PRIMARY RESEARCH ARTICLE





Interactive climate factors restrict future increases in spring productivity of temperate and boreal trees

Constantin M. Zohner¹ | Lidong Mo¹ | Thomas A. M. Pugh^{2,3} | Jean-Francois Bastin^{1,4} | Thomas W. Crowther¹

Correspondence

Constantin M. Zohner, Institute of Integrative Biology, ETH Zurich (Swiss Federal Institute of Technology), Universitätsstrasse 16, 8092 Zurich, Switzerland. Email: constantin.zohner@t-online.de

Funding information

ETH Zurich Postdoctoral Fellowship; China Scholarship Council; DOB Ecology; Plantfor-the-Planet; German Federal Ministry for Economic Cooperation and Development; European Union's Horizon 2020, Grant/ Award Number: 758873

Abstract

Climate warming is currently advancing spring leaf-out of temperate and boreal trees, enhancing net primary productivity (NPP) of forests. However, it remains unclear whether this trend will continue, preventing for accurate projections of ecosystem functioning and climate feedbacks. Several ecophysiological mechanisms have been proposed to regulate the timing of leaf emergence in response to changing environmental cues, but the relative importance of those mechanisms remains unclear. Here, we use 727,401 direct phenological observations of common European forest trees to examine the dominant controls on leaf-out. Using the emerging mechanisms, we forecast future trajectories of spring arrival and evaluate the consequences for forest carbon dynamics. By representing hypothesized relationships with autumn temperature, winter chilling, and the timing of spring onset, we accurately predicted reductions in the advance of leaf-out. There was a strong consensus between our empirical model and existing process-based models, revealing that the advance in leaf-out will not exceed 2 weeks over the rest of the century. We further estimate that, under a 'business-as-usual' climate scenario, earlier spring arrival will enhance NPP of temperate and boreal forests by ~0.2 Gt per year at the end of the century. In contrast, previous estimates based on a simple degree-day model range around 0.8 Gt. As such, the expected NPP is drastically reduced in our updated model relative to previous estimates—by a total of ~25 Gt over the rest of the century. These findings reveal important environmental constraints on the productivity of broad-leaved deciduous trees and highlight that shifting spring phenology is unlikely to slow the rate of warming by offsetting anthropogenic carbon emissions.

KEYWORDS

carbon cycle, climate change, phenology, spring leaf-out, temperate forests, terrestrial carbon sink

1 | INTRODUCTION

Shifts in the timing of annual growth cycles in temperate and boreal trees have direct impacts on global biogeochemical cycles (Keenan et al., 2014; Richardson et al., 2010), species distribution

patterns (Chuine, 2010), and ultimately feedback to the climate system by affecting the atmospheric carbon budget (Richardson et al., 2013). There is broad consensus that warming trends over the past decades have led to an earlier arrival of spring leaf emergence in Northern Hemisphere trees, a trend that is enhancing global primary productivity under climate change (Keenan et al., 2014; Menzel & Fabian, 1999; Zohner & Renner, 2014). Depending

Constantin M. Zohner and Lidong Mo should be considered joint first author.

Glob Change Biol. 2020;00:1–14. wileyonlinelibrary.com/journal/gcb © 2020 John Wiley & Sons Ltd

¹Institute of Integrative Biology, ETH Zurich (Swiss Federal Institute of Technology), Zurich, Switzerland

²School of Geography, Earth and Environmental Sciences, University of Birmingham, Birmingham, UK

³Birmingham Institute of Forest Research, University of Birmingham, Birmingham, UK

⁴Department of Applied Ecology and Environmental Biology, Faculty of Bioscience Engineering, Ghent University, Ghent, Belgium

on species and location, leaf emergence has advanced by 3-8 days for every degree increase in air temperature (Cook et al., 2012; Menzel & Fabian, 1999; Zohner & Renner, 2014). However, a growing body of evidence suggests that this past trend cannot be used to predict future responses because other environmental factors may constrain the future advances in spring phenology (Laube et al., 2014; Polgar, Gallinat, & Primack, 2014; Zohner, Benito, Fridley, Svenning, & Renner, 2017; Zohner, Benito, Svenning, & Renner, 2016). Apart from spring temperature, most trees rely on additional factors, including winter chilling and day length, that are likely to become limiting in the future (Laube et al., 2014; Polgar et al., 2014; Zohner et al., 2016, 2017). Yet, a lack of information about the existence, or relative importance of these drivers, translates to high uncertainty in model predictions of future forest phenology (Basler, 2016). Given that each day advance in spring leaf unfolding of deciduous trees translates to an increase in net ecosystem carbon uptake of 4.5 g C/m² (Keenan et al., 2014), untangling these mechanisms is critical for improving confidence in future climate projections.

Three main factors—autumn temperatures (Fu et al., 2014; Heide, 2003), winter chilling (Laube et al., 2014; Luedeling, Girvetz, Semenov, & Brown, 2011; Yu, Luedeling, & Xu, 2010; Zohner

et al., 2017), and day length (Heide, 1993a, 1993b; Körner & Basler, 2010)—have been proposed to control spring leaf-out by modulating the amount of warming that trees require to leaf-out. These factors serve trees as a safety mechanism to prevent precocious leaf-out in case of an early warm spell when the risk of nightly freezing is still high (Körner & Basler, 2010; Zohner, Mo, Renner, et al., 2020; Zohner, Mo, Sebald, & Renner, 2020). Each of these factors is therefore likely to counteract the advances in spring leaf-out under a warming climate. Specifically, as the climate warms, the accumulated warming required for leaves to emerge is expected to increase because (a) warmer autumn temperatures delay the initiation of dormancy (Fu et al., 2014; Heide, 2003); (b) warmer winters lead to reduced chilling accumulation (Fu et al., 2015; Zohner & Renner, 2014); and (c) days when spring warming occurs are becoming shorter (Fu et al., 2019; Heide, 1993b; Vitasse & Basler, 2013; Zohner & Renner, 2015; Figure 1).

The potential effects of these separate environmental drivers have been identified using controlled climate chamber experiments with pot plants or twig cuttings (Laube et al., 2014; Polgar et al., 2014; Zohner et al., 2016). These studies provide valuable mechanistic insights, but they do not necessarily reflect the behaviour of mature trees under natural growing conditions (Vitasse, 2013).

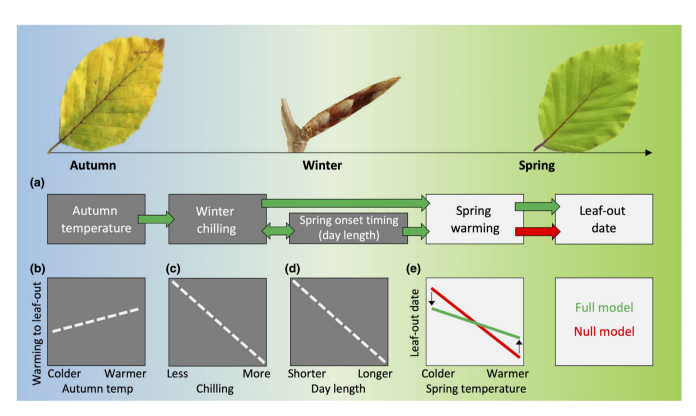


FIGURE 1 Testing for interactive climate effects on the timing of spring leaf-out. (a) In the *full model* (green), the amount of warming required to leaf-out is directly affected by winter chilling and the timing of spring onset (SO; day length when spring warming occurs). In addition, winter chilling interacts with the timing of SO, and autumn temperatures affect winter chilling accumulation. In the *null model* (red), leaf-out is solely driven by spring warming. (b-d) The interactive effects among climate factors should lead to an increase in warming requirements under warmer autumns (b), reduced chilling (c), and an earlier SO (d). (e) Under cold spring conditions, leaf-out should occur earlier than expected from the *null model* because long days and long chilling reduce the amount of warming required to leaf-out; under warm spring conditions, leaf-out should occur later than expected from the *null model* because short days and short chilling increase the amount of warming required to leaf-out

Although the inclusion of these hypothesized mechanisms can improve the performance of mechanistic phenological models, the exact nature, and relative importance, of these mechanisms remain untested under natural conditions (Fu et al., 2019). As such, we cannot represent these mechanisms in global biogeochemical models to predict the consequences for future forest productivity. Parameterizing phenological models and translating their effects into global biogeochemical models require direct empirical evidence about the effects of these dominant environmental drivers in mature trees exposed to real-world changes under natural environmental conditions (Chen, Melaas, Gray, Friedl, & Richardson, 2016).

To represent the important phenological mechanisms into larger biogeochemical models, we need unifying evidence for the strength and direction of these ecological parameters. Empirically testing the influence of these environmental constraints is also vital for avoiding overparameterization in global biogeochemical models, which need to rely on simple submodels to represent plant physiological processes. To date, dynamic global vegetation models, such as LPJ-GUESS, cannot reflect the complex dynamics that are represented in specialized phenology models. As such, they often only account for spring phenology using a simple degree-day-chilling relationship, neglecting the important physiological mechanisms that are likely to restrict the advance of spring phenology in the future. These models are thus likely to vastly overestimate the advances in spring phenology over the rest of the century. Addressing this huge source of uncertainty necessitates that we generate simple empirical parameters for the combined roles of autumn temperature, winter chilling, and day length.

In this study, we aim to bridge the gap between specialized phenological models and global vegetation models by developing a simple, empirical model to evaluate the key mechanisms represented in process-based models. Using a massive in situ database of forest leaf-out observations, we determine the interactive effects of autumn temperature, winter chilling and spring day-length variation on thermal requirements to leaf-out in mature forest trees. We then use the observed relationships to train statistical predictions of future spring arrival. By comparing the performance of this empirical model with all available process-based models from the phenological literature, we show that it adequately reflects the dominant drivers of spring phenology, and predicts spring leaf-out with as much accuracy as existing mechanistic models. In addition, we use forecasts of future temperatures to project the future changes in spring phenology under two climate change scenarios ('CO₂ stabilization' scenario, RCP 4.5 and 'business-as-usual', RCP 8.5). With high confidence in our 'simple' empirical model performance, we could then use the calculated coefficients to train a global dynamic vegetation model to more accurately reflect the future changes in spring phenology. Ultimately, this big-data approach enables us to test the effects of interacting climate drivers, benchmark model projections, and evaluate how these mechanisms influence global dynamic vegetation model predictions of future phenology and global net primary productivity (NPP).

2 | METHODS

2.1 | Data set

In situ observations of leaf-out date were obtained from the Pan European Phenology network (Templ et al., 2018), which provides open-access phenological data for Europe (mainly Germany, Switzerland, and Austria). We selected leaf-out records of 9 common European tree species (7 deciduous angiosperms, 1 deciduous conifer, 1 evergreen conifer) at 4,165 sites (see Figure S1 for site locations). For the seven angiosperms, leaf-out was defined as the date when unfolded leaves, pushed out all the way to the petiole, were visible on the respective individual (BBCH 11, Biologische Bundesanstalt, Bundessortenamt und Chemische Industrie). For the two conifers, *Larix decidua* and *Picea abies*, leaf-out was defined as the date when the first needles started to separate ('mouse-ear stage'; BBCH 10).

Information on temperature parameters was derived from a grid-ded climate data set of daily minimum and maximum temperatures at 0.5° spatial resolution (approximately 50 km; Beer et al., 2014). We additionally tested the CRU/NCAR data set (https://crudata.uea. ac.uk/cru/data/ncep/), which also contains daily minimum and maximum temperatures at 0.5° spatial resolution and obtained very similar results (R^2 for degree-days extracted from Beer et al. [2014] versus CRU/NCAR data set = 0.94). Future predictions of daily maximum and minimum temperatures were based on two different climate warming scenarios (RCP 4.5 and 8.5; Beer et al., 2014).

2.2 | Data cleaning

Following Vitasse, Signarbieux, and Fu (2017), we removed (a) leafout dates that deviated from an individual's median more than three times the median absolute deviation (moderately conservative threshold), (b) leaf-out dates for which the accumulated degree-days deviated from an individual's median degree-days more than three times the median absolute deviation, and (c) individuals for which the standard deviation of phenological observations across years was higher than 15. This data cleaning removed 10% of the data, resulting in a total of 24,650 time-series and 727,401 phenological observations (individuals × years), with a median time-series length of 29 years (minimally 15 years, maximally 63 years).

2.3 | Environmental parameters

Accumulated warming to leaf-out was calculated as the growing degree-days (GDD; using 5°C as base temperature) from 1 January until the date of leaf unfolding. We also tested a temperature threshold of 0°C, which produced very similar results. Here, we only report the results using the threshold of 5°C. To calculate degree-days, we approximated hourly temperature values with a sine curve based on daily maximum ($T_{\rm max}$) and minimum temperatures ($T_{\rm min}$; Equation 1), subtracted 5 (base temperature) from each value, then set all values

below the base temperature to zero (because negative development is biologically not possible), and finally calculated the mean of all 24 values for each day, weighting day-time values (= time when sun is above the horizon) three times more than night-time values. This weighting was done because the effect of day-time temperature on leaf unfolding is ~3 times higher than that of night-time temperature (Fu et al., 2016; Piao et al., 2015).

Winter chilling, reflecting the sum of chilling from 1 October until the mean leaf-out date of each individual, was calculated in two ways (either temperatures below 5°C or between 0 and 5°C) to reflect two possibilities proposed in the literature (Coville, 1920; Fu et al., 2015; Hunter & Lechowicz, 1992). Temperature ($T_{\rm hr}$) at any time of the day (time_{day}) was simulated with a sine curve based on daily maximum ($T_{\rm max}$) and minimum temperatures ($T_{\rm min}$) using the following equation:

$$T_{\rm hr} = \frac{\left(T_{\rm max} - T_{\rm min}\right)}{2} * \sin\left(\frac{\pi}{12} * {\rm time}_{\rm day} - \frac{\pi}{2}\right) + \frac{\left(T_{\rm max} + T_{\rm min}\right)}{2}. \quad (1)$$

This allowed us to calculate the daily proportion of chilling, rather than using a simple presence/absence classification based on daily mean temperatures (e.g. Fu et al., 2015). Multiple studies have reported that temperatures slightly above freezing are most effective in satisfying chilling requirements and assume that effective chilling temperatures range between 0 and 5°C (Coville, 1920; Vitasse et al., 2017):

$$Chill_{hr} = 1 \quad \text{if } 0 \le T \le 5, \tag{2}$$

where chilling (Chill_h,) at any given time of the day depends on the temperature (*T*). We then calculated daily chilling proportions, e.g. a day in which in 75% of the time temperatures are between 0 and 5°C translates to 0.75 chilling days.

In addition, we calculated winter chilling including all temperatures below or equal to 5° C (Fu et al., 2015) as follows:

$$Chill_{hr} = 1 \quad \text{if } T \le 5. \tag{3}$$

To calculate the timing of spring onset (SO) for each year, we first needed to define a date reflecting the onset of spring warming. To do so, for each site and species combination, we calculated the average degree-days accumulating before leaf-out. SO each year was then defined as the day length at the date when the average degree-days to leaf-out at the respective site were reached (Forsythe, Rykiel, Stahl, Wu, & Schoolfield, 1995). SO thus reflects how early spring warming occurred each year:

$$SO = 24 - \frac{24}{\pi} \cos^{-1} \left[\frac{\sin \frac{0.8333\pi}{180} + \sin \frac{L\pi}{180} \sin \varphi}{\cos \frac{L\pi}{180} * \cos \varphi} \right], \tag{4}$$

$$\varphi = \sin^{-1}(0.29795 * \cos \theta),$$
 (5)

$$\theta = 0.2163108 + 2 * tan^{-1} (0.9671396 * tan (0.00860 * (DOY - 186))),$$
 (6)

where L is the latitude of the phenological site and DOY is the day of year when the average degree-days to leaf-out at each site were reached

Autumn temperatures in the year preceding leaf unfolding were calculated as the mean temperatures of the months September and October, September–November, or October and November.

For each species and site, we also analysed the relationship between spring temperatures and leaf-out dates (Figure S2). Spring temperature for each year and individual was defined as the average temperature during the 60 days prior to the average leaf-out date of an individual.

2.4 | Analysis

To test for the importance of autumn temperatures, winter chilling, and spring day length on warming required to leaf-out at each site, we applied univariate regression models over time at the individual level (Figure 2). To visualize the correlations for each species, we removed noise that is due to between-site variation using mixed effects models (R-package Ime4; Figure S3). We calculated chilling in two ways (Equations 2 and 3), and, in all nine species, the effect of chilling on the amount of warming required to leaf-out was significantly higher when choosing the second option (all temperatures ≤5°C satisfy chilling requirements; Figure 2b). To remove possible covariate effects of day length, we also applied partial correlation analyses between winter chilling and spring warming and obtained similar results, i.e. in all nine species, partial correlation coefficients were higher when using all temperatures ≤5°C to calculate winter chilling. Similarly, we tested which temperature period in autumn best predicts the amount of warming required to leaf-out, and, for each time series, the autumn temperature period that yielded the highest correlation coefficient was chosen for multivariate modelling.

We used breakpoint analysis (Richardson et al., 2018), based on the residual sums of squares, to test whether the effect of the timing of SO or winter chilling on required accumulated warming is linear or whether the observed response is flattening beyond a threshold. In 70% and 76% of all time series, a linear model was preferred over a breakpoint model for the effect of the timing of SO and winter chilling, respectively, on required accumulated warming. For the 30% and 25% of time series in which a breakpoint was inferred, we investigated whether steeper slopes are preferred with an earlier arrival of spring warming or decreasing chilling. For the timing of SO, a steeper slope at earlier dates was preferred for only 15% of pixels, while the opposite pattern also was preferred for 15% of pixels. For chilling, a steeper slope under low chilling was only inferred for 13% of pixels, while the opposite pattern was inferred for 11%. We thus rejected the hypothesis that the effect of the timing of SO or winter chilling on the amount of warming required to leaf-out is nonlinear, i.e. increases with earlier arrival of spring warming or decreasing chilling.

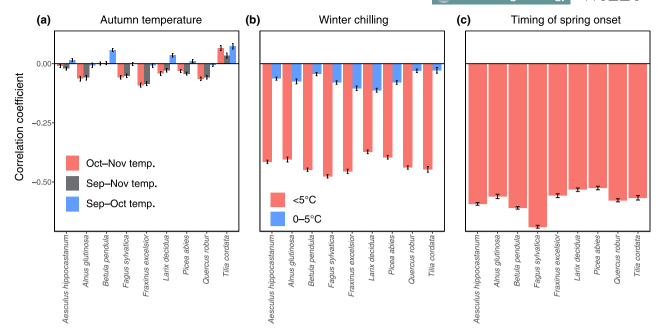


FIGURE 2 The effects of autumn temperature (a), winter chilling (b), and the timing of spring onset (SO; c) on accumulated warming required to leaf-out. Pearson correlation coefficients (±2 SE) are shown for each parameter. (a) The mean temperatures of the months October and November, September-November, or September and October were used to calculate autumn temperatures. (b) Two different temperature ranges were used to calculate winter chilling: all temperatures below 5°C (red) or temperatures between 0 and 5°C (turquoise). (c) The relationship between the timing of SO (day length when spring warming occurs) and accumulated warming required to leaf-out. Number of analysed time series per species: Aesculus hippocastanum, 3,703; Alnus glutinosa, 1,841; Betula pendula, 3,663; Fagus sylvatica, 3,091; Fraxinus excelsior, 2,178; Larix decidua, 2,644; Picea abies, 2,942; Quercus robur, 3,152; Tilia cordata, 1,436

After we had chosen the best autumn period and chilling model for each species, we modelled individual warming requirements using multivariate linear models. Sixteen models were tested against each other (Figure S4a). The models always included winter chilling and day length as fixed effects. Additionally, we either included or excluded autumn temperatures as explanatory variable. We also tested for an interaction term between day length and winter chilling because day-length and chilling cues can interact, with long days substituting for insufficient chilling and vice versa (Vitasse & Basler, 2013; Zohner & Renner, 2015). We also tested models including chilling and the timing of SO as exponential terms (which did not affect model precision and projections; Figure S4). In addition to our multivariate model (hereafter referred to as full model), we applied a chilling model (which has previously been implemented in the LPJ-GUESS dynamic global vegetation model), in which the amount of warming required to leaf-out is solely affected by winter chilling (Equation 7), and a null model, in which leaf-out is solely driven by spring warming (degree-day accumulation) to test for the importance of these individual mechanisms.

By contrast to more complex phenological models, the starting date of degree-day accumulation was not fitted to the observed data and instead fixed to the first day of the year, allowing for easy incorporation into large-scale vegetation models. This also ensures that the *null model* (warming-only model) is not confounded by other factors because fitting a starting date of degree-day accumulation implicitly accounts for winter chilling and/or day length by determining when plants become susceptible to spring warming.

All models were fitted separately to individuals because we were interested in temporal patterns within individuals (rather than spatial patterns among individuals), and spring warming, day-length, and chilling requirements differ among individuals (Zohner, Mo, & Renner, 2018).

2.4.1 | Process-based phenological models

We ran 17 parameterized process-based phenological models from the literature to test the overall performance of our *full model* against existing models. We used the R-package PHENOR (Hufkens, Basler, Milliman, Melaas, & Richardson, 2018) to calibrate the models. Model parameters were optimized using the GenSA algorithm (Xiang, Gubian, Suomela, & Hoeng, 2013), combining both the Boltzmann machine- and faster Cauchy machine-simulated annealing approaches for fast optimizations (Tsallis & Stariolo, 1996). According to Hufkens et al. (2018), the number of iterations was set to 40,000 with a starting temperature of 10,000°C.

2.4.2 | Model evaluation

To judge the performance of phenological models, previous studies either relied solely on root mean-squared errors (RMSEs) of observed versus predicted leaf-out dates (Basler, 2016; Fu, Campioli, Oijen, Deckmyn, & Janssens, 2012; Vitasse, Schneider, Rixen, Christen, & Rebetez, 2018) or additionally

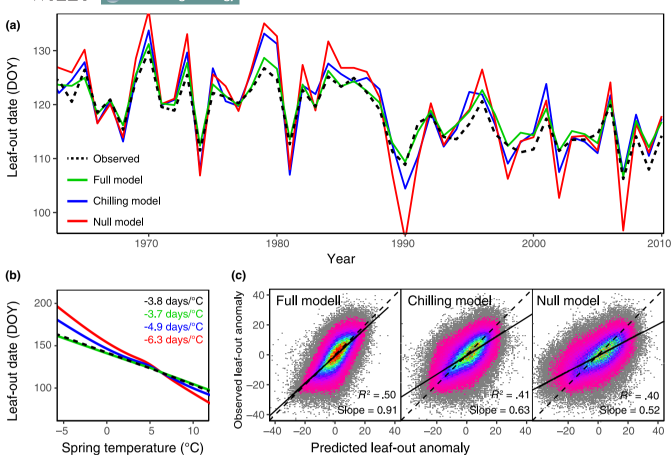
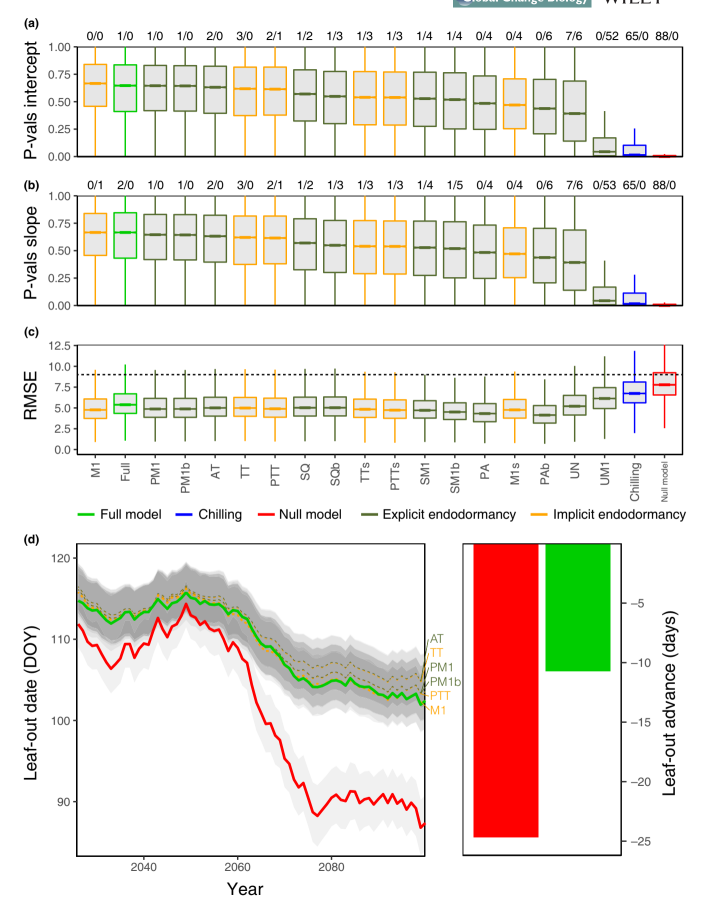


FIGURE 3 Leaf-out date predictions based on the empirical relationships between required accumulated warming and autumn temperature, winter chilling, and the timing of spring onset (see Figure 1). (a, b) Observed and empirically modelled leaf-out dates using tenfold crossvalidations in response to year (a) and spring temperature (b) averaged across all nine study species (observed leaf-out = black lines; full model = green lines; chilling model = blue lines; null model = red lines). See Figures S6 and S7 for species-specific plots. Loess smoothing curves in (b) are based on random effects models to control for differences among sites. (c) Observed versus predicted leaf-out dates of the full model, the chilling model, and the null model. Solid lines show linear regression fit; dashed lines show the 1:1 line. For the chilling model and the null model, the intercept differed significantly from 0 and the slope differed from 1 (p < .05). To standardize among sites, observed and predicted leaf-out dates are shown as anomalies, i.e. as deviation from the mean observed leaf-out date at each site

evaluated model predictions by comparing predicted (in the y-axis) versus observed (in the x-axis) leaf-out dates (Delpierre et al., 2009; Hufkens et al., 2018; Schaber & Badeck, 2003). However, such regression to evaluate models is incorrect, leading to erroneous estimates of the slope and intercept (Piñeiro, Perelman, Guerschman, & Paruelo, 2008). Especially in directional models such as spring phenological projections, where

future climate conditions will lead to ever earlier occurrence dates, models need to be evaluated by analysing intercept and slope components of observed (in the y-axis) versus predicted dates (in the x-axis). To do so, we conducted Wald test-based comparisons (Fox, 2016) using the linear Hypothesis function in the R-package car, allowing us to test for each individual site whether the slopes and intercepts of observed versus predicted

FIGURE 4 Model evaluation and future projections of central European leaf-out dates. (a-c) Model comparison of the three empirical models applied in this study (green = full model, blue = chilling model, red = null model) and 17 process-based models from the literature. (a) Significance values reporting whether the slope of observed versus predicted leaf-out dates differs from 1. Numbers above indicate the percentages of sites for which the model slopes were significantly (p < .05) smaller (= overprediction) or larger than 1 (= underprediction). (b) Significance values reporting whether the intercept of observed versus predicted leaf-out dates differs from 0. Numbers above indicate the percentages of sites for which the model intercepts were significantly larger (= overprediction) or smaller than 0 (= underprediction). (c) Root-mean-squared errors of models. The dashed line shows the average RMSE expected under a null model where leaf-out dates do not differ among years. (d) Future leaf-out projections (15 year moving averages for nine species) under the RCP 8.5 climate scenario, based on the seven best-performing models and the null model. The grey area indicates one SE either side of the mean. Right panel shows estimated advances in leaf-out by the end of the 21st century (2080–2100) compared to the average leaf-out dates between 1990 and 2010 according to the full model (green) and the null model (red)



leaf-out dates differ significantly from 1 and 0, respectively (Figure 4a,b). For each species, we also obtained the overall model fit (R^2 values) and RMSEs for observed versus predicted values (Figures 3c and 4c; Figure S5). Next, we applied tenfold crossvalidations (Stone, 1974), and tested whether projected leaf-out dates capture (a) observed temporal trends and (b) the observed sensitivity of leaf-out dates to spring temperatures (Figure 3a,b; Figures S6 and S7). To calculate temperature sensitivity trends based on time series, we had to remove noise that is due to between-site variation. This was done by adjusting the data using mixed effects modelling available through the R-package lme4.

2.4.3 | Future projections of spring leaf-out

To examine how the analysed ecological mechanisms influence future projections of spring leaf-out, we extrapolated the timing of spring leaf-out until 2100 using two future climate scenarios ('CO₂ stabilization' scenario, RCP 4.5 and 'business-as-usual', RCP 8.5; Figure S8). Specifically, for each scenario, we ran statistical extrapolations of future leaf-out dates, based on the seven best-performing phenology models, including our *full model*, and the simple *null model* accounting solely for temperature accumulation. Future projections of daily minimum and maximum temperatures came from (Beer et al., 2014; Figure S8). Emissions in the RCP 4.5 climate scenario peak around 2040 and then decline. In the RCP 8.5 climate scenario emissions continue to rise throughout the 21st century.

2.4.4 | Land-surface flux projections

We used LPJ-GUESS, a dynamic global vegetation model (Smith et al., 2014), to simulate the effects of shifting spring phenology on temperate and boreal forest NPP. LPJ-GUESS represents vegetation growth and dynamics using a mixture of plant functional types that respond to forcing from the climate (temperature, precipitation, incoming shortwave radiation), atmospheric ${\rm CO_2}$ mixing ratios, and soil type. The successional structure of vegetation is simulated using multiple (here 10) replicate patches in each grid cell, which are subject to stochastic processes of establishment and mortality. Photosynthesis, respiration, stomatal conductance, and phenology in LPJ-GUESS are simulated on a daily time step.

Limitations in availability of the necessary driving data and requirements for parsimony to operate at large scales mean that common process-based phenological models cannot easily be incorporated into global vegetation models such as LPJ-GUESS. Instead, in common with most other such models (Clark et al., 2011; Krinner et al., 2005), spring phenology was represented by an exponential relationship between GDD to leaf-out and the length of the chilling period (chilling model). In LPJ-GUESS, the

relationship was formulated as follows (Sykes, Prentice, & Cramer, 1996):

$$GDD^{\circ} = \alpha + \beta e^{-\kappa C}, \tag{7}$$

where *C* is the length of the chilling period and α , β , and κ are constants specific to plant functional types.

Based on our empirical findings we replaced this equation by the following (*full model*):

$$GDD^{\circ} = \alpha + \beta C + \gamma D + \delta CD, \tag{8}$$

where C is the length of the chilling period, D is the timing of spring warming, CD is the interaction between chilling and the timing of spring warming, and α , β , γ , and δ are coefficients specific to plant functional types (Table S1). The length of the chilling period was defined as the number of days <5°C from 1 October; the timing of spring warming was defined relative to a degree-day threshold (Table S1). We calculated a specific SO for each functional type because the needle-leaved summer-green species Larix decidua, for example, flushes earlier than many broad-leaved summer-green trees. Three functional types of trees (BSI, broad-leaved summer-green shade-intolerant; BST, broad-leaved summer-green shade-tolerant; NS, needle-leaved summer-green) were present in our species sampling. Following Niinemets and Valladares (2006), Fagus sylvatica and Tilia cordata were treated as shade tolerant and Aesculus hippocastanum, Alnus glutinosa, Betula pendula, Fraxinus excelsior, and Quercus robur as shade intolerant. Leaf-out phenology of Picea abies was not included in LPJ-GUESS because, in evergreen species, onset of photosynthetic activity in spring is not dependent on the flushing of new buds. In addition to the deciduous plant functional types described above, LPJ-GUESS simulations also included a temperate needle-leaved evergreen tree, a boreal needle-leaved evergreen shade-tolerant tree, a boreal needle-leaved evergreen shade-intolerant tree, and a C3 grass (Smith et al., 2014), with the distributions of each functional type governed by model internal processes of competition. All simulations were run as potential natural vegetation (i.e. without land management) and the outputs were masked and rescaled to areas above 23°N latitude currently covered by temperate and boreal forest, whereby forest cover was provided by Hansen et al. (2013).

Daily climate forcing data came from the r1i1p1 ensemble member of the IPSL-CM5A-LR model from CMIP5 (Taylor, Stouffer, & Meehl, 2012) for 1850–2099 following the RCP 8.5 scenario, bias-corrected to 1960–1999 WATCH climate (Hempel, Frieler, Warszawski, Schewe, & Piontek, 2013), as prepared for the ISI-MIP2 project. Atmospheric $\rm CO_2$ mixing ratios were as prescribed for the RCP 8.5 scenario of CMIP5, and N deposition data were taken from Lamarque et al. (2013). Simulations were spun up for 500 years using recycled, detrended 1850–1879 climate, and 1850 atmospheric $\rm CO_2$ mixing ratio and N deposition. They were then run under fully transient environmental forcings from 1850 to 2099. The spatial resolution was 0.5° × 0.5°. In total, four simulations were conducted: simulations with the original and updated phenology algorithms, and two further simulations in which, for each of the algorithms, leaf-out

dates from 2010 onwards were forced by mean 2001–2010 daily temperatures in each grid cell, so as to provide a baseline from which to identify the effects of the phenology algorithm on the carbon cycle.

3 | RESULTS

3.1 | The environmental drivers of spring leaf-out

Our linear univariate models showed that, while autumn temperatures had a relatively minor effect, both winter chilling (p < .001; correlation coefficient = 0.4–0.5) and day length (p < .001; correlation coefficient = 0.5–0.7) had consistent negative effects on accumulated warming required to leaf-out across all species (Figure 2; Figure S3). When chilling was calculated using all temperatures below 5°C, the model outperformed an equivalent model in which effective chilling temperatures range between 0 and 5°C (Figure 2b).

The best-performing multivariate model (lowest Akaike information criterion and highest R^2) included chilling and the timing of SO as fixed effects and an interaction between winter chilling and the timing of SO (Figure 3; Figure S4a). Across all species, this *full model* adequately predicted the accumulated warming required to leaf-out across 727,401 observations over 63 years (average R^2 and RMSE = .5 and 5.5, respectively; Figures 3c and 4c; Figures S4a and S5a).

On average, across all species, observed leaf-out dates advanced by 3.8 ± 0.1 days per each degree increase in air temperature. The *full model* performed well in predicting this temperature sensitivity, predicting 3.7 ± 0.2 days/°C. In contrast, the *chilling* and *null model* overestimated leaf emergence, predicting 4.9 ± 0.2 and 6.3 ± 0.2 days/°C, respectively (Figure 3b).

3.2 | Evaluating phenology model performance

Compared to all existing phenology models, our empirical model performed well in predicting leaf emergence over the last 15 years of leaf-out observations, explaining over 50% of the variation in spring leaf emergence over 727,401 observations. This was only marginally worse explanatory power than the best available phenology models (see RMSE values in Figure 4c). Our full model also showed high model accuracy, with predictions fitting close to the 1:1 line in predicted versus observed plots (Figure 4c). As such, the intercept and slope components of observed versus predicted comparisons of leaf-out dates for our full model were among the least likely to differ from 1 and 0, respectively, with a significant (p < .05) deviation only found for <2%of sites (Figure 4a,b). Four of the other process-based models showed an equally low proportion of significant sites with exceptionally high model accuracy. Model accuracy was slightly lower for 11 models (2%-6% significant sites), while the remaining 4 models all performed considerably worse (13%-88% significant sites; Figure 4a,b). The bestperforming model was the M1 model both in terms of model explanatory power and accuracy.

3.3 | Future projections of spring leaf-out

For both climate scenarios, the seven best models (including our *full model*) gave very similar future predictions, estimating a ~60% reduction in the phenological response rates to global warming compared to what would be expected if spring warming was the sole driver of

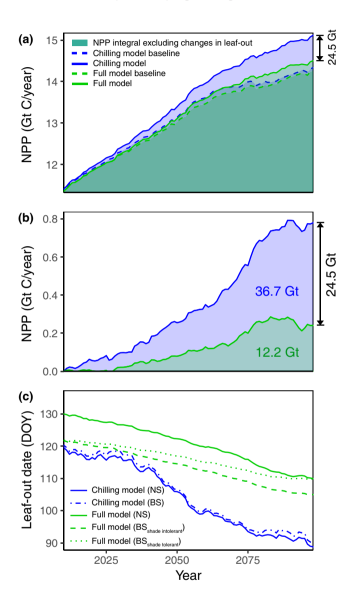


FIGURE 5 Effects of leaf-out changes in Northern Hemisphere deciduous trees on net primary productivity (NPP). (a) Annual forest NPP (above 23°N latitude) over the 21st century, simulating spring leaf-out times with the *chilling model* (solid blue line) or the *full model* (solid green line). Dashed lines show the baselines assuming no leaf-out changes in the future (phenology fixed at years 2001–2010). (b) Increases in NPP that are solely caused by leaf-out shifts simulated with the *chilling model* and the *full model*. Arrows in (a) and (b) show the cumulative difference in NPP between the standard LPJ-GUESS model (including the *chilling model*) and the updated model (including our *full model*). (c) Differences in average leaf-out times of Northern Hemisphere deciduous trees in forests above 23°N latitude simulated with the *chilling model* and the *full model*. Plant functional types: NS, needle-leaved summer-green; BS, broad-leaved summer-green (either shade tolerant or intolerant)

leaf-out phenology (i.e. the *null model*; Figure 4d). While the *null model* predicted a 25 day earlier leaf unfolding by the end of the 21st century under a 'business-as-usual' scenario, the best-performing models estimated advances of only 11 days. Our *full model* projected similar responses for all species, with the exception of *F. sylvatica* (Figure S9), which is expected to advance leaf-out dates less than the other species because pronounced chilling and day-length constraints (Figure 2) cause a lower temperature sensitivity (3.0 days/°C) compared to the other study species (Figures S2 and S7).

3.4 | Changes in forest productivity

The standard LPJ-GUESS model (including a simple chilling–degree-day function to predict spring phenology) estimated that, under a 'business-as-usual' climate scenario, earlier spring arrival will enhance forest NPP above 23°N latitude by ~0.8 Gt carbon per year at the end of the century, resulting in a total increase in cumulative spring NPP of 37 Gt carbon over the rest of the century. In contrast, the updated model, including the new empirically derived information about the ecological constraints on spring phenology (Table S1), estimated that earlier spring arrival will enhance forest NPP by only ~0.2 Gt per year at the end of the century, resulting in a total increase of only 12 Gt over the rest of the century (Figure 5).

4 | DISCUSSION

Our analyses show that, across all nine tested species, winter chilling and the timing of SO have consistent negative effects on the accumulated warming required to leaf-out (Figure 2; Figure S3). In line with previous studies (Heide, 1993b; Vitasse & Basler, 2013; Zohner et al., 2016), European beech showed the strongest sensitivity to chilling and the timing of SO (Figure 2b,c), but the limiting effects of both variables were consistent across all species. As such, although spring warming is likely to increase over the rest of the century, the reductions in winter chilling and the timing of SO are likely to constrain the advance in spring leaf emergence over the rest of the century. These limiting mechanisms may be an important safety strategy against precocious leaf development under future spring climates that overall will be warmer but also more variable, counterintuitively increasing trees' risk of late frost damage to their young leaves in many Eurasian temperate forests (Zohner, Mo, Renner, et al., 2020). In those regions where late frost risk is strongly increasing with climate change, conservative, late-flushing species or populations with pronounced chilling and day-length requirements will be least likely to experience leaf frost damage during spring (Vitasse et al., 2018; Zohner, Mo, Sebald, et al., 2020).

While our findings suggest that the timing of the onset of spring warming represents a strong control on leaf emergence across all nine studied tree species (see Fu et al., 2019 for a more detailed test of this relationship), it remains unclear what is ultimately driving this relationship. A possible explanation for the negative relationship between the

amount of warming required to leaf-out and the 'earliness' of SO is shortening day length. Yet, experimental studies revealed that only in a few species, such as *F. sylvatica*, does day length have an effect on spring leaf-out timing (Laube et al., 2014; Zohner et al., 2016). It is therefore also possible that the time effect we detect here could ultimately be driven by mechanisms other than day length, such as time per se (sensed through an internal clock) or changes in spectral light composition (Brelsford & Robson, 2018). Our results do not give mechanistic insights that would allow us to disentangle the mechanisms by which plants sense the time of the year, but they provide important evidence that both winter chilling and the timing of the onset of spring warming modulate the amount of warming required to leaf-out, thereby restricting future advances in spring leaf-out under climate change.

In contrast to previous suggestions (Fu et al., 2015; Vitasse & Basler, 2013; Vitasse et al., 2017), our results suggest that below-zero temperatures are effective in fulfilling chilling requirements. The model in which chilling was calculated using all temperatures below 5°C outperformed an equivalent model in which effective chilling temperatures ranged between 0 and 5°C (Figure 2b). Our results further show that autumn temperatures have a negligible effect on next year's leaf-out dates (Figure 2a). Yet, autumn temperatures might be of increasing importance in the future if continued autumn warming will further delay the initiation of dormancy, thereby leading to a reduction in winter chilling.

To predict the amount of warming required for each tree to leafout, we ran multivariate models, including all three factors (autumn temperature, winter chilling, and the timing of SO) and the interactions between them. The best model included chilling and the timing of SO as fixed effects, and an interaction between winter chilling and the timing of SO (Figure S4a). This interaction term is supported by experimental studies showing that winter chilling can substitute for day length and vice versa (Heide, 1993a, 1993b; Laube et al., 2014; Zohner et al., 2016; Zohner & Renner, 2015). The coefficients in these empirical models reveal parameters for each of the dominant environmental drivers of spring phenology that are necessary for predicting changes in leaf-out over time.

To test for the importance of these ecological mechanisms, we compared the predictions of our full model (including spring warming, timing of SO, and winter chilling) against similar empirical models that lack these mechanisms. Specifically, we compared the performance of our full-model to a simple 'null model', which included only spring warming, and a 'chilling model' (see Equation 7)-including spring warming and winter chilling—which has previously been implemented in the LPJ-GUESS dynamic global vegetation model. Our full model performed well in predicting the observed temperature sensitivity of 3.8 ± 0.1 days per each degree increase in air temperature, predicting 3.7 ± 0.2 days/°C. In contrast, because they lack the ecological mechanisms that might restrict future advances in spring leaf-out, the chilling and null model overestimated leaf emergence, predicting temperature sensitivities of 4.9 ± 0.2 and 6.3 ± 0.2 days/°C, respectively (Figure 3b). The inclusion of all three mechanisms therefore vastly improved model accuracy, but, more importantly, this reduced the overestimation of spring

leaf-emergence in extremely warm years (Figure 3a). This demonstrates that the combined roles of winter chilling, the timing of SO, and spring warming need to be accounted for in predictions of future tree phenology and productivity.

We also compared the performance of our full model against 17 process-based models from the literature to evaluate whether our full empirical model is capturing the mechanisms in existing state-of-the-art phenology models (Figure 4). We stress that, even though some of these models are called 'ecodormancy models' (suggesting that they solely consider spring warming as a factor), all of these models at least implicitly account for winter chilling-/day length-induced endodormancy release by fitting specific starting dates of degree-day accumulation to the data (we therefore refer to them as explicit or implicit endodormancy models hereafter). Although fitting a specific starting date of degree-day accumulation cannot reflect the gradual transition from endo- to ecodormancy (see e.g. figure 2 in Zohner & Renner, 2015), these models all directly or indirectly represent the ecological mechanisms that we have evaluated in our full model.

By accurately representing the three dominant factors regulating spring leaf-out, our simple empirical model performed as well as the best-performing phenology models. In doing so, our statistical approach can provide a benchmark, revealing which mechanistic models are most accurately representing the ecophysiological mechanisms regulating spring leaf-out. Compared to all existing phenology models, our empirical model had only marginally worse explanatory power than the best available phenology models (Figure 4c) and excelled in terms of model accuracy (intercept and slope components of observed versus predicted leaf-out dates; Figure 4a,b). Four of the other process-based models showed an equally high model accuracy, with the M1 model performing best. This high predictive accuracy of the top 4 process-based models is in direct contrast to previous studies, which suggested low performance across all phenology models (Basler, 2016). This distinction is likely to arise from our focus on model accuracy (i.e. slope estimates) rather than model fit (i.e. RMSE), and the test if predicted values (in the x-axis) reflect observations (in the y-axis), not vice versa (Piñeiro et al., 2008; see Section 2).

Our simple empirical model was trained under current climate conditions, which can lead to uncertainties in future projections if environmental conditions fall outside the model training range. Yet, as expected from the high predictive accuracy of the top models, the seven best models gave very similar future predictions, with our full model and the best-performing M1 model representing the same leaf-out trajectories (Figure 4d). Compared to our null model, in which spring warming was the sole driver of leaf-out phenology, the top models estimated an ~60% or 14 day reduction in the phenological responses to global warming (Figure 4d). This demonstrates that, despite different parameters and assumptions, there is a broad consensus among phenology models-including our full model. As such, our simple regression model can serve to provide basic parameters that can easily be incorporated into large-scale vegetation models and Earth system models to project future terrestrial vegetation carbon dynamics. More complex phenological models rely on spatially explicit parameter-optimization

algorithms to account for endodormancy release. Capturing the spatial variation across temperate and boreal forests would require large amounts of spatially uniform phenological data to train these models. Such data do not currently exist and would require a huge coordinated sampling effort. In contrast, our regression model offers a parsimonious approach, reflecting the main mechanisms triggering spring phenology without the limitations of model overparameterization. This approach can therefore provide projections of increased veracity without inflating structural uncertainty, which remains the main cause of divergence in vegetation model projections (Nishina et al., 2015). Our model can thus provide the empirical relationships that are needed to underpin future projections of spring phenology, and its impacts on terrestrial vegetation carbon dynamics.

To finally comprehend how our leaf-out predictions will affect future projections of NPP, we used a dynamic global vegetation model (LPJ-GUESS). Previously, spring phenology was implemented as a function of degree-days and winter chilling (see chilling model in Figures 3-5; Sykes et al., 1996). We parameterized the phenology algorithm using the empirically derived relationships with the timing of SO, and the updated estimates of winter chilling (Table S1). These changes drastically reduced the projected increases in forest productivity over the rest of this century. Specifically, the standard LPJ-GUESS model (including chilling-only) estimates that cumulative forest NPP will increase over the rest of the century by a total of 37 Gt carbon as a result of earlier spring leaf-out. However, the updated model, including the new empirically derived information about the ecological constraints on spring phenology, estimates an increase of only 12 Gt over the same time period (Figure 5). These differences highlight the need for an improved representation of plant phenology when predicting vegetation dynamics and the terrestrial carbon cycle. The high predictive accuracy of state-of-the-art phenology models we detect here demonstrates that it is possible to adequately represent the main environmental drivers of phenology, and future efforts should thus be directed toward integrating these relevant drivers within boreal, temperate, and tropical ecosystems in global vegetation models.

5 | CONCLUSIONS

Our big-data approach enables us to test the effects of the three main ecological factors—winter chilling, day length, and spring warming—that regulate the timing of spring leaf emergence in temperate and boreal forest trees. A simple statistical model reflecting these interactive ecological drivers performed as well as the best existing phenology models at predicting spring leaf-out over 24,650 individual time series, highlighting that these mechanisms are critical for representing future changes in spring leaf-out. Although spring warming is likely to increase over the rest of the century, the reductions in winter chilling and an earlier timing of spring warming are likely to constrain the future advances in spring leaf emergence. Our statistical model reveals unifying parameters that can be used to represent these important

phenological mechanisms in larger biogeochemical models. By representing this information into a global dynamic vegetation model, we find that the expected increases in temperate forest NPP over the rest of the century are substantially reduced (by 0.6 Gt carbon per year at the end of the 21st century) relative to previous expectations. These results have direct implications for future climate projections, highlighting that forest productivity will be increasingly constrained by factors apart from air temperature in the future.

ACKNOWLEDGEMENTS

This work was supported by grants to C.M.Z. from the ETH Zurich Postdoctoral Fellowship Programme, L.M. from the China Scholarship Council, and T.W.C. from DOB Ecology, Plant-for-the-Planet and the German Federal Ministry for Economic Cooperation and Development. T.A.M.P. acknowledges funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (grant agreement No 758873, TreeMort) (This is paper number 45 of the Birmingham Institute of Forest Research).

CONFLICT OF INTEREST

The authors declare that there are no competing interests.

AUTHOR CONTRIBUTIONS

The study was conceived and developed by C.M.Z. Statistical analysis was performed by L.M. and C.M.Z. LPJ-GUESS simulations were run by T.A.M.P. The manuscript was written by C.M.Z. with assistance from T.W.C. All other authors reviewed and provided input on the manuscript.

DATA AVAILABILITY STATEMENT

All data used for this study are freely available through the Pan European Phenology project (www.PEP725.eu).

ORCID

Constantin M. Zohner https://orcid.org/0000-0002-8302-4854

Jean-Francois Bastin https://orcid.org/0000-0003-2602-7247

REFERENCES

- Basler, D. (2016). Evaluating phenological models for the prediction of leaf-out dates in six temperate tree species across central Europe. Agricultural and Forest Meteorology, 217, 10–21. https://doi.org/10.1016/j.agrformet.2015.11.007
- Beer, C., Weber, U., Tomelleri, E., Carvalhais, N., Mahecha, M., & Reichstein, M. (2014). Harmonized European long-term climate data for assessing the effect of changing temporal variability on land-atmosphere CO₂ fluxes. *Journal of Climate*, 27(13), 4815–4834. https://doi.org/10.1175/JCLI-D-13-00543.1
- Brelsford, C. C., & Robson, T. M. (2018). Blue light advances bud burst in branches of three deciduous tree species under short-day conditions. *Trees*, 32(4), 1157–1164. https://doi.org/10.1007/s00468-018-1684-1
- Chen, M., Melaas, E. K., Gray, J. M., Friedl, M. A., & Richardson, A. D. (2016). A new seasonal-deciduous spring phenology submodel in the

- Community Land Model 4.5: Impacts on carbon and water cycling under future climate scenarios. *Global Change Biology*, 22(11), 3675–3688. https://doi.org/10.1111/gcb.13326
- Chuine, I. (2010). Why does phenology drive species distribution? *Philosophical Transactions of the Royal Society B: Biological Sciences*, 365(1555), 3149–3160. https://doi.org/10.1098/rstb.2010.0142
- Clark, D. B., Mercado, L. M., Sitch, S., Jones, C. D., Gedney, N., Best, M. J., ... Cox, P. M. (2011). The Joint UK Land Environment Simulator (JULES), model description Part 2: Carbon fluxes and vegetation dynamics. Geoscientific Model Development, 4(3), 701–722. https://doi.org/10.5194/gmd-4-701-2011
- Cook, B. I., Wolkovich, E. M., Davies, T. J., Ault, T. R., Betancourt, J. L., Allen, J. M., ... Travers, S. E. (2012). Sensitivity of spring phenology to warming across temporal and spatial climate gradients in two independent databases. *Ecosystems*, 15(8), 1283–1294. https://doi. org/10.1007/s10021-012-9584-5
- Coville, F. V. (1920). The influence of cold in stimulating the growth of plants. *Proceedings of the National Academy of Sciences of the United States of America*, 6(2), 151–160. https://doi.org/10.1175/1520-0493 (1920)48<643b:TIOCIS>2.0.CO;2
- Delpierre, N., Dufrêne, E., Soudani, K., Ulrich, E., Cecchini, S., Boé, J., & François, C. (2009). Modelling interannual and spatial variability of leaf senescence for three deciduous tree species in France. Agricultural and Forest Meteorology, 149(6-7), 938-948. https://doi.org/10.1016/j.agrformet.2008.11.014
- Forsythe, W. C., Rykiel, E. J., Stahl, R. S., Wu, H., & Schoolfield, R. M. (1995). A model comparison for daylength as a function of latitude and day of year. *Ecological Modelling*, 80(1), 87–95. https://doi.org/10.1016/0304-3800(94)00034-F
- Fox, J. (2016). Applied regression analysis and generalized linear models. Hamilton, Canada: McMaster University.
- Fu, Y. H., Campioli, M., Van Oijen, M., Deckmyn, G., & Janssens, I. A. (2012). Bayesian comparison of six different temperature-based budburst models for four temperate tree species. *Ecological Modelling*, 230, 92–100. https://doi.org/10.1016/j.ecolm odel.2012.01.010
- Fu, Y. H., Campioli, M., Vitasse, Y., De Boeck, H. J., Van den Berge, J., AbdElgawad, H., ... Janssens, I. A. (2014). Variation in leaf flushing date influences autumnal senescence and next year's flushing date in two temperate tree species. Proceedings of the National Academy of Sciences of the United States of America, 111(20), 7355–7360. https:// doi.org/10.1073/pnas.1321727111
- Fu, Y. H., Liu, Y., De Boeck, H. J., Menzel, A., Nijs, I., Peaucelle, M., ... Janssens, I. A. (2016). Three times greater weight of daytime than of night-time temperature on leaf unfolding phenology in temperate trees. New Phytologist, 212(3), 590–597. https://doi.org/10.1111/ nph.14073
- Fu, Y. H., Zhang, X., Piao, S., Hao, F., Geng, X., Vitasse, Y., ... Janssens, I. A. (2019). Daylength helps temperate deciduous trees to leaf-out at the optimal time. *Global Change Biology*, 25(7), 2410–2418. https://doi.org/10.1111/gcb.14633
- Fu, Y. H., Zhao, H., Piao, S., Peaucelle, M., Peng, S., Zhou, G., ... Janssens, I. A. (2015). Declining global warming effects on the phenology of spring leaf unfolding. *Nature*, 526, 104. https://doi.org/10.1038/ nature15402
- Hansen, M. C., Potapov, P. V., Moore, R., Hancher, M., Turubanova, S. A., Tyukavina, A., ... Townshend, J. R. G. (2013). High-resolution global maps of 21st-century forest cover change. *Science*, 342(6160), 850– 853. https://doi.org/10.1126/science.1244693
- Heide, O. M. (1993a). Daylength and thermal time responses of budburst during dormancy release in some northern deciduous trees. *Physiologia Plantarum*, 88(4), 531–540.
- Heide, O. M. (1993b). Dormancy release in beech buds (*Fagus sylvatica*) requires both chilling and long days. *Physiologia Plantarum*, 89(1), 187–191. https://doi.org/10.1111/j.1399-3054.1993.tb01804.x

- Heide, O. M. (2003). High autumn temperature delays spring bud burst in boreal trees, counterbalancing the effect of climatic warming. *Tree Physiology*, 23(13), 931–936. https://doi.org/10.1093/treephys/ 23 13 931
- Hempel, S., Frieler, K., Warszawski, L., Schewe, J., & Piontek, F. (2013).
 A trend-preserving bias correction The ISI-MIP approach. Earth
 System Dynamics, 4(2), 219–236. https://doi.org/10.5194/esd-4-219-2013
- Hufkens, K., Basler, D., Milliman, T., Melaas, E. K., & Richardson, A. D. (2018). An integrated phenology modelling framework in r. Methods in Ecology and Evolution, 9(5), 1276–1285. https://doi.org/10.1111/2041-210X.12970
- Hunter, A. F., & Lechowicz, M. J. (1992). Predicting the timing of budburst in temperate trees. *Journal of Applied Ecology*, 29, 597–604. https://doi.org/10.2307/2404467
- Keenan, T. F., Gray, J., Friedl, M. A., Toomey, M., Bohrer, G., Hollinger, D. Y., ... Richardson, A. D. (2014). Net carbon uptake has increased through warming-induced changes in temperate forest phenology. Nature Climate Change, 4(7), 598–604. https://doi.org/10.1038/nclimate2253
- Körner, C., & Basler, D. (2010). Phenology under global warming. *Science*, 327(5972), 1461–1462. https://doi.org/10.1126/science.1186473
- Krinner, G., Viovy, N., de Noblet-Ducoudré, N., Ogée, J., Polcher, J., Friedlingstein, P., ... Prentice, I. C. (2005). A dynamic global vegetation model for studies of the coupled atmosphere-biosphere system. Global Biogeochemical Cycles, 19(1), 1–33. https://doi.org/ 10.1029/2003GB002199
- Lamarque, J.-F., Dentener, F., McConnell, J., Ro, C.-U., Shaw, M., Vet, R., ... Nolan, M. (2013). Multi-model mean nitrogen and sulfur deposition from the atmospheric chemistry and climate model intercomparison project (ACCMIP): Evaluation of historical and projected future changes. Atmospheric Chemistry and Physics, 13(16), 7997–8018. https://doi.org/10.5194/acp-13-7997-2013
- Laube, J., Sparks, T. H., Estrella, N., Höfler, J., Ankerst, D. P., & Menzel, A. (2014). Chilling outweighs photoperiod in preventing precocious spring development. *Global Change Biology*, 20(1), 170–182. https://doi.org/10.1111/gcb.12360
- Luedeling, E., Girvetz, E. H., Semenov, M. A., & Brown, P. H. (2011).
 Climate change affects winter chill for temperate fruit and nut trees. PLoS ONE, 6(5), e20155. https://doi.org/10.1371/journal.pone.0020155
- Menzel, A., & Fabian, P. (1999). Growing season extended in Europe. *Nature*, 397(6721), 659. https://doi.org/10.1038/17709
- Niinemets, Ü., & Valladares, F. (2006). Tolerance to shade, drought, and waterlogging of temperate Northern Hemisphere trees and shrubs. *Ecological Monographs*, 76(4), 521–547. https://doi.org/10.1890/001 2-9615(2006)076[0521:TTSDAW]2.0.CO;2
- Nishina, K., Ito, A., Falloon, P., Friend, A. D., Beerling, D. J., Ciais, P., ... Yokohata, T. (2015). Decomposing uncertainties in the future terrestrial carbon budget associated with emission scenarios, climate projections, and ecosystem simulations using the ISI-MIP results. *Earth System Dynamics*, 6(2), 435–445. https://doi.org/10.5194/ esd-6-435-2015
- Piao, S., Tan, J., Chen, A., Fu, Y. H., Ciais, P., Liu, Q., ... Peñuelas, J. (2015). Leaf onset in the northern hemisphere triggered by daytime temperature. *Nature Communications*, 6(1). https://doi.org/10.1038/ncomms7911
- Piñeiro, G., Perelman, S., Guerschman, J. P., & Paruelo, J. M. (2008). How to evaluate models: Observed vs. predicted or predicted vs. observed? *Ecological Modelling*, 216(3-4), 316-322. https://doi. org/10.1016/j.ecolmodel.2008.05.006
- Polgar, C., Gallinat, A., & Primack, R. B. (2014). Drivers of leaf-out phenology and their implications for species invasions: Insights from Thoreau's Concord. *New Phytologist*, 202(1), 106–115. https://doi.org/10.1111/nph.12647

- Richardson, A. D., Andy Black, T., Ciais, P., Delbart, N., Friedl, M. A., Gobron, N., ... Varlagin, A. (2010). Influence of spring and autumn phenological transitions on forest ecosystem productivity. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 365(1555), 3227–3246. https://doi.org/10.1098/rstb.2010.0102
- Richardson, A. D., Hufkens, K., Milliman, T., Aubrecht, D. M., Furze, M. E., Seyednasrollah, B., ... Hanson, P. J. (2018). Ecosystem warming extends vegetation activity but heightens vulnerability to cold temperatures. *Nature*, 560(7718), 368–371. https://doi.org/10.1038/s41586-018-0399-1
- Richardson, A. D., Keenan, T. F., Migliavacca, M., Ryu, Y., Sonnentag, O., & Toomey, M. (2013). Climate change, phenology, and phenological control of vegetation feedbacks to the climate system. *Agricultural and Forest Meteorology*, 169, 156–173. https://doi.org/10.1016/j.agrformet.2012.09.012
- Schaber, J., & Badeck, F. W. (2003). Physiology-based phenology models for forest tree species in Germany. *International Journal of Biometeorology*, 47(4), 193–201. https://doi.org/10.1007/s00484-003-0171-5
- Smith, B., Wärlind, D., Arneth, A., Hickler, T., Leadley, P., Siltberg, J., & Zaehle, S. (2014). Implications of incorporating N cycling and N limitations on primary production in an individual-based dynamic vegetation model. *Biogeosciences*, 11(7), 2027–2054. https://doi.org/10.5194/bg-11-2027-2014
- Stone, M. (1974). Cross-validatory choice and assessment of statistical predictions. *Proceedings of the Royal Society B: Biological Sciences*, 36(2), 111–174.
- Sykes, M. T., Prentice, I. C., & Cramer, W. (1996). A bioclimatic model for the potential distributions of north European tree species under present and future climates. *Journal of Biogeography*, 23(2), 203–233.
- Taylor, K. E., Stouffer, R. J., & Meehl, G. A. (2012). An overview of CMIP5 and the experiment design. Bulletin of the American Meteorological Society, 93(4), 485-498. https://doi.org/10.1175/ BAMS-D-11-00094.1
- Templ, B., Koch, E., Bolmgren, K., Ungersböck, M., Paul, A., Scheifinger, H., ... Zust, A. (2018). Pan European Phenological database (PEP725): A single point of access for European data. *International Journal of Biometeorology*, 62(6), 1109–1113. https://doi.org/10.1007/s00484-018-1512-8
- Tsallis, C., & Stariolo, D. A. (1996). Generalized simulated annealing. *Physica A: Statistical Mechanics and Its Applications*, 233(1–2), 395–406. https://doi.org/10.1016/S0378-4371(96)00271-3
- Vitasse, Y. (2013). Ontogenic changes rather than difference in temperature cause understory trees to leaf out earlier. *New Phytologist*, 198(1), 149–155. https://doi.org/10.1111/nph.12130
- Vitasse, Y., & Basler, D. (2013). What role for photoperiod in the bud burst phenology of European beech. *European Journal of Forest Research*, 132(1), 1–8. https://doi.org/10.1007/s10342-012-0661-2
- Vitasse, Y., Schneider, L., Rixen, C., Christen, D., & Rebetez, M. (2018). Increase in the risk of exposure of forest and fruit trees to spring frosts at higher elevations in Switzerland over the last four decades. Agricultural and Forest Meteorology, 248, 60-69. https://doi. org/10.1016/j.agrformet.2017.09.005
- Vitasse, Y., Signarbieux, C., & Fu, Y. H. (2017). Global warming leads to more uniform spring phenology across elevations. *Proceedings of the National Academy of Sciences of the United States of America*, 115(5), 1004–1008. https://doi.org/10.1073/pnas.1717342115
- Xiang, Y., Gubian, S., Suomela, B., & Hoeng, J. (2013). Generalized simulated annealing for global optimization: The GenSA Package. *R Journal*, 5(June), 13–28. https://doi.org/10.1007/s10792-010-9404-x
- Yu, H., Luedeling, E., & Xu, J. (2010). Winter and spring warming result in delayed spring phenology on the Tibetan Plateau. *Proceedings of the National Academy of Sciences of the United States of America*, 107(51), 22151–22156. https://doi.org/10.1073/pnas.1012490107
- Zohner, C. M., Benito, B. M., Fridley, J. D., Svenning, J. C., & Renner, S. S. (2017). Spring predictability explains different leaf-out strategies

- in the woody floras of North America, Europe and East Asia. *Ecology Letters*, 20(4), 452–460. https://doi.org/10.1111/ele.12746
- Zohner, C. M., Benito, B. M., Svenning, J. C., & Renner, S. S. (2016). Day length unlikely to constrain climate-driven shifts in leaf-out times of northern woody plants. *Nature Climate Change*, 6(12), 1120–1123. https://doi.org/10.1038/nclimate3138
- Zohner, C. M., Mo, L., & Renner, S. S. (2018). Global warming reduces leaf-out and flowering synchrony among individuals. *eLife*, 7, e40214. https://doi.org/10.7554/eLife.40214
- Zohner, C. M., Mo, L., Renner, S. S., Svenning, J.-C., Vitasse, Y., Benito, B. M., ... Crowther, T. W. (2020). Late spring-frost risk between 1959 and 2017 decreased in North America, but increased in Europe and Asia. Proceedings of the National Academy of Sciences of the United States of America. https://doi.org/10.1073/pnas.1920816117
- Zohner, C. M., Mo, L., Sebald, V., & Renner, S. S. (2020). Leaf-out in northern ecotypes of wide-ranging trees requires less spring warming, enhancing the risk of spring frost damage at cold range limits. *Global Ecology and Biogeography*. https://doi.org/10.1111/geb.13088
- Zohner, C. M., & Renner, S. S. (2014). Common garden comparison of the leaf-out phenology of woody species from different native climates, combined with herbarium records, forecasts long-term

- change. *Ecology Letters*, 17(8), 1016–1025. https://doi.org/10.1111/ele.12308
- Zohner, C. M., & Renner, S. S. (2015). Perception of photoperiod in individual buds of mature trees regulates leaf-out. *New Phytologist*, 208(4), 1023–1030. https://doi.org/10.1111/nph.13510

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Zohner CM, Mo L, Pugh TAM, Bastin J-F, Crowther TW. Interactive climate factors restrict future increases in spring productivity of temperate and boreal trees. *Glob Change Biol.* 2020;00:1–14. https://doi.org/10.1111/gcb.15098