Climate warming disrupts mast seeding and its fitness benefits in European beech

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**Introductory paragraph**

Many plants benefit from synchronous year-to-year variation in seed production, called masting. Masting benefits plants because it increases the efficiency of pollination and satiates predators, which reduces seed loss. Here, using a 39 year-long dataset, we show that climate warming over recent decades has increased seed production of European beech, but decreased the year-to-year variability of seed production and the reproductive synchrony among individuals. Consequently, the benefits that the plants gained from masting has declined. While climate warming was associated with increased reproductive effort, we demonstrate that less effective pollination and greater losses of seeds to predators offset any benefits to the plants. This shows that an apparently simple benefit of climate warming unravels because of complex ecological interactions. Our results indicate that in masting systems, the main beneficiaries of climate-driven increases in seed production are seed predators, not plants.

**Introduction**

Anthropogenic environmental changes exert increasing pressure on forests world-wide1. Research on the effects of environmental change on trees has focused on the impacts on tree growth, carbon sequestration, mortality, or phenology2–4. There have been few studies of the impacts of environmental change on the reproductive ecology of trees, yet long-term changes in reproduction determine the ability of trees to disperse to shifting habitats and recolonize sites after disturbance5. What is more, reproduction of trees is likely to be sensitive to climate change, especially in the case of mast seeding (or masting) species – i.e. those that reproduce through spatially synchronous and temporally variable seed production – due to strong correlations of seed production with annual variation in weather6–8.

Numerous plant species, including the majority of forest-forming tree species in temperate zones, reproduce through mast seeding9,10. While masting plants are expected to be sensitive to global change11–13, the direction of that change is a source of controversy. Predictions range from an increase, to a decrease, to an unchanged strength of masting (i.e. interannual variability and synchrony) in response to climate change11,14,15, depending on the sensitivity of masting to climate and associated resources. However, the strength of masting is crucial in terms of plant fitness and forest regeneration, since masting is a life history trade-off between missed reproductive opportunities in low-seed years and enhanced pollination efficiency and decreased seed predation in mast years16–18. Studies that measure seed production, pollination efficiency, and seed predation for long enough to capture potential change are the key to understanding how anthropogenic climate change will affect plant reproduction and thus global vegetation dynamics, but such studies are virtually non-existent. Here, we report on a unique 39-year study of 139 individuals of European beech (*Fagus sylvatica*) across twelve sites in England that documents how the success of this major European forest-forming tree species, including seed production, pre-dispersal seed predation by *Cydia* *fagiglandana* (Lepidoptera), and pollination success, has changed throughout the last four decades.

Mast seeding enhances plant fitness through economies of scale that decrease the cost of reproduction per surviving offspring12,19. This mainly happens through two mechanisms. The first is that large and synchronized flowering effort enhances pollination success, because pollination efficiency is increased when the density of flowers is high (density dependent, pollination efficiency hypothesis) 16,20,21. Secondly, masting decreases seed predation by starving predator populations in years of low-seed production, and then satiating them in high-seed years (predator satiation hypothesis) 22–24. Predator satiation is crucially dependent on the sequence of high and low years, whereas pollination efficiency is affected by the size, not sequence, of high-flowering years25. Therefore, the effects of global change will act differently on these two economies of scale, depending on how it alters plant reproductive variability and reproductive synchrony. This, in turn, depends on the causal relationship between masting and weather cues. Notably, if plants respond to the temperature difference between the two previous summers, as suggested by the ΔT model, then masting should be relatively insensitive to increased mean temperatures (though sensitive to changes in temperature fluctuation)14. By contrast, if seed production is driven by interactions between plant internal resource dynamics and weather cues such as warm absolute temperatures, more frequent cueing will decrease synchrony among plants, reducing population-level year-to-year variation in seed crops17,26. Similarly, according to the environmental stress hypothesis, resource-augmented plants should increase mean seed production, and their reproduction should become less variable over time25,27. If global environmental change disrupts masting patterns, the long-term regeneration of masting plants may decrease at the same time as global warming requires increasingly rapid geographic range shifts to keep species within suitable climate zones28,29.

**Results**

Our 39-year dataset of seed production in 139 European beech trees indicated that mean reproductive effort increased during the study period, but mast seeding became progressively weaker (Fig. 1). Mean annual seed production per tree increased (Fig 1a, z = 4.29, p < 0.001), while the probability of a tree having a year of zero seed production decreased over time (z = -2.29, p = 0.003). Accordingly, sliding window analysis indicated that interannual variation in reproduction measured by the coefficient of variation (CV)30 decreased 40% at the population level (CVp) from around 1.15 to 0.70 (Fig 1b, z = -4.93, p < 0.001), and at individual-tree level (CVi) from around 1.30 to 0.80 (Fig 1c, z = -9.74, p < 0.001). Furthermore, synchrony, measured by mean cross-correlation of seed production among trees, also decreased by 30% from over 0.85 to around 0.60, both at the within-site (Si) (Fig 1d, z = -4.66, p < 0.001) and among-site level (Sp) (Fig 1e, z = -7.27, p < 0.001). Thus, population-level variation in seeding across years decreased because individual trees varied less from year to year, and trees were more poorly synchronized.

The analysis of abiotic drivers of seed production supported mathematical models of the role of environmental cues/vetoes in mast seeding, which predict that more frequent occurrence of weather cues for reproduction should result in a decrease in both variability and synchrony of seed crops17,26. In the case of European beech, seed production correlates positively with warm summer temperature one year prior to seed dispersal (Supplementary Table 1), likely by promoting flower initiation8,31 (the AIC analysis provided less support for the ΔT model as the driver of masting in our population: ΔAIC = 130.5, see Supplementary Table 1). Our analysis of temporal contributions attributed the temporal change in seed production largely to increasing temperatures (Fig. 2). Sensitivity of seed production to increasing growing season mean temperature was 0.64 ± 0.44 [mean ± s.e.m.] (seeds per 7-min count per tree yr-1for each °C yr-1), while sensitivity to increasing summer temperature a year prior to seed dispersal was 1.30 ± 0.90 (seeds per 7-min count tree yr-1for each °C yr-1). This suggests that the increase in mean seed production was caused by two parallel mechanisms. First, warmer growing season temperatures are likely to enhance average seed output through lengthening of the growing season and increased photosynthesis32. Second, more frequent warm summers in recent years apparently trigger large flowering events more frequently, and simultaneously reduce the frequency of years with no reproduction. The resource budget model predicts that more frequent weather cueing should weaken the reinforcing dynamics of stored resources on among-tree synchrony of reproductive variation17,26 – a pattern consistent with our data. While other variables, such as summer temperature two years before seed fall or nitrogen deposition in the past 5 years, significantly explained variance of annual seed production in our beech populations (Supplementary Table 1), they did not correlate with the temporal trends in seed production (Fig. 2).

Unfortunately for the trees, desynchronization and more regular seeding weakened the masting benefits that economies of scale provided to European beech (see Supplementary Table 2 for outputs of all models). The proportion of pre-dispersal seed predation by *Cydia* sharply decreased with increasing ratio of the previous to the current year seed production (Fig. 3, z = -8.90, p < 0.001), indicating a strong starvation effect (numerical response) of beech masting on the moth population. The slope of that relationship did not change with time, but the intercept progressively increased with time (z = 9.72, p < 0.001, Fig. 3). Thus, while masting still starved insects during years of low seed production, a lesser effect on predator numbers was seen in recent years (Fig. 3). Moreover, in accordance with the predator satiation hypothesis, weaker predator starvation resulted in less effective predator satiation: the proportion of predated seeds decreased with the number of seeds produced, but the slope of that relationship became less steep in recent years (interaction term seed production × year: z = 2.66, p = 0.008, Fig. 3).

In the case of pollination efficiency economy of scale, the proportion of successfully pollinated seeds increased with the summed reproductive effort of conspecifics in the population, and that effect was stronger in years when the synchrony of flowering was higher (Fig. 3, Supplementary Table 2). In the model that included both summed conspecific flowering effort and synchrony, the effect of year was not significant (p = 0.07). Thus, while large and synchronized reproductive effort always increased pollination efficiency, such large and synchronized flowering efforts happened less often in recent years, lowering the mean pollination efficiency (see below).

As a consequence of these weakened economies of scale, the estimated proportion of seeds predated by insects increased dramatically from 1% at the beginning of the study to over 40% in the final years (linear effect: z = 6.07, p < 0.001, quadratic effect: p = -3.42, p < 0.001, Fig. 4). Simultaneously, the pollination efficiency decreased from 50% to 33% (z = -2.20, p = 0.03, Fig. 4). Thus, the large temporal increase in total seed production (Fig. 4, β = 0.04 ± 0.01 [SE]) became weaker once only pollinated seeds were considered (β = 0.03 ± 0.01, z = 4.21, p < 0.001), and that positive trend almost disappeared for only pollinated and not predated seeds (β = 0.02 ± 0.01, z = 2.22, p = 0.04, Fig. 4). Overall, the increased predation pressure and decreased pollination efficiency led to 2.6-fold decrease in the probability that a female flower becomes a sound seed. This probability was estimated to equal 47% (95% confidence interval: 31% – 65%) at the beginning of the study period, and decreased to 18% (7% – 36%) in 2010s (Fig. 4). This implies that the costs to European beech per viable seed more than doubled during the last four decades as economies of scale became increasingly ineffective (Fig. 4d).

**Discussion**

Altered mean seed production induced by global changes has been reported in few previous studies, and has found both increase33,34, and decrease27 in reproductive effort. How this translates into recruitment potential was, however, unknown since this crucially depends on parallel changes in variability and synchrony of reproduction and their consequences for seed predation and pollination efficiency 7,13. Our study is the first to show that a long-term increase in mean seed production is accompanied by a decrease in synchrony and year-to-year variability of reproduction, and that these are associated with elevated costs. The apparently improved performance of European beech trees in a warmer world is offset by the breakdown of selectively beneficial strategies to avoid seed predators and improve pollination. Hence, the benefits of higher reproductive effort by the trees are largely offset by the effects of lower pollination success and higher pre-dispersal seed predation.

The temporal decrease in variability and synchrony of seed production provides the first documented natural experiment of how long-term changes in climate alter seed crop patterns and thereby alter the effectiveness of economies of scale. The predator satiation hypothesis consists of two parts, in which the starvation of seed predators in years of low seed production results in the satiation of reduced populations of predators in subsequent mast years35,36. More regular seeding of European beech, and the reduction in the frequency of seed failures in this study weakened the starvation effect which in turn lowered the effectiveness of pre-dispersal predator satiation. Such an effect is in line with past experimental and observational studies that found warming or nitrogen deposition was associated with increased mean seed production but also higher seed predation13,37. Furthermore, the disappearance of highly-synchronised mast years in recent years reduced the efficiency of pollination, and thus the proportion of successfully pollinated flowers. Together, the breakdown in economies of scale resulted in only small reproductive benefits to the studied trees, disproportionate to the size of the increase in long-term reproductive effort. Worryingly, successful beech regeneration also requires the successful starvation and satiation of *post*-dispersal seed predators, especially small mammals38,39. Here we show that predator satiation became less efficient in the case of a pre-dispersal insect seed predator, but if post-dispersal predator satiation also became less effective, as we would predict, the net effect on surviving seeds (Fig. 4) could easily be negative overall. This is an important area for further research.

Climate warming was the major driver of decreased variability and synchrony of reproduction in European beech. Even though our statistical analyses do not directly prove causality, the results support the dominant role of increased temperatures in explaining the trends in masting pattern. First, our analysis suggested that the trend in increasing seed production is driven by warming temperatures in the preceding summer, a well-documented cue for flowering in European beech8,31,40. The resource budget models of masting17,41 predict that more frequent occurrences of the weather cue will decrease individual variability and desynchronize flowering by weakening the reinforcing effects that resource dynamics have on reproductive synchrony26,42, which jointly decrease population-level seed production variability. This mechanism is supported empirically in European beech, as we found that increased mean seed production is associated both with significantly lower CVi and with lower synchrony (Si). Second, the important effect on mean seed production of increasing mean temperatures during growing seasons supports the environmental stress hypothesis, that predicts masting plants will produce on average more seeds with lower variability when resource availability is high10,25,27. According to this hypothesis, plants growing under favourable conditions will be able to more rapidly accumulate the resources required for reproduction and, therefore, present a more regular pattern in seed production – a pattern consistent with the lower CVi in our data. The environmental stress hypothesis does not explicitly predict the effect of resource augmentation on reproductive synchrony. Nevertheless, warmer growing season temperatures and associated increases in resource availability potentially makes resources less limiting for reproduction, weakening the positive effect of plant internal resource dynamics on synchronising masting17,26,43.

The changes in masting patterns reported in this study will have profound implications for the long-term future of European beech. In northern Europe where beech is predicted to expand northwards under climate warming44, our results hint that a breakdown in masting might result in recruitment limitation. This emphasises the importance of including realistic reproduction processes within forest models45. Similar processes may explain observed declines in beech recruitment elsewhere in Europe46,47. More generally, our results imply potential climate-driven changes in the dynamics of beech-dominated ecosystems, including the spread of Lyme disease and Hantavirus by rodents dependent on beech seeds48–50, habitat selection of ground-nesting birds51, and population dynamics of small mammal predators52. Similar changes are likely to occur in other masting species, with other negative conservation consequences53, but such long-term datasets as the one reported here are extremely rare27. Thus, experiments to better understand the mechanisms underlying masting, and consequently better predict the consequences of a changing climate for plant reproductive patterns and global vegetation dynamics, should become a research priority54. The net benefits of climate warming in the European beech system studied here accrue largely to an invertebrate seed predator, which suggests that ecological interactions may determine the real-world consequences of global change.

**Methods**

**Study species.** European beech (*Fagus sylvatica* L.) is a major forest-forming species in temperate Europe. The seeds are eaten and dispersed by a suite of vertebrates37,55, and destroyed by a seed-eating moth, *Cydia fagiglandana* Z. (Tortricidae). Because fruit and seed coats develop if pollination occurs and unpollinated fruits lack a seed (kernel)56, relatively accurate estimates of pollination can be made from seed production data. European beech produces flowers in the spring, which are fertilized and develop into mature fruit in the same year as they were pollinated. Flower induction happens in the summer prior to the flowering year.

**Data collection.** We sampled seed production in 139 beech trees located at 12 sites spaced across England annually for 39 years (1980 – 2018) (Supplementary Figure 1). The ground below each tree was searched for seeds for 7 minutes, and seeds were later classified as sound, or empty with formed pericarps (not pollinated), or damaged by *Cydia* sp. moth. Detailed description of sites and procedures is given in56.

Nitrogen deposition data for each site was estimated using the FRAME model (Fine Resolution Atmospheric Multipollutant Exchange58), an atmospheric chemistry and transport model, to produce estimates of N deposition59. Deposition data was available at a 5 x 5km resolution and produced estimates of both total reduced and oxidised N (kg N ha-1 yr-1) over deciduous forest for three snapshots (1970, 1990, 2010). Linear interpolation was used to estimate annual deposition. Data was provided by CEH Edinburgh. Mean monthly maximum temperature and monthly sum precipitation for each site was obtained from the corresponding 0.25˚ grid cell of the E-OBS dataset (version 19.0e)60.

**Analysis.**

**Trends in seed production.** We started our analysis by asking whether and how seed production patterns of European beech changed over the last 39 years. We built a zero-inflated, negative binomial family generalized linear mixed model (GLMM) with autoregressive order-1 autocorrelation structure, and included total yearly seed production per tree as a response while year was included as fixed effect in both in count and logit part of the model. The model included tree ID and site as random intercepts.

**Temporal contributions and sensitivities of changes.** To attribute the temporal trends in seed production to its possible drivers, we used the temporal contribution method61,62. First, using the tree ID and site as the random intercepts and an autoregressive order-1 autocorrelation structure, we modelled seed production as a function of weather cues identified as important for beech seeding in past studies63,64, i.e. mean max temperature and summed precipitation in summer (June and July) one and two years before seed dispersal. We also included mean temperature in the growing season (May-August) and nitrogen deposition, i.e. the cumulative totals of the previous 5 years to test whether resource augmentation drives the trend. We then used the full model to predict the change of the response variables during the study period (1980–2018). We first calculated the observed trend (slope estimate ± standard error of the slope estimate) in our data using GLMMs with random intercepts and temporal autocorrelation structure (see above, Trends in seed production). We then calculated the trend predicted by the full model and the trends predicted by the same model but maintaining the predictors constant one at a time (e.g., summer temperature is held constant, using the median values per site, while all other predictors change according to the observations). The difference between the observed trend and when one variable was controlled was the contribution of that predictor variable to the change in the response variable. The difference between all individual contributions and the observed trend were considered to be unknown contributions. Finally, we calculated the average seed production sensitivities to predictor changes by dividing the temporal contributions by the trends of the predictor variables. All errors were calculated using the error-propagation method61.

**Changes in masting behaviour (time series).** To explore whether changes in mean seed production were accompanied by changes in annual variation and among-plant synchrony in reproduction we calculated variability and synchrony on a 10-year sliding window with 1-year step size from 1980 to 2018 for each tree (for plant-level analysis) or site (for site-level analysis) separately. We used the coefficient of variation (CV) as a measure of temporal variability, and the mean Pearson cross-correlation as a measure of reproductive synchrony. CV was calculated both at tree- (CVi) and population-level (CVp)30. Similarly, synchrony was calculated as a mean pairwise correlation of seed production of an individual plant with all other individuals observed at the site (within-site synchrony, Si), or as a mean pairwise correlation of mean site-level seed production with all other sites (among-populations synchrony, Sp). Next, these rolling masting metrics were included as a response in linear mixed models with autoregressive order-1 autocorrelation structure and tree ID and site (for CVi and Si) or only site (CVp and Sp) included as random intercepts.

**Economies of scale.** We explored whether beech masting provides fitness benefit through economies of scale, and whether that benefit changed with time. To do so, we built three binomial family GLMMs: two with proportion of seeds predated (predator satiation economy of scale), and another with proportion of seeds pollinated (pollination efficiency economy of scale) included as a response, all per tree per year. In all models, we used tree ID and site as random intercepts, observation-level random effect to account for overdispersion, and autoregressive order-1 autocorrelation structure. In the first predator satiation model, the fixed effect included tree-level yearly seed production (satiation effect or functional response of the predator), while in the second, the fixed effect included the ratio of the number of seeds produced in the current year to the number produced in the previous year (starvation effect or the numerical response of the predator). In the pollination efficiency model, fixed effects included summed number of total seeds produced by all conspecific trees at a particular site, the within site within year synchrony of seeding (calculated as coefficient of variation), and their interaction term. In all models, we also included the quadratic terms of fixed effects which were retained or dropped based on the standard AIC criteria65. Finally, to test for temporal patterns in economies of scale we included the second-order interactions of the above-listed fixed factors with year. These were removed from the final models if not significant.

**Temporal changes in pollinated and unpredated seeds, and costs of reproduction**. In the final step, we explored how the proportion of pollinated and predated seeds changed over time, how this translated into the production of “pollinated”, and “pollinated and not predated” seeds, and how all these affected beech reproductive costs. The temporal change in proportion of predated and pollinated seeds was tested using binomial GLMMs with similar structure as in tests of economies of scale, but using year as a fixed factor. In the case of proportion of predated seeds, we also included the previous year seed production as a covariate as this strongly affects *Cydia* population dynamics (see Results). In turn, temporal change in the production of “pollinated”, and “pollinated and not predated” seeds was modelled similarly as in the models exploring temporal changes in total seed production, but we used tree-level number of pollinated or pollinated and not predated seeds as a response. The change in reproductive costs was calculated as 1-S, where S was calculated as probability that a seed is pollinated times probability that it will avoid *Cydia* predation. We used parameter estimates from the above-described GLMMs and obtained standard errors with parametric bootstrapping, that is, sampling from the distributions defined by the mean and standard error of each coefficient to obtain a joint distribution for the derived variables.

All statistics were run in R (version 3.4.1), mixed models were fitted via glmmTMB66. Model validation was run with the DHARMa package 67. We used the windowscanr package for rolling window analysis68. In models in which we detected heteroscedasticity of residuals, the dispersion was modelled as a function of fixed factors using the dispformula function.

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**Authors’ contributions.** MB conceived the study and drafted the manuscript; MB led the analysis with input from AHP and DK; PT, JL, AHP collected and managed the data. All authors interpreted the results, revised the text, and provided critical feedback and helped shape the final text.

**Conflict of interest.** Authors declare no competing interests as defined by Nature Research, or other interests that might be perceived to influence the results and/or discussion reported in this paper.

**Data availability.**

The data that support the findings of this study are available on request from the corresponding author (MB).

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Figure 1. Temporal trends in European beech (*Fagus sylvatica*) seed production in England, population- and individual-level variability (CV), and within- and among-site synchrony of reproduction for 12 sites and 139 trees (1980–2018). (a) Observed population-level seed production per year per 7-minute search periods, and fitted long term mean. Significant fitted trends in temporal change in (b) population-level CVp, (c) mean individual CVi, (d) synchrony among trees within a site (Si), and (e) synchrony among sites (Sp). Trends were calculated using GLMMs, with the site and trees as random intercepts and year as a fixed effect. Models also included an autoregressive order-1 autocorrelation structure. Shading around prediction lines indicates the 95% confidence intervals. See Methods for further details.

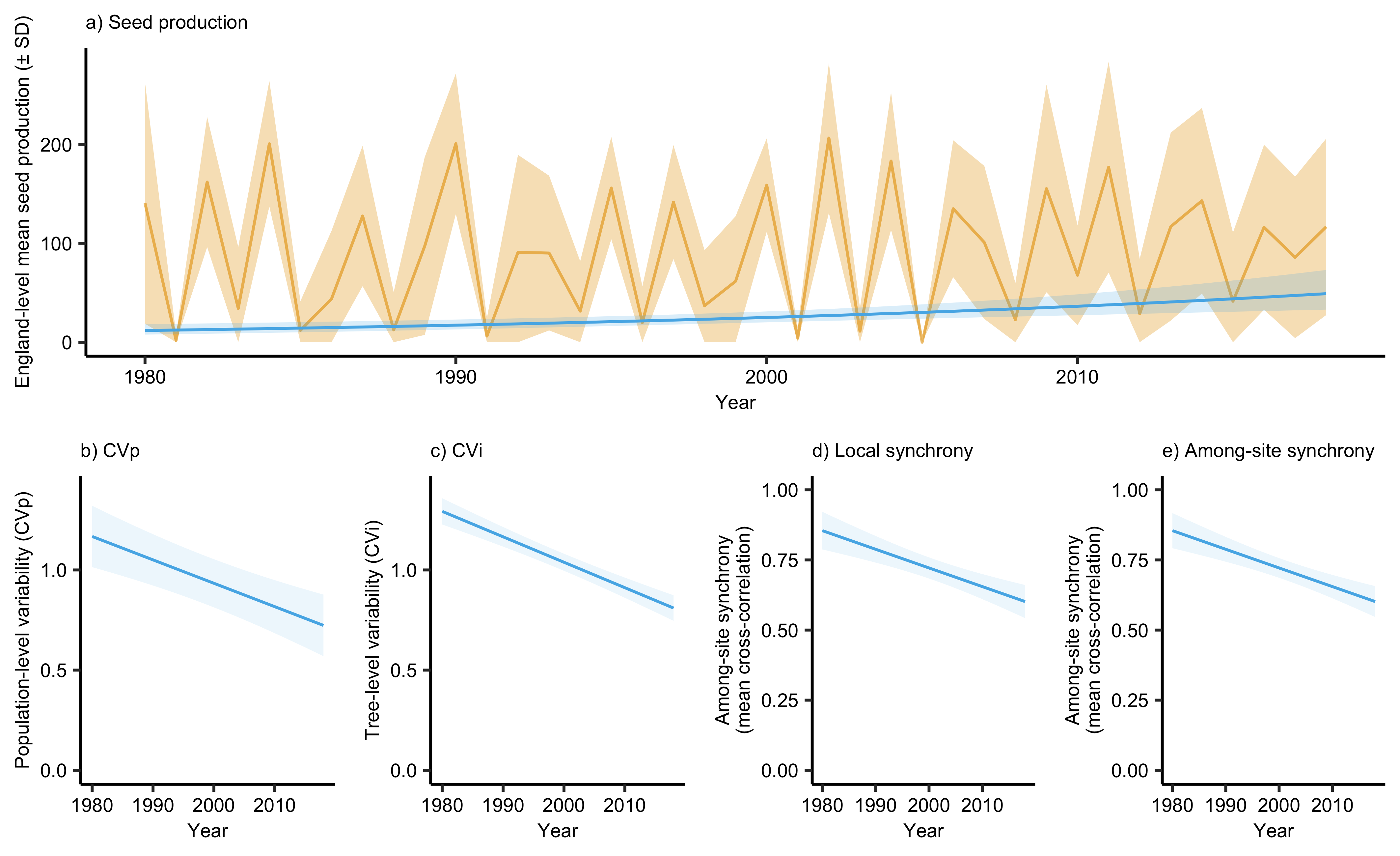


Figure 2. Temporal contribution of the predictor variables. The analysis, based on 12 sites and 139 trees (1980–2018), suggested that increasing temperature is the main contributor to the observed increase in European beech (*Fagus sylvatica*) seed production (in seeds per tree per year per 7-minute search periods). The difference between the modelled contributions and the observed trends has been considered as an unknown contribution to the temporal variation seed production. The temporal trends of the predictors are shown in square brackets. Error bars for associated contributions indicate standard errors. TempT1 is the mean max June-July temperate in the year prior seed dispersal, while TempT2 two years before. PrecipT1 is summed June-July precipitation in the year prior seed dispersal. Units are °C for temperature, kg ha−1 yr−1 for N deposition, mm for precipitation. See Methods for information about the methodology used to calculate the contributions. Significance levels (two-sided t-test): + P = 0.10; \*P = 0.04; \*\*\*P < 0.001.

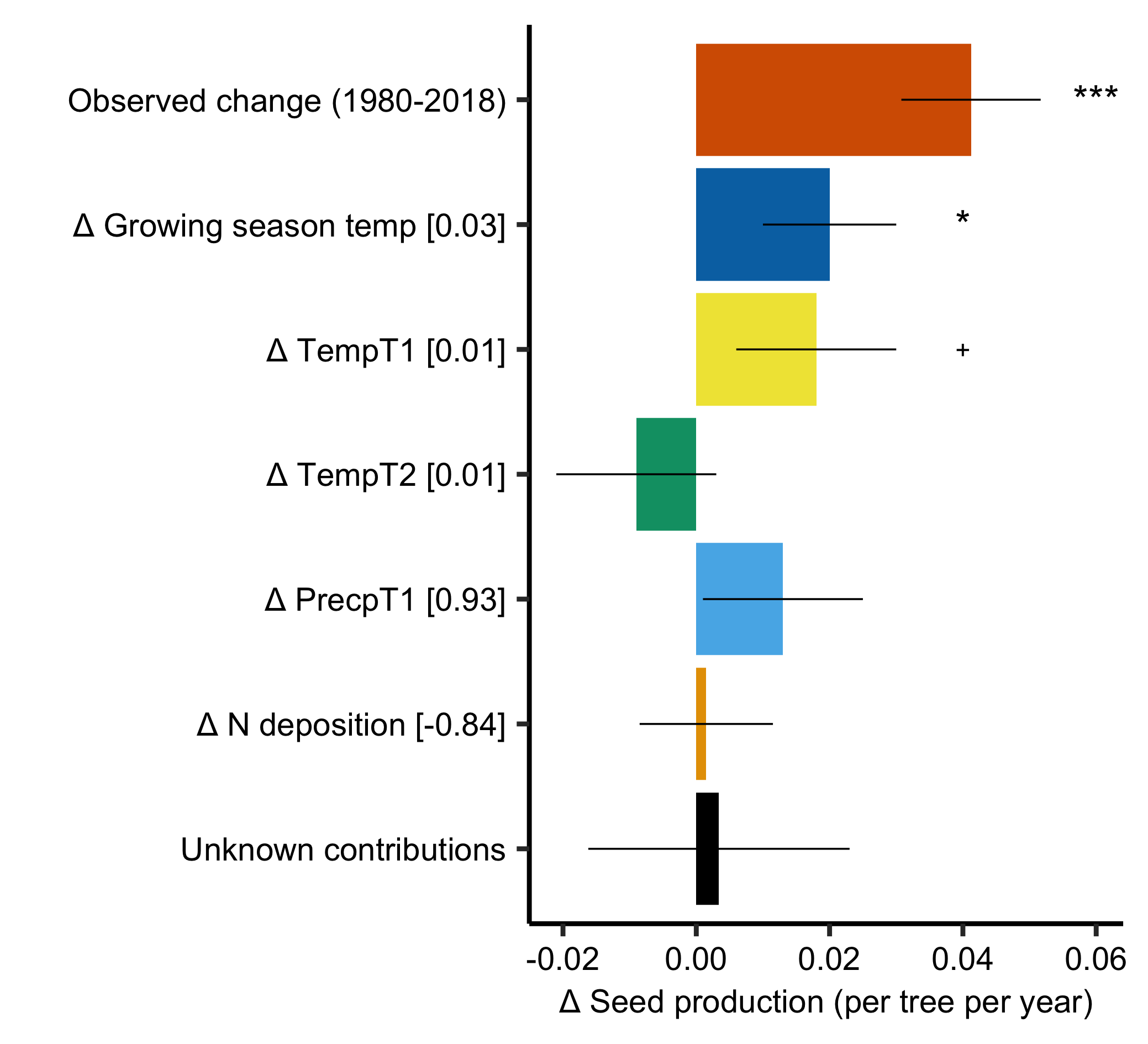


Figure 3. Weakening benefits to European beech from mast seeding. The proportion of seed production of each tree infested by *Cydia* sp. moths vs. (a) the current seed production per tree, and (b) the ratio of seed production in the current vs. the previous year. Different color lines show predictions for early (before 2000, black) and late (after 2000, yellow) monitoring periods. In later years, predation was higher for a given level of seed crop variation. (c) The proportion of successfully pollinated seeds on a plant vs the summed reproductive effort of all other conspecific trees at a particular study site. Different color lines show estimates for low (CV > 0.5, yellow) and high (CV < 0.5, green) within-site synchrony of flowering. When sites were more highly synchronous, pollination increased for a given flowering effort. Synchrony was calculated as CV of within-site among-tree flowering effort (thus, smaller value indicates higher among-tree synchrony of seeding). The lines are based on the significant binomial GLMMs predictions, while shading indicates the 95% confidence intervals, and is based on 39-yr data set of 139 beech trees spaced across 12 sites in England. All models included the interaction term between the predictor showed at the x-axis and year. In the case of pollination efficiency, once synchrony and flowering effort were included, year effect became not significant (see Table S2).

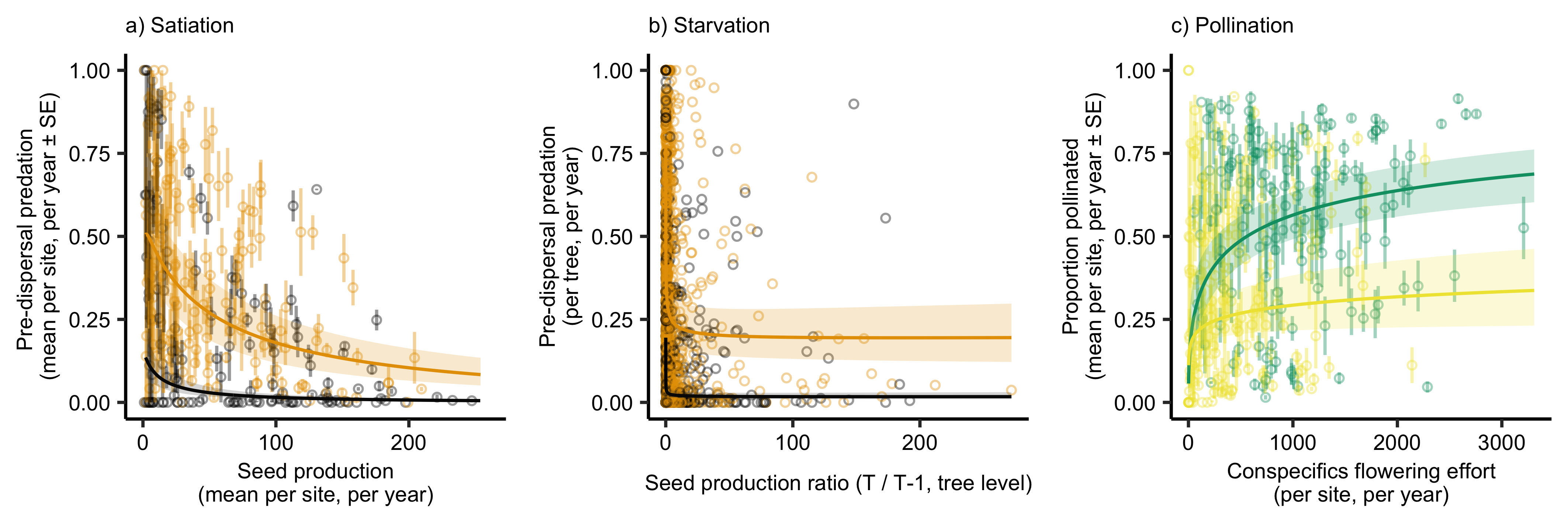


Figure 4. Temporal trends in European beech (a) total and effective seed production, (b) pollination efficiency, (c) seed predation, and (d) economies of scale, for 12 sites and 139 trees (1980–2018). Trends were calculated using GLMMs, with the site and trees as a random intercepts and year as a fixed effect. Models also used an autoregressive order-1 autocorrelation structure. Shading indicates the 95% confidence intervals of the means. In (a) temporal trend was calculated based on either all seeds produced, only the fraction of pollinated seeds, or only pollinated and not-predated seeds. See Methods for further details. D) shows means and 95% confidence intervals.

