(23), 9374–9379. 525 526 http://doi.org/10.5061/dryad.d5f1r.1 527 Bobbink, R., Hicks, K., Galloway, J., Spranger, T., Alkemade, R., Ashmore, M., ... De Vries, W. (2010). 528 Global assessment of nitrogen deposition effects on terrestrial plant diversity: a synthesis. 529 *Ecological Applications*, 20(1), 30–59. http://doi.org/10.1890/08-1140.1 530 Boulanger, V., Dupouey, J.-L., Archaux, F., Badeau, V., Baltzinger, C., Chevalier, R., ... Ulrich, E. (2017). 531 Ungulates increase forest plant species richness to the benefit of non-forest specialists. Global Change Biology, (March 2017), 485–495. http://doi.org/10.1111/gcb.13899 532 533 Coudun, C., & Gégout, J. (2007). Quantitative prediction of the distribution and abundance of Vaccinium myrtillus with climatic and edaphic factors. Journal of Vegetation Science, 18(4), 534 517-524. http://doi.org/10.1658/1100-9233(2007)18[517:QPOTDA]2.0.CO;2 535 536 Dahlgren, J. P., Eriksson, O., Bolmgren, K., Strindell, M., & Ehrlen, J. (2006). Specific leaf area as a 537 superior predictor of changes in field layer abundance during forest succession. Journal of
- 538
 Vegetation
 Science,
 17(5),
 577–582.
 http://doi.org/10.1658/1100

 539
 9233(2006)17[577:SLAAAS]2.0.CO;2
 9233(2006)17[577:SLAAAS]2.0.CO;2
- 540 De Frenne, P., Rodríguez-Sánchez, F., Coomes, D. A., Baeten, L., Verstraeten, G., Vellend, M., ...
- 541 Verheyen, K. (2013). Microclimate moderates plant responses to macroclimate warming.
- 542 Proceedings of the National Academy of Sciences of the United States of America, 110(46),
- 543 18561–18565. http://doi.org/10.1073/pnas.1311190110
- 544 De Frenne, P., Rodríguez-Sánchez, F., De Schrijver, A., Coomes, D. a., Hermy, M., Vangansbeke, P.,

- 545 & Verheyen, K. (2015). Light accelerates plant responses to warming. *Nature Plants*, 1(art.
 546 15110), 1–3. http://doi.org/10.1038/nplants.2015.110
- 547 De Schrijver, A., De Frenne, P., Ampoorter, E., Van Nevel, L., Demey, A., Wuyts, K., & Verheyen, K.
 548 (2011). Cumulative nitrogen input drives species loss in terrestrial ecosystems. *Global Ecology*549 *and Biogeography*, 20(6), 803–816. http://doi.org/10.1111/j.1466-8238.2011.00652.x
- Degen, T., Devillez, F., & Jacquemart, A.-L. (2005). Gaps promote plant diversity in beech forests
 (*Luzulo-Fagetum*), North Vosges, France. *Annals of Forest Science*, 62(5), 429–440.
 http://doi.org/10.1051/forest
- den Ouden, J. (2000). *The role of bracken (*Pteridium aquilinum) *in forest dynamics*. Wageningen
 University, the Netherlands.
- Diekmann, M. (2003). Species indicator values as an important tool in applied plant ecology a
 review. *Basic and Applied Ecology*, 4(6), 493–506. http://doi.org/10.1078/1439-1791-00185
- 557 Diekmann, M., & Lawesson, J. E. (1999). Shifts in ecological behaviour of herbaceous forest species

along a transect from Northern Central to North Europe. *Folia Geobotanica*, *34*(1), 127–141.

- 559 Dirnböck, T., Grandin, U., Bernhardt-Römermann, M., Beudert, B., Canullo, R., Forsius, M., ...
- 560 Uzieblo, A. K. (2014). Forest floor vegetation response to nitrogen deposition in Europe. *Global*
- 561 *Change Biology, 20*(2), 429–440. http://doi.org/10.1111/gcb.12440
- 562 Dupouey, J. L., Dambrine, E., Laffite, J. D., & Moares, C. (2002). Irreversible impact of past land use 563 on forest soils and biodiversity. *Ecology*, *83*(11), 2978–2984.
- 564 Duprè, C., Stevens, C. J., Ranke, T., Bleeker, A., Peppler-Lisbach, C., Gowing, D. J. G., ... Diekmann, M.
- 565 (2010). Changes in species richness and composition in European acidic grasslands over the
- past 70 years: The contribution of cumulative atmospheric nitrogen deposition. *Global Change*
- 567 *Biology*, *16*(1), 344–357. http://doi.org/10.1111/j.1365-2486.2009.01982.x

- 568 Ehrlén, J., & Lehtilä, K. (2002). How perennial are perennial plants? *Oikos*, *98*(2), 308–322.
 569 http://doi.org/10.1034/j.1600-0706.2002.980212.x
- 570 Ellenberg, H., Weber, H., Düll, R., Wirth, V., & Werner, W. (2001). *Zeigerwerte von Pflanzen in* 571 *Mitteleuropa. Scripta Geobotanica* (3rd ed.). Goltze, Göttingen.
- Fischer, H. S. (2015). On the combination of species cover values from different vegetation layers.
 Applied Vegetation Science, *18*(1), 169–170. http://doi.org/10.1111/avsc.12130
- 574 França, F., Louzada, J., Korasaki, V., Griffiths, H., Silveira, J. M., Barlow, J., & Nally, R. Mac. (2016).
- 575 Do space-for-time assessments underestimate the impacts of logging on tropical biodiversity?
- 576 An Amazonian case study using dung beetles. *Journal of Applied Ecology*, *53*(4), 1098–1105.
- 577 http://doi.org/10.1111/1365-2664.12657
- 578 Gaudio, N., Balandier, P., & Marquier, A. (2008). Light-dependent development of two competitive
- 579 species (Rubus idaeus, Cytisus scoparius) colonizing gaps in temperate forest. Annals of Forest

580 *Science*, *65*(1), 104. http://doi.org/10.1051/forest

- 581 Gelman, A., & Hill, J. (2007). *Data Analysis Using Regression and Multilevel/Hierarchical Models*.
 582 New York: Cambridge University Press.
- George, L. O., & Bazzaz, F. A. (1999). The fern understory as an ecological filter: emergence and
 establishment of canopy-tree seedlings. *Ecology*, *80*(3), 833–845.
 http://doi.org/10.1890/0012-9658(1999)080[0833:TFUAAE]2.0.CO;2
- 586 Gilliam, F. S. (2007). The ecological significance of the herbaceous layer in temperate forest 587 ecosystems. *Bioscience*, *57*(10), 845–858. http://doi.org/10.1641/B571007
- 588 Gilliam, F. S., Welch, N. T., Phillips, A. H., Billmyer, J. H., Peterjohn, W. T., Fowler, Z. K., ... Adams, M.
- 589 B. (2016). Twenty-five-year response of the herbaceous layer of a temperate hardwood forest
- to elevated nitrogen deposition. *Ecosphere*, 7(4), 1–16. http://doi.org/10.1002/ecs2.1250

- 591 Godefroid, S., Massant, W., & Koedam, N. (2005). Variation in the herb species response and the
- 592 humus quality across a 200-year chronosequence of beech and oak plantations in Belgium.

593 *Ecography*, *28*(2), 223–235. http://doi.org/10.1111/j.0906-7590.2005.03877.x

- Grace, J. B., Anderson, T. M., Olff, H., & Scheiner, S. M. (2010). On the specification of structural
 equation models for ecological systems. *Ecological Monographs*, *80*(1), 67–87.
 http://doi.org/10.1890/09-0464.1
- Härdtle, W., Oheimb, G. Von, & Westphal, C. (2003). The effects of light and soil conditions on the
 species richness of the ground vegetation of deciduous forests in northern Germany
 (Schleswig-Holstein). *Forest Ecology and Management*, 182(1–3), 327–338.
 http://doi.org/10.1016/S0378-1127(03)00091-4
- Harris, I., Jones, P. D., Osborn, T. J., & Lister, D. H. (2014). Updated high-resolution grids of monthly
 climatic observations the CRU TS3.10 Dataset. *International Journal of Climatology*, *34*(3),
 603 623–642. http://doi.org/10.1002/joc.3711
- Hedwall, P. O., & Brunet, J. (2016). Trait variations of ground flora species disentangle the effects of
- global change and altered land-use in Swedish forests during 20 years. *Global Change Biology*,
- 606 22(12), 4038–4047. http://doi.org/10.1111/gcb.13329
- Hedwall, P. O., Skoglund, J., & Linder, S. (2015). Interactions with successional stage and nutrient
 status determines the life-form-specific effects of increased soil temperature on boreal forest
 floor vegetation. *Ecology and Evolution*, *5*(4), 948–960. http://doi.org/10.1002/ece3.1412
- 610 Hester, A. J., Miles, J., & Gimingham, C. H. (1991). Succession from heather moorland to birch
- 611 woodland . II . Growth and competition between *Vaccinium myrtillus* , *Deschampsia flexuosa*
- 612 and Agrostis capillaris. Journal of Ecology, 79(2), 317–327. http://doi.org/10.2307/2260715
- Isaac, N. J. B., Girardello, M., Brereton, T. M., & Roy, D. B. (2011). Butterfly abundance in a warming

- 614 climate : patterns in space and time are not congruent. Journal of Insect Conservation, 15(1–
- 615 2), 233–240. http://doi.org/10.1007/s10841-010-9340-0
- Johnson, E. A., & Miyanishi, K. (2008). Testing the assumptions of chronosequences in succession.

617 *Ecology Letters*, *11*(5), 419–431. http://doi.org/10.1111/j.1461-0248.2008.01173.x

- 618 Kapfer, J., Hédl, R., Jurasinski, G., Kopecký, M., Schei, F. H., & Grytnes, J. (2017). Resurveying
- 619 historical vegetation data opportunities and challenges. Applied Vegetation Science, 20(2),

620 164–171. http://doi.org/10.1111/avsc.12269

- 621 Kelemen, K., Mihók, B., Gálhidy, L., & Standovár, T. (2012). Dynamic response of herbaceous
- 622 vegetation to gap opening in a central European beech stand. *Silva Fennica*, 46(1), 53–65.
- 623 http://doi.org/10.14214/sf.65
- 624 Kirby, K. J., & Thomas, R. C. (2000). Changes in the ground flora in Wytham Woods, southern England
- 625 from 1974 to 1991 implications for nature conservation. *Journal of Vegetation Science*, 11(6),

626 871–880. http://doi.org/10.2307/3236557

- Kopecký, M., Hédl, R., & Szabó, P. (2013). Non-random extinctions dominate plant community
 changes in abandoned coppices. *Journal of Applied Ecology*, *50*(1), 79–87.
 http://doi.org/10.1111/1365-2664.12010
- Lauenroth, W. K., & Sala, O. E. (1992). Long-Term Forage Production of North American Shortgrass
 Steppe. *Ecological Applications*, 2(4), 397–403. http://doi.org/10.2307/1941874
- 632 Lenoir, J., Graae, B. J., Aarrestad, P. A., Alsos, I. G., Armbruster, W. S., Austrheim, G., ... Svenning, J.-
- 633 C. (2013). Local temperatures inferred from plant communities suggest strong spatial buffering
- of climate warming across Northern Europe. *Global Change Biology*, *19*(5), 1470–1481.
- 635 http://doi.org/10.1111/gcb.12129
- Li, M. H., Du, Z., Pan, H. L., Yan, C. F., Xiao, W. F., & Lei, J. P. (2012). Effects of neighboring woody

- 637 plants on target trees with emphasis on effects of understorey shrubs on overstorey physiology
- 638 in forest communities: a mini-review. *Community Ecology*, *13*(1), 117–128.
 639 http://doi.org/10.1556/ComEc.13.2012.1.14
- 640 Marage, D., & Gégout, J. C. (2009). Importance of soil nutrients in the distribution of forest 641 communities on a large geographical scale. *Global Ecology and Biogeography*, *18*(1), 88–97.
- 642 http://doi.org/10.1111/j.1466-8238.2008.00428.x
- Milberg, P., Bergstedt, J., Fridman, J., Odell, G., & Westerberg, L. (2008). Observer bias and random
 variation in vegetation monitoring data. *Journal of Vegetation Science*, *19*(5), 633–644.
 http://doi.org/10.3170/2008-8-18423
- Moser, B., Schütz, M., & Hindenlang, K. E. (2006). Importance of alternative food resources for
 browsing by roe deer on deciduous trees: The role of food availability and species quality.
 Forest Ecology and Management, 226(1–3), 248–255.
 http://doi.org/10.1016/j.foreco.2006.01.045
- Naaf, T., & Wulf, M. (2007). Effects of gap size, light and herbivory on the herb layer vegetation in
 European beech forest gaps. *Forest Ecology and Management*, 244(1–3), 141–149.
 http://doi.org/10.1016/j.foreco.2007.04.020
- Nakagawa, S., & Schielzeth, H. (2013). A general and simple method for obtaining R2 from
 generalized linear mixed-effects models. *Methods in Ecology and Evolution*, 4(2), 133–142.
 http://doi.org/10.1111/j.2041-210x.2012.00261.x
- Nieto-lugilde, D., Lenoir, J., Abdulhak, S., Aeschimann, D., Dullinger, S., Gégout, J., ... Svenning, J. 656 657 (2015). Tree cover at fine and coarse spatial grains interacts with shade tolerance to shape 658 plant species distributions across the Alps. Ecography, 38(6), 578-589. http://doi.org/10.1111/ecog.00954 659

- Nilsson, M. C., & Wardle, D. A. (2005). Understorey vegetation as a forest ecosystem driver:
 evidence from the northern Swedish boreal forest. *Frontiers in Ecology and the Environment*,
 3(8), 421–428. http://doi.org/10.1890/100071
- Pearman, P. B., Guisan, A., Broennimann, O., & Randin, C. F. (2008). Niche dynamics in space and
- 664 time. *Trends in Ecology & Evolution*, 23(3), 149–158. http://doi.org/10.1016/j.tree.2007.11.005
- 665 Perring, M. P., Bernhardt-Römermann, M., Baeten, L., Midolo, G., Blondeel, H., Depauw, L., ...
- 666 Verheyen, K. (2018). Global environmental change effects on plant community composition
- 667 trajectories depend upon management legacies. *Global Change Biology*, (June 2017), 1722–
- 668 1740. http://doi.org/10.1111/gcb.14030
- 669 Perring, M. P., De Frenne, P., Baeten, L., Maes, S. L., Depauw, L., Blondeel, H., ... Verheyen, K. (2016).
- 670 Global environmental change effects on ecosystems: The importance of land-use legacies.

671 *Global Change Biology*, 22(4), 1361–1371. http://doi.org/10.1111/gcb.13146

- 672 Peterken, G. F. (1996). Natural Woodland. Ecology and Conservation in Northern Temperate
 673 Regions. *Cambridge University Press, Cambridge*.
- 674 Pickett, S. (1989). Space-for-time substitution as an alternative to long-term studies. In *Long-Term*
- 675 *Studies in Ecology: Approaches and Alternatives* (pp. 110–135). New York: Springer.
- R Core Team. (2017). R: A language and environment for statistical computing. Vienna, Austria: R
 Foundation for Statistical Computing. Retrieved from https://www.r-project.org/
- 678 Raduła, M. W., Szymura, T. H., & Szymura, M. (2018). Topographic wetness index explains soil
- 679 moisture better than bioindication with Ellenberg's indicator values. *Ecological Indicators*,
- 680 *85*(March 2017), 172–179. http://doi.org/10.1016/j.ecolind.2017.10.011
- Randin, C. F., Jaccard, H., Vittoz, P., Yoccoz, N. G., & Guisan, A. (2009). Land use improves spatial
- 682 predictions of mountain plant abundance but not presence-absence. *Journal of Vegetation*

- 683 Science, 20(6), 996–1008. http://doi.org/10.1111/j.1654-1103.2009.01098.x
- 684 Ricard, J.-P., & Messier, C. (1996). Abundance, growth and allometry of red raspberry (*Rubus idaeus*
- L.) along a natural light gradient in a northern hardwood forest. *Forest Ecology and Management*, *81*(1–3), 153–160. http://doi.org/10.1016/0378-1127(95)03643-1
- 687 Rogister, J. (1978). *De ekologische mR- en mN-waarden van de kruidlaag en de humuskwaliteit van*688 *bosplantengezelschappen*. Groenendaal-Hoeilaart.
- 689 Rolo, V., Olivier, P. I., Guldemond, R. A. R., & van Aarde, R. J. (2016). Validating space-for-time
- 690 substitution in a new-growth coastal dune forest. *Applied Vegetation Science*, *19*(2), 235–243.
- 691 http://doi.org/10.1111/avsc.12210
- Rooney, T. P. (2001). Deer impacts on forest ecosystems: a North American perspective. *Forestry*,
 74(3), 201–208. http://doi.org/10.1093/forestry/74.3.201
- Royo, A. A., & Carson, W. P. (2006). On the formation of dense understory layers in forests
 worldwide: consequences and implications for forest dynamics, biodiversity, and succession.
- 696 *Canadian Journal of Forest Research, 36*(6), 1345–1362. http://doi.org/10.1139/x06-025
- 697 Schielzeth, H. (2010). Simple means to improve the interpretability of regression coefficients.
- 698 Methods in Ecology and Evolution, 1, 103–113. http://doi.org/10.1111/j.2041-
- 699 210X.2010.00012.x
- Taylor, K., Rowland, A. P., & Jones, H. E. (2001). *Molinia caerulea* (L.) Moench. *Journal of Ecology*, *89*(1), 126–144. http://doi.org/10.1046/j.1365-2745.2001.00534.x
- 702 Thrippleton, T., Bugmann, H., Kramer-priewasser, K., & Snell, R. S. (2016). Herbaceous Understorey :
- An Overlooked Player in Forest Landscape Dynamics? *Ecosystems*, 19(7), 1240–1254.
- 704 http://doi.org/10.1007/s10021-016-9999-5

- Van Couwenberghe, R., Collet, C., Lacombe, E., & Gégout, J. (2011). Abundance response of western
 European forest species along canopy openness and soil pH gradients. *Forest Ecology and Management*, 262(8), 1483–1490. http://doi.org/10.1016/j.foreco.2011.06.049
- Verheyen, K., Baeten, L., De Frenne, P., Bernhardt-Römermann, M., Brunet, J., Cornelis, J., ...
 Verstraeten, G. (2012). Driving factors behind the eutrophication signal in understorey plant
 communities of deciduous temperate forests. *Journal of Ecology*, *100*(2), 352–365.
 http://doi.org/10.1111/j.1365-2745.2011.01928.x
- Verheyen, K., De Frenne, P., Baeten, L., Waller, D. M., Hédl, R., Perring, M. P., ... BernhardtRömermann, M. (2017). Combining biodiversity resurveys across regions to advance global

714 change research. *BioScience*, 67(1), 73–83. http://doi.org/10.1093/biosci/biw150

- Vild, O., Hédl, R., Kopecký, M., Szabó, P., Suchánková, S., & Zouhar, V. (2017). The paradox of long term ungulate impact : increase of plant species richness in a temperate forest. *Applied Vegetation Science*, *20*(2), 282–292. http://doi.org/10.1111/avsc.12289
- Wagner, S., Fischer, H., & Huth, F. (2011). Canopy effects on vegetation caused by harvesting and
 regeneration treatments. *European Journal of Forest Research*, *130*(1), 17–40.
 http://doi.org/10.1007/s10342-010-0378-z
- Walker, L. R., Wardle, D. A., Bardgett, R. D., & Clarkson, B. D. (2010). The use of chronosequences in
 studies of ecological succession and soil development. *Journal of Ecology*, *98*(4), 725–736.
 http://doi.org/10.1111/j.1365-2745.2010.01664.x
- Wasof, S., Lenoir, J., Gallet-Moron, E., Jamoneau, A., Brunet, J., Cousins, S. a. O., ... Decocq, G. (2013).
 Ecological niche shifts of understorey plants along a latitudinal gradient of temperate forests
 in north-western Europe. *Global Ecology and Biogeography*, *22*(10), 1130–1140.
 http://doi.org/10.1111/geb.12073

- Weber, H. (2002). Zeigerwerte der Rubus-Arten. In H. Ellenberg, H. Weber, R. Düll, V. Wirth, & W.
 Werner (Eds.), *Zeigerwerte von Pflanzen in Mitteleuropa* (3rd ed., pp. 167–174). Göttingen: E.
 Goltze & Co.
- Zuur, A. F., Ieno, E. N., Walker, N. J., Saveliev, A. A., & Smith, G. M. (2009). *Mixed Effect Models and Extensions in Ecology with R.* New York, USA: Springer. http://doi.org/10.1007/978-0-38787458-6

734 Figure captions

Fig. 1: Map showing the location of the 36 data sets included in this study (the numbers refer to
data set IDs in Table S2). All data sets are included in the forestREplot network, a global database of
understorey resurvey plots from temperate forests.

738 Fig. 2: Results of species-specific models predicting the cover abundance of six understorey plant species in response to spatial gradients in local-scale and large-scale explanatory variables. Points 739 740 representation of the estimated effects (+/- 95% CI) of the explanatory variables for the spatial models using contemporary resurvey data (x-axis) and historical survey data (y-axis); only the 741 742 explanatory variables that were retained after model selection are shown. The line represents the 743 1:1 line of no difference. Different shapes represent the six study species and colours the different explanatory variables. In parentheses R²_{conditional} values for historical and resurvey models are given, 744 745 respectively.

Fig. 3: Predicted cover abundance change of six understorey plant species in response to temporal changes in several environmental explanatory variables. Predictions were based on species abundance changes along spatial environmental gradients (contemporary spatial models) and on actually observed temporal changes in abundance. For each explanatory variable separately, we predicted the change in species cover (as log ratio) to the observed mean change in that predictor

between the two survey times. Additionally, predictions of cover change were made in the absence of environmental change (No change). The dotted line represents these predictions using the temporal models. Explanatory variables that were retained in the most parsimonious contemporary spatial models are annotated in grey. The lines indicate the 95% confidence intervals around the predictions. Red: temporal predictions; Blue: predictions using SFT approach. Humus quality (Hms), Ellenberg F value (EIVF), Overstorey cover (OS), Mean annual temperature (MAT), Mean annual precipitation (MAP) and mean N deposition rate (Ndep).





759 Fig. 1







764 Fig.3 (colour)

- 765 Table 1: Results of a literature review, showing the understorey species that are reported in > 5
- publications as having negative effects on tree regeneration. Species in bold were selected for this
- study based on their presence and abundance in our data.

Species	# Publications	# Resurvey plots [# data sets]	# Historical plots [# data sets]
Rubus fruticosus agg.	20	651 [27]	502 [24]
Deschampsia flexuosa	12	246 [21]	274 [19]
Rubus idaeus	8	403 [29]	365 [21]
Pteridium aquilinum	8	228 [19]	228 [21]
Molinia caerulea	7	139 [16]	135 [16]
Agrostis capillaris	7	87 [11]	78 [14]
Deschampsia cespitosa	7	359 [27]	406 [29]
Vaccinium myrtillus	6	266 [20]	310 [20]
Calamagrostis epigeios	6	96 [9]	62 [9]

Table 2: Overview of the environmental conditions in the plots used in this study and how they changed over time. Mean values of all used explanatory variables over the contemporary and historical plots in which at least one study species is present are given. Mean changes between surveys are calculated for the subset of plots in which at least one species is present at both survey times. Values between brackets are the 20th and 80th percentiles of the range of change over time.

	Historical	Contemporary	Mean change (20 th , 80 th) (%)
Hms (EIV _N x EIV _R)	23.16	24.11	12.01 (-12.26, 38.66)
EIV _F	5.20	5.33	1.22 (-4.10, 7.47)
OS (%)	75.18	74.10	-0.73 (-17.15, 20.09)
MAT (°C)	8.65	9.90	15.30 (10.66, 20.49)
MAP (mm)	806.13	810.13	2.55 (-1.26, 7.29)
Ndep (kg/ha/year)	8.47	17.68	214.87 (-10.45, 505.81)

775 Appendix A: Supplementary data

- 776 **Table 1**: Species that are reported in scientific literature to be competitive towards tree seedlings in
- temperate European forests. Ranking according to number of publications.

Species or genus	Publications	#
		Publications
Rubus fruticosus agg.	(1–20)	20
Deschampsia flexuosa	(2, 3, 10, 21–29)	12
Rubus idaeus	(2, 9, 10, 14, 23, 30–32)	9
Pteridium aquilinum	(2, 8, 13, 21, 33–36)	8
Molinia caerulea	(2, 3, 21, 27, 30, 33, 37)	7
Agrostis capillaris	(2, 10, 38–42)	7
Deschampsia cespitosa	(2, 23, 24, 27, 43–45)	7
Vaccinium myrtillus	(2, 3, 21, 26, 46, 47)	6
Calamagrostis epigeios	(2, 3, 10, 20, 21, 27)	6
Epilobium angustifolium	(2, 9, 10, 48)	4
Calluna vulgaris	(2, 3, 33)	3
Holcus lanatus	(28, 38, 39)	3
Lolium perenne	(2, 28, 49)	3
Carex brizoides	(2, 27, 50)	3
Juncus effusus	(2, 3, 10)	3
Festuca rubra	(39, 42, 51)	3
Calamagrostis villosa	(27, 46, 48)	2
Cytisus scoparius	(2, 38)	3
Urtica dioica	(2, 9)	2
Cirsium vulgare	(2, 28)	2
Epilobium ciliatum	(2, 28)	2
Poa annua	(2, 28)	2
Holcus mollis	(10, 38)	2
Agrostis stolonifera	(43, 45)	2
Dactylis glomerata	(39, 52)	2
Robinia pseudoacacia	(2, 53)	2
Betula pubescens	(2, 41)	2
Sambucus nigra	(2, 23)	2
Calamagrostis arundinacea	(48)	1
Brachypodium pinnatum	(54)	1
Fagus sylvatica	(55)	1
Allium ursinum	(56)	1
Senecio ovatus	(56)	1
Persicaria maculosa	(28)	1
Rumex obtusifolius	(28)	1
Arrhenatherum elatius	(39)	1
Carex sylvatica	(3)	1
Carex digitata	(3)	1
Vaccinium vitis-idaea	(21)	1
Vaccinium uliginosum	(21)	1
Trientalis europaea	(21)	1
Poa pratensis	(51)	1
Agrostis canina	(51)	1
Poa trivialis	(28)	1

Table 1: continued.

Brachypodium sylvaticum	(2)	1
Chenopodium album	(2)	1
Galium aparine	(2)	1
Conyza canadensis	(2)	1
Rumex crispus	(2)	1
Ulex europaeus	(2)	1
Impatiens glandulifera	(2)	1
Elytrigia repens	(2)	1
Dryopteris filix-mas	(2)	1
Polygonum aviculare	(2)	1
Sinapis arvensis	(2)	1
Rorippa sylvestris	(2)	1
Convolvulus arvensis	(2)	1
Galium aparine	(2)	1
Sonchus arvensis	(2)	1
Cirsium arvense	(2)	1
Artemisia vulgaris	(2)	1
Tripleurospermum maritimum	(2)	1
Clematis vitalba	(2)	1
Phytolacca americana	(2)	1
Adenostyles alliariae	(2)	1
Filipendula ulmaria	(2)	1
Rubus caesius	(2)	1
Amorpha fruticosa	(2)	1
Cynodon dactylon	(2)	1
Sorghum halepense	(2)	1
Hedera helix	(2)	1
Solidago canadensis/gigantea	(2)	1
Athyrium filix-femina	(2)	1
Rubus hirtus	(2)	1
Heracleum mantegazzianum	(2)	1
Arbutus unedo	(2)	1
Erica arborea	(2)	1
Empetrum nigrum	(2)	1
Thlaspi arvense	(2)	1
Equisetum arvense	(2)	1

778

1. Balandier P, et al. (2013) Architecture, cover and light interception by bramble (*Rubus fruticosus*): A

780 common understorey weed in temperate forests. *Forestry* 86(1):39–46.

781 2. Willoughby I, et al. (2009) *Forest vegetation management in Europe: Current practice and future*

782 requirements.

783 3. Dodet M, Collet C, Frochot H, Wehrlen L (2011) Tree regeneration and plant species diversity

784 responses to vegetation control following a major windthrow in mixed broadleaved stands. *Eur J For*

785 *Res* 130(1):41–53.

- Gessler A, Keitel C, Nahm M, Rennenberg H (2004) Water shortage affects the water and nitrogen
 balance in Central European beech forests. *Plant Biol* 6(3):289–298.
- Harmer R, Kiewitt A, Morgan G (2012) Can overstorey retention be used to control bramble (*Rubus fruticosus* L. agg.) during regeneration of forests? *Forestry* 85(1):135–144.
- 790 6. Harmer R, Kiewitt A, Morgan G (2012) Effects of overstorey retention on ash regeneration and
- bramble growth during conversion of a pine plantation to native broadleaved woodland. *Eur J For Res* 131(6):1833–1843.
- 793 7. Harmer R (2011) The effect of plant competition and simulated summer browsing by deer on tree
 794 regeneration. *J Appl Ecol* 38(5):1094–1103.
- Harmer R, Morgan G (2007) Development of *Quercus robur* advance regeneration following canopy
 reduction in an oak woodland. *Forestry* 80(2):137–149.
- 9. Lin N, Bartsch N, Vor T (2014) Long-term effects of gap creation and liming on understory vegetation
 with a focus on tree regeneration in a European beech (*Fagus sylvatica* L.) forest. *Ann For Res*57(2):233–246.
- Petritan IC, von Lüpke B, Petritan AM (2012) Response of planted beech (*Fagus sylvatica* L.) and
 Douglas-fir (*Pseudotsuga menziesii* (Mirb.) Franco) saplings to herbaceous and small shrubs control

802 on clearcuts. *J For Res* 17(2):164–174.

- 803 11. Willoughby IH, Harmer R, Morgan GW, Peace AJ (2013) Triclopyr applied in the winter dormant
- season can give effective control of bramble (*Rubus fruticosus* L. agg.) without damaging young tree
 seedlings or other non-target vegetation. *Forestry* 86(1):59–69.
- 806 12. Harmer R, Kiewitt A, Morgan G, Gill R (2010) Does the development of bramble (*Rubus fruticosus* L.
- 807 agg.) facilitate the growth and establishment of tree seedlings in woodlands by reducing deer
- 808 browsing damage? *Forestry* 83(1):93–102.
- 809 13. Harmer R, Morgan G, Beauchamp K (2011) Restocking with broadleaved species during the
- 810 conversion of *Tsuga heterophylla* plantations to native woodland using natural regeneration. *Eur J*

811 For Res 130(2):161–171.

812 14. Jensen AM, Götmark F, Löf M (2012) Shrubs protect oak seedlings against ungulate browsing in

813 temperate broadleaved forests of conservation interest: A field experiment. For Ecol Manage

814 266:187-193.

- 815 15. Ammer C, Schall P, Wördehoff R, Lamatsch K, Bachmann M (2011) Does tree seedling growth and 816 survival require weeding of Himalayan balsam (Impatiens glandulifera)? Eur J For Res 130(1):107-116.
- 817
- 818 Fotelli MN, Gessler A, Peuke AD, Rennenberg H (2001) Drought affects the competitive interactions 16. 819 between Fagus sylvatica seedlings and an early successional species, Rubus fruticosus: responses of 820 growth, water status and delta13C composition. New Phytol 151(2):427–435.
- 821 Fotelli MN, Rennenberg H, Geßler A (2002) Effects of drought on the competitive interference of an 17.
- 822 early successional species (Rubus fruticosus) on Fagus sylvatica L. seedlings: 15N uptake and
- 823 partitioning, responses of amino acids and other N compounds. *Plant Biol* 4(3):311–320.
- 824 18. Fotelli MN, Rudolph P, Rennenberg H, Geßler A (2005) Irradiance and temperature affect the
- 825 competitive interference of blackberry on the physiology of European beech seedlings. New Phytol 826 165(2):453-462.
- 827 19. Mountford EP, Savill PS, Bebber DP (2006) Patterns of regeneration and ground vegetation
- 828 associated with canopy gaps in a managed beechwood in southern England. Forestry 79(4):389–408.
- 829 20. Kelemen K, Mihók B, Gálhidy L, Standovár T (2012) Dynamic response of herbaceous vegetation to
- 830 gap opening in a central European beech stand. Silva Fenn 46(1):53-65.
- 831 21. Dzwonko Z, Loster S, Gawroński S (2015) Impact of fire severity on soil properties and the
- 832 development of tree and shrub species in a Scots pine moist forest site in southern Poland. For Ecol 833 *Manage* 342:56–63.
- 834 22. Jönsson AM, Nihlgård B (2004) Slash pile burning at a Norway spruce clear-cut in Southern Sweden. 835 Water Air Soil Pollut 158(1):127–135.
- 836 23. Löf M, Gemmel P, Nilsson U, Welander NT (1998) The influence of site preparation on growth in
- 837 Quercus robur L. seedlings in a southern Sweden clear-cut and shelterwood. For Ecol Manage 838 109(1-3):241-249.
- 839 24. Mason WL, Edwards C, Hale SE (2004) Survival and early seedling growth of conifers with different

840 shade tolerance in a Sitka spruce spacing trial and relationship to understorey light climate. *Silva*

841 Fenn 38(4):357–370.

- 842 25. Nilsson U, Örlander G (1999) Vegetation management on grass-dominated clearcuts planted with
 843 Norway spruce in southern Sweden. *Can J For Res* 29(7):1015–1026.
- 844 26. Nilsson U, Örlander G (1999) Water uptake by planted Picea abies in relation to competing field
- 845 vegetation and seedling rooting depth on two grass-dominated sites in southern Sweden. *Scand J*846 *For Res* 14(4):312–319.
- Wagner S, Fischer H, Huth F (2011) Canopy effects on vegetation caused by harvesting and
 regeneration treatments. *Eur J For Res* 130(1):17–40.
- 849 28. Willoughby I, Clay D V, Dixon FL, Morgan GW (2006) The effect of competition from different weed
 850 species on the growth of *Betula pendula* seedlings. *Can J For Res* 36(8):1900–1912.
- 29. Jarvis APG (1964) Interference by *Deschampsia flexuosa* (L.) Trin. *Oikos* 15(1):56–78.
- Balandier P, Collet C, Miller JH, Reynolds PE, Zedaker SM (2006) Designing forest vegetation
 management strategies based on the mechanisms and dynamics of crop tree competition by
- 854 neighbouring vegetation. *Forestry* 79(1):3–27.
- 855 31. Mayer P, Abs C, Fischer A (2004) Colonisation by vascular plants after soil disturbance in the

Bavarian Forest - Key factors and relevance for forest dynamics. *For Ecol Manage* 188(1–3):279–289.

- 857 32. Jensen AM, Löf M, Gardiner ES (2011) Effects of above- and below-ground competition from shrubs
- 858 on photosynthesis, transpiration and growth in *Quercus robur* L. seedlings. *Environ Exp Bot*
- 859 71(3):367–375.
- Gaudio N, et al. (2011) Light-mediated influence of three understorey species (*Calluna vulgaris*,
 Pteridium aquilinum, Molinia caerulea) on the growth of *Pinus sylvestris* seedlings. *Eur J For Res* 130(1):77–89.
- 863 34. den Ouden J (2000) The role of bracken (*Pteridium aquilinum*) in forest dynamics. Dissertation
 864 (Wageningen University, the Netherlands).
- 865 35. Collet C, Ningre F, Frochot H (1998) Modifying the microclimate around young oaks through
- 866 vegetation manipulation: Effects on seedling growth and branching. For Ecol Manage 110(1–3):249–

867 262.

868 36. Humphrey J, Swaine M (1997) Factors Affecting the Natural Regeneration of *Quercus* in Scottish
869 Oakwoods . I . Competition from *Pteridium Aquilinum*. J Appl Ecol 34(3):577–584.

870 37. Pages JP, Michalet R (2003) A test of the indirect facilitation model in a temperate hardwood forest
871 of the northern French Alps. *J Ecol* 91(6):932–940.

872 38. Provendier D, Balandier P (2008) Compared effects of competition by grasses (Graminoids) and
873 broom (Cytisus scoparius) on growth and functional traits of beech saplings (Fagus sylvatica). Ann
874 For Sci 65(5):510(1-9).

875 39. Coll L, Balandier P, Picon-Cochard C (2004) Morphological and physiological responses of beech

876 (Fagus sylvatica) seedlings to grass-induced belowground competition. *Tree Physiol* 24(1):45–54.

40. Emmett BA, Anderson JM, Hornung M (1991) Nitrogen sinks following two intensities of harvesting

in a Sitka spruce forest (N. Wales) and the effect on the establishment of the next crop. *For Ecol*

879 *Manage* 41(1–2):81–93.

Kelly DL (2002) The regeneration of Quercus petraea (sessile oak) in southwest Ireland: A 25-year
experimental study. *For Ecol Manage* 166(1–3):207–226.

Vandenberghe C, Freléchoux F, Gandallah F, Buttler A (2006) Competitive effects of herbaceous
vegetation on tree seedling emergence, growth and survival: Does gap size matter? *J Veg Sci*17(4):481–488.

885 43. Collet C, Frochot H (1996) Effects of interspecific competition on periodic shoot elongation in oak
886 seelings. *Can J For Res* 26(11):1934–1942.

44. Chaar H, Colin F, Collet C (1997) Effects of environmental factors on the shoot development of
Quercus petraea seedlings. A methodological approach. *For Ecol Manage* 97(2):119–131.

Kollet C, Frochot H, Guehl J-M (1996) Effect of two forest grasses differing in their growth dynamics
on water relations and the growth of Quercus petraea seedlings. *Can J Bot* 74(10):1555–1561.

46. Kirchner K, Kathke S, Bruelheide H (2011) The interaction of gap age and microsite for herb layer

892 species in a near-natural spruce forest. *J Veg Sci* 22(1):85–95.

893 47. Mirschel F, Zerbe S, Jansen F (2011) Driving factors for natural tree rejuvenation in anthropogenic

894		pine (<i>Pinus sylvestris</i> L.) forests of NE Germany. <i>For Ecol Manage</i> 261(3):683–694.
895	48.	Budzáková M, Galvánek D, Littera P, Šibík J (2013) The wind and fire disturbance in central European
896		mountain spruce forests: The regeneration after four years. Acta Soc Bot Pol 82(1):13–24.
897	49.	Picon-Cochard C, Nsourou-Obame A, Collet C, Guehl JM, Ferhi A (2001) Competition for water
898		between walnut seedlings (Juglans regia) and rye grass (Lolium perenne) assessed by carbon isotope
899		discrimination and delta180 enrichment. <i>Tree Physiol</i> 21(2–3):183–191.
900	50.	Major KC, Nosko P, Kuehne C, Campbell D, Bauhus J (2013) Regeneration dynamics of non-native
901		northern red oak (Quercus rubra L.) populations as influenced by environmental factors: A case
902		study in managed hardwood forests of southwestern Germany. For Ecol Manage 291:144–153.
903	51.	Harmer R, Robertson M (2003) Seedling root growth of six broadleaved tree species grown in
904		competition with grass under irrigated nursery conditions. <i>Ann For Sci</i> 60(7):601–608.
905	52.	Bloor JMG, Leadley PW, Barthes L (2008) Responses of <i>Fraxinus excelsior</i> seedlings to grass-induced
906		above- and below-ground competition. <i>Plant Ecol</i> 194(2):293–304.
907	53.	Kawaletz H, et al. (2013) Back to the roots: how do seedlings of native tree species react to the
908		competition by exotic species? Ann For Sci 71(3):337–347.
909	54.	Catorci A, Scapin W, Tardella FM, Vitanzi A (2012) Seedling survival and dynamics of upper
910		timberline in central Apennines. <i>Polish J Ecol</i> 60(1):79–94.
911	55.	Collet C, Piboule A, Leroy O, Frochot H (2008) Advance Fagus sylvatica and Acer pseudoplatanus

912 seedlings dominate tree regeneration in a mixed broadleaved former coppice-with-standards forest.
913 *Forestry* 81(2):135–150.

914 56. Diaci J, Adamic T, Rozman A (2012) Gap recruitment and partitioning in an old-growth beech forest

915 of the Dinaric Mountains: Influences of light regime, herb competition and browsing. *For Ecol*

916 *Manage* 285:20–28.

Table 2: Details of the 36 data sets included in this study. The ID of data set refers to the location on the map in Fig. 1

ID	Country	Study region	Lat (°N)	Long (°E)	Study area (ha)	Plot size (range; m ²)	Initial survey year	Recent survey year
1	В	Gaume	49.6	5.6	1000	50-400	1953-1963	2008
2	В	Binnen-Vlaanderen	51.1	3.5	30000	100-200	1977-1983	2009
3	В	Zoerselbos	51.3	4.7	30	100	1982	2008
4	В	Meerdaalwoud	50.8	4.7	1319	125-225	1954	2000
5	В	Tournibus	50.3	4.6	228	100	1967	2005
6	S	Dalby	55.7	13.3	36	1 (16 canopy)	1935	2010
7	D	Elbe-Weser	53.4	9.2	750000	100-400	1986-1989	2008
8	CZ	Milovice Wood	48.8	16.7	2100	500	1953	2006
9	CZ	Rychlebské hory Mts.	50.3	17.1	4800	315	1941-1943	1998-1999
10	CZ	Milíčovský les	50.0	14.5	93	50-625	1986	2008
11	СН	Switzerland	47.0	7.5	1500000	100-400	1940-1965	1998
12	F	Hirson	50.0	4.1	> 1000	500-800	1956-1965	1996-1999
13	F	Andigny	50.0	3.6	> 3000	500-800	1957-1963	1995-1996
14	NL	Speulderbos	52.3	5.7	1000	100-250	1957-1959	1987-1988
15	IRL	Killarney National Park	52.0	-9.6	350	8	1991	2011
16	D	Göttingen, Carici-Fag.	51.3	9.8	4000	30-400	1955-1960	2011-2012
17	D	Göttingen, Hordelymo-Fag.	51.6	10.0	4000	75-400	1955-1967	2009
18	А	Zöbelboden	47.8	14.4	90	100-100	1993	2005-2010
19	D	Brandenburg	52.2	13.6	295	100-400	1963-1965	2012
20	SK	Slovakia, South-West	48.4	17.3	25000	500	1966-1972	2007
21	SK	Slovakia, Central	48.3	19.4	70000	500	1964-1973	2005-2007
22	SK	Slovakia, North-East	49.2	21.9	40000	500	1965-1974	2006
23	CZ	České středohoří	50.6	14.1	8700	500	1965	2012
24	CZ	Krumlov Wood	49.1	16.4	3300	400	1964-1968	2012
25	CZ	Hodonínská Dúbrava	48.9	17.1	3600	400	1965	2012
26	PL	Białowieża	52.8	23.9	4747	100-200	1966	2012
27	F/CH	Jura	46.8	6.4	2268600	200-400	1989	2007

Table 2: Continued information

28	D	Göttingen, Hünstollen	51.6	10.0	56	100-250	1992	2012
29	PL	Sanocko-Turczańskie	49.5	22.4	25000	100-400	1972-1973	2005-2007
		Mountains						
30	PL	Bazaltowa Mt	51.0	16.1	110	200-400	1992-1994	2010-2014
31	PL	Buki Sudeckie beech forest	50.9	16.0	174.42	100-160	1990	2014
32	PL	Trzebnickie Hills	51.3	17.2	25	200	1962	2011-2012
33	D	Prignitz	53.1	12.3	282340	400	1954-1960	2014
34	S	Öland	56.7	16.5	134700	225	1988	2014
35	D	Brandenburg Nord	53.0	13.5	700000	200-800	1963-1964	2014
36	D	Brandenburg Süd	51.8	13.8	500000	400	1960-1966	2014

Table 2: Continued information on each study region.

ID	Altitude (range; m a.s.l.)	Soil texture	Bedrock type	Dominant tree species
1	267-372	sand, loamy sand, sandy loam	sandstone (with variable calcareous content)	Fagus sylvatica, Quercus spp.
2	5-79	sand, sandy loam, loam	ΝΑ	Populus spp., Quercus robur, Fraxinus excelsior, Acer
				pseudoplatanus
3	20-20	sand	ΝΑ	Quercus robur, Pinus sylvestris
4	62-104	loam	Tertiary sandy formations	Quercus robur, Acer pseudoplatanus
5	226-274	loam	sandstone, shale, siltstone	Quercus spp., Carpinus betulus, Fraxinus excelsior
6	50-75	loamy clay	chalk	Fraxinus excelsior, Ulmus glabra, Quercus robur, Fagus
				sylvatica
7	14-54	loam	NA	Carpinus betulus, Fraxinus excelsior, Quercus robur
8	220-320	clay to silt	loess	Quercus robur, Carpinus betulus, Acer campestre
9	380-730	loam	gneiss, granite	Fagus sylvatica
10	270-300	loam	eolic, fluvial sediments	Quercus robur, Quercus petraea, Tilia cordata,
				Carpinus betulus
11	400-780	brown (forest) soil	chalk	Fagus sylvatica, Fraxinus excelsior, Quercus spp.
12	150-330	gravels (alluvia) to deep loess	schists	Quercus spp., Fagus sylvatica
13	145-175	loess, sand	chalk, Thanetian sand	Quercus robur, Alnus glutinosa
14	60-60	sand to loamy sand (fine to medium)	NA	Fagus sylvatica, Quercus robur

Table 2: Continued.

4.5	20.420	hannes and the shall are stated	alal ward as welstawa	0
15	30-120	brown earth-shallow peat	old red sandstone	Quercus petraea
16	200-420	clay, silt	limestone	Fagus sylvatica
17	290-420	clay, silt	limestone	Fagus sylvatica
18	623-846	sandy loam, high coarse fraction	dolomite (Hauptdolomit)	Fagus sylvatica
19	50-60	Niedermoortorf, Anmoorgley	ΝΑ	Alnus glutinosa, Carpinus betulus, Pinus sylvestris,
				Fraxinus excelsior
20	203-418	loam, clay, silt on gley soils	granodiorite, loess loam	Quercus petraea
21	226-595	loam	andesite, loess loam	Quercus petraea
22	310-618	clay-loam, loam	flysch slates	Fagus sylvatica
23	220-620	variable	basalt	Quercus petraea, Fagus sylvatica, Tilia cordata,
				Carpinus betulus, Acer spp.
24	210-400	loam (sandy loam)	granite, granodiorite	Quercus petraea, Tilia cordata, Carpinus betulus
25	165-231	sand	eolic sand	Quercus robur, Tilia cordata
26	159-172	sand with admixture of clay and silt	ΝΑ	Carpinus betulus
27	550-1320	ΝΑ	calcareous	Abies alba
28	327-422	clay, silt	limestone, red clay	Fagus sylvatica
29	400-650	clay	Carpathian flysch, alternating marine deposits of clavstones, shales, sandstones	Fagus sylvatica
30	300-360	rubble, clay	Basalt, greenstone	Quercus petraea
31	440-525	clay	shale pericytes	Fagus sylvatica

32	140-230	sand	Quarternary deposits	Abies alba
33	26-112	sand, loam	glacial deposits (Pleistocene), glaco-fluvial sands (Holocene)	Quercus robur, Fagus sylvatica, Alnus glutinosa, Carpinus betulus, Fraxinus excelsior
34	4-35	variable	Slate, limestone	Quercus robur, Acer platanoides, Fraxinus excelsior, Ulmus spp.
35	15-115	sand	glacial deposits	Quercus spp.
36	25-155	sand	glacial deposits	Quercus spp., Pinus sylvestris

Table 2: Continued.

931 Table 3-8: Parameter estimates and test statistics of spatial models fitted using recent and initial plot data from summary output in R statistics using "Kenward-

932	Roger"	' approximation. R ²	values were cal	culated followin	g the method of	of Nakagawa a	nd Schielzeth (2013)
					<u> </u>			

Deschampsia flexuosa								
Recent		Old						
Variable	Estimate	SE	t value	Pr(> t)	Estimate	SE	t value	Pr(> t)
Intercept	0.92	0.24	3.78	0.001	1.41	0.21	6.57	< 0.001
Humus quality	-0.35	0.10	-3.57	< 0.001	-0.43	0.11	-3.96	< 0.001
Soil moisture (EIV _F)	-0.40	0.11	-3.55	< 0.001	0.03	0.10	0.32	0.751
Overstorey cover	-0.46	0.10	-4.39	< 0.001	0.21	0.12	1.74	< 0.05
Random intercept (StDev)		R²			Random ir	ntercept (StDev)	R²	
Region	Residual	Marginal	Conditional	-	Region	Residual	Marginal	Conditional
0.86	1.72	0.13	0.42		0.4955	2.21	0.06	0.24

Molinia caerulea										
Recent					Old					
Variable	Estimate	SE	t-value	Pr(> t)	Estimate	SE	t-value	Pr(> t)		
Intercept	-0.61	0.18	-3.40	0.002	2.27	0.38	5.79	< 0.001		
Humus quality	-1.03048	0.11447	-8.911	< 0.001	-0.49	0.19	-2.46	0.016		
Soil moisture (EIV _F)	0.91	0.3	3.00	0.003	0.58	0.25	2.20	0.03		
Soil moisture ²	-0.55	0.15	-3.76	< 0.001	-0.46	0.16	-2.78	0.006		
Overstorey cover	-0.35	0.16	-2.21	0.03	-0.49	0.17	-2.93	0.004		
Random intercept (StDev)		R²			Random in	itercept (StDev)	R²			
Region	Residual	Marginal	Conditional	-	Region	Residual	Marginal	Conditional		
2.00	2.20	0.13	0.54		1.12	2.53	0.16	0.42		

Pteridium aquilin	um										
Recent						Old					
Variable	Estimate	SE	t-value	e Pr(> t)		Estimate	e SE		t-value	Pr(> t)
Intercept	1.52	0.19	7.73	< 0.001		1.36	0.2	20	6.74	< 0.001	
Humus quality	-0.57	0.12	-4.77	< 0.001		-0.82	0.1	11	-7.05	< 0.001	
Random intercept	: (StDev)	R²				Random	intercept	(StDev)	R²		
Region	Residual	Marginal	Condit	tional		Region	Re	sidual	Marginal	Conditi	onal
0.37	1.99	0.12	0.26			0.44	2.0)7	0.21	0.35	
Rubus fruticosus a	agg.										
Recent							Old				
Variable	Esti	mate SE		t-value	Pr(> t)		Estimate	SE		t-value	Pr(> t)
Intercept	0.7	1 0.2	22	3.20	0.004		0.74	0.23		3.17	0.005
Overstorey cover	-0.2	28 0.0	07	-4.13	< 0.001		-0.19	0.07		-2.72	0.007
Random intercept	: (StDev)	R²			_	_	Random	intercept (S	tDev)	R²	
Region	Res	idual M	arginal	Conditional	-		Region	Resi	dual	Marginal	Conditional
1.10	2.2	1 0.0	03	0.37			0.97	1.78		0.01	0.36

Rubus idaeus									
Recent					Old				
Variable	Estimate	SE	t-value	Pr(> t)	Estimate	SE	t-value	Pr(> t)	
Intercept	0.30	0.23	1.30	ns	0.23	0.24	0.90	ns	
Overstorey cover	-0.15	0.07	-2.05	0.04	-0.01	0.06	-0.26	ns	
Mean N dep	-0.30	0.14	-2.21	0.03	-0.16	0.24	-0.67	ns	
Random intercept (StDev) R ²				Random int	Random intercept (StDev)		R ²		
Region	Residual	Marginal	Conditional	_	Region	Residual	Marginal	Conditional	
1.13	1.31	0.04	0.49		0.89	1.02	0.01	0.47	
Vaccinium myrtillu	15								
Recent					Old				
Variable	Estimate	SE	t-value	Pr(> t)	Estimate	SE	t-value	Pr(> t)	
Intercept	1.25	0.26	4.75	< 0.001	1.30	0.21	6.07	< 0.001	
	-1 01	0.17	-5.80	< 0.001	-0.72	0.17	-4.24	< 0.001	
Humus quality	1.01								
Humus quality Random intercept	(StDev)	R ²			Random inte	rcept (StDev)	R ²		
Random intercept	(StDev) Residual	R ² Marginal	Conditional		Random inte Region	rcept (StDev) Residual	R² Marginal	Conditional	

940 Fig. 1: Results of species-specific models predicting the cover abundance of six understorey plant species in 941 response to spatial gradients in local-scale and large-scale explanatory variables. Points representation of 942 the estimated effects (+/- 2SE) of the explanatory variables for the spatial models using contemporary 943 resurvey data (x-axis) and historical survey data (y-axis). The line represents the 1:1 line of no difference. 944 Different shapes represent the six study species and colours the different explanatory variables.



Fig.2: Predicted cover abundance change of six understorey plant species in response to temporal changes in several environmental explanatory variables.
Predictions were based on species abundance changes along spatial environmental gradients (contemporary spatial models) and on actually observed temporal
changes in abundance. For each explanatory variable separately, we predicted the change in species cover (as log ratio) to (A) the 20% percentile and (B) 80%
percentile of the observed change in the plot-pairs between historical and contemporary resurveys. Additionally, predictions of cover change were made in the
absence of environmental change (No change). The dotted line represents these predictions using the temporal models. Explanatory variables that were retained
in the most parsimonious contemporary spatial models are annotated in grey. The lines indicate the 95% confidence intervals around the predictions. Red:
temporal predictions; Blue: predictions using SFT approach. Humus quality (Hms), Ellenberg F value (EIVF), Overstorey cover (OS), Mean annual temperature

- 954 (MAT), Mean annual precipitation (MAP) and mean N deposition rate (Ndep).
- 955 (A)







959 Appendix B: Detailed information on the temporal vs. space-for-time approach

960 We test if species abundance changes over contemporary spatial environmental gradients allow predicting 961 how environmental change over time cause species to change their abundances (research question 3). 962 First, we modelled observed changes in species abundance over time, by fitting regression models for each 963 species using the subset of plots in which a species was present at both survey times for each species 964 (Temporal model). For each plot-pair, temporal change was characterized as the natural logarithm of the 965 ratio between the contemporary resurvey and the historical survey. We did this for the response variable 966 (percentage cover), and for all the explanatory variables used in the spatial models. Similar as for the spatial 967 models, we used a multilevel modelling approach with random effects for 'data set'.

968
$$\ln\left(\frac{y_r}{y_i}\right) = \beta_0^T + \beta_1^T \ln\left(\frac{x_r}{x_i}\right) + \dots + random part$$

969

979

970 In this model, y_r and y_i are the percentage cover of one of the study species at the contemporary and 971 historical survey resp., x_r and x_i are one of the six explanatory variables at the contemporary and historical 972 survey resp. and *random part* denotes the effect of 'data set' and residual error. The intercept β_0^T catches 973 the average temporal change in abundance not accounted for by the used explanatory variables.

974 Second, these temporal models then allowed making predictions of abundance change over time in

975 response to a particular change in environmental conditions. For each predictor variable separately, we

976 predicted the change in species cover (as log ratio) for a realistic change in the predictor (also as log ratio):

977 here we used the observed mean change in the predictor between the two survey times, $\bar{r_{\chi}} = \frac{\bar{x_r - x_l}}{x_l}$ (Table).

978
$$\widehat{y^T} = \ln\left(\frac{y_r}{y_\iota}\right) = \beta_0^T + \beta_1^T \ln\left(\frac{x_r}{x_\iota}\right) = \beta_0^T + \beta_1^T \ln(1 + \overline{r_x})$$

Temporal prediction

Additionally, predictions were made for the mean change in abundance in the absence of environmentalchange:

982
$$\widehat{y^T} = \beta_0^T$$

'No change' prediction

984

985 but based on the species' abundance patterns along spatial gradients, that is, using the following 986 contemporary spatial model: $\ln(y) = \beta_0 + \beta_1 \ln(x_1) + \dots + \beta_6 \ln(x_6) + random part$ 987 988 **Spatial model** Using the parameter estimates of the spatial model, we predicted the cover abundance for the mean of 989 each predictor value ($\overline{x_{1,\dots,6}}$) over all the contemporary plots in which at least one species is present: 990 991 $\widehat{\ln(y_1)} = \beta_0 + \beta_1 \ln(\overline{x_1}) + \dots + \beta_6 \ln(\overline{x_6})$ 992 Equation [2] 993 Similarly, we made a second prediction of the species cover, but one of the predictor variables, here $\overline{x_1}$, 994 was increased by the same mean change in that predictor between the two surveys as in the temporal 995 predictions, i.e. $\overline{r_{x_1}} = \overline{r_x}$. All other predictor variables were kept at their mean values as in Equation [2]. $\widehat{\ln(y_r)} = \beta_0 + \beta_1 \ln\left(\overline{x_1}(1 + \overline{x_1})\right) + \beta_2 \ln(\overline{x_2}) + \dots + \beta_6 \ln(\overline{x_6})$ 996 997 Equation [3] 998 Equation [2] and [3] were then combined in a log ratio: $\widehat{y^{SFT}} = \widehat{\ln\left(\frac{y_r}{y_l}\right)} = \widehat{\ln(y_r)} - \widehat{\ln(y_l)}$ 999 $= \left[\beta_0 + \beta_1 \ln\left(\overline{x_1}\left(1 + \overline{r_{x_1}}\right)\right) + \beta_2 \ln(\overline{x_2}) + \dots + \beta_6 \ln(\overline{x_6})\right] - \left[\beta_0 + \beta_1 \ln(\overline{x_1}) + \beta_2 \ln(\overline{x_2}) + \dots + \beta_6 \ln(\overline{x_6})\right]$ 1000 $=\beta_1 \ln(1+\overline{r_{x_1}})$ 1001 1002 Space-for-time prediction

Third, we made similar predictions of temporal change in abundance in response to environmental change,

- 1003 In this way, we were able to compare the predicted change in the species' abundances for the actual
- 1004 temporal vs. the spatial model, based on the same change $\overline{r_x}$ in an environmental predictor x.