# Why longer seasons with climate change may not increase tree growth

E.M. Wolkovich<sup>1</sup>, Ailene K. Ettinger<sup>2</sup>, Alana Chin<sup>3</sup>, Catherine J. Chamberlain<sup>4</sup>, Frederik Baumgarten<sup>1</sup>, Kavya Pradhan<sup>5,6</sup>, Rubén D. Manzanedo<sup>7–9</sup> & Janneke Hille Ris Lambers<sup>7</sup>

# January 26, 2025

Author contributions: All authors conceived of the idea, prepared the literature review, edited the manuscript and contributed to designing figures; in addition EMW wrote the manuscript, AKE, AC, CJC and EMW synthesized the literature review, RD led developing Fig. 1, CJC developed Fig 2, AC, RD and EMW led developing the Box figure and Fig. 3.

 $<sup>^1</sup>$  Forest & Conservation Sciences, Faculty of Forestry, University of British Columbia, 2424 Main Mall, Vancouver, BC V6T 1Z4, Canada

<sup>&</sup>lt;sup>2</sup> The Nature Conservancy of Washington, 74 Wall Street, Seattle, WA USA

<sup>&</sup>lt;sup>3</sup> California State Polytechnic University, Humboldt. Department of Biological Sciences. Arcata, CA, USA

<sup>&</sup>lt;sup>4</sup> The Nature Conservancy, 334 Blackwell St Ste 300, Durham, NC, USA

<sup>&</sup>lt;sup>5</sup> Department of Biology, University of Washington, Seattle, WA, 98195

<sup>&</sup>lt;sup>6</sup> Moore Center for Science, Conservation International, Arlington, VA, 22202

 $<sup>^7</sup>$  Institute for Integrative Biology, ETH Zürich, 8092 Zürich, Switzerland

 $<sup>^{8}</sup>$  Institute of Plant Sciences, University of Bern, Bern, Switzerland

 $<sup>^{8}</sup>$ Oeschger Center for Climate Change Research, University of Bern, Bern, Switzerland

1 Abstract

A number of recent studies have challenged the assumption that longer growing seasons lead to increased tree growth, raising concerns that forecasts of future climate change—which include increased carbon storage through this assumption—may be overly optimistic. In a review of recent literature, we found that 58% of studies supported the assumption of increased growth with longer seasons, while 36% of studies did not. Diverging results remained when holding methodology constant, which suggests the current major challenge is to understand what underlies this widespread variation. Studies have proposed a suite of hypotheses for why longer growing seasons may not always increase tree growth, including drought-related constraints and internal limits. These hypotheses and their underlying mechanisms, however, were generally tested in different ways by different fields on different species, making comparisons difficult. We outline how bridging these current divides while simultaneously integrating evolutionary history and ecological theory could yield new advances and build a unified model across species for when longer seasons will—or will not—lead to greater tree growth, with major forecasting implications.

#### 16 Introduction

The idea that longer growing seasons lead to increased plant growth is an intuitive tenet across multiple fields of biology, including physiology, dendrochronology and ecosystem ecology (Nobel et al., 1983; Frank et al., 2022). It is also a foundational assumption of many global carbon cycle models (e.g. Ito et al., 2020; Friedlingstein et al., 2022). These models project that continued anthropogenic warming will be partly offset by increased carbon sequestration as warming lengthens growing seasons in many forests (Friedlingstein et al., 2022), an assumption supported by ecosystem-scale studies (Chen et al., 1999; Keenan et al., 2014; Finzi et al., 2020).

Yet recent work has questioned this longstanding assumption (e.g. Dow et al., 2022; Green & Keenan, 2022; Silvestro et al., 2023), with potentially large implications for future climate change. These recent studies challenge decades of research reporting increased growth with longer seasons, from observations along elevational and latitudinal gradients (Myneni et al., 1997; Berdanier & Klein, 2011; King et al., 2013; Cuapio-Hernández et al., 2022), classic experiments in lab settings (Went, 1957), to trends in ecosystem fluxes with warming (Chen et al., 1999; Keenan et al., 2014; Finzi et al., 2020). Proposed mechanisms for the apparent disconnect are diverse (Fig. 1), including the complex nature of climate change (e.g. drought or heat stress, Dow et al., 2022) and internal limits on plant growth (Zohner et al., 2023).

Here we review how different fields have studied the relationship between growing season length and tree growth to identify the potential mechanisms that unite—and could disconnect—these processes. Working across multiple definitions of growth and growing season length (see Box), we find results that suggest substantial variation in growth × season length relationships across species. We also find a pervasive disciplinary split between studies, which often test different mechanisms on different species. Current work often implicitly ignores the importance of

community and phylogenetic ecology to plant growth (e.g. Grime, 1977; Webb et al., 2002; Ávila-Lovera et al., 2023), which could aid the search for a universal model when studying different species. We show how increased cross-disciplinary efforts to build a model across species would allow the field to rapidly develop a framework to predict when, where and how climate change may increase tree growth.

### Evidence that longer seasons increase plant growth, or not

The idea that time limits growth is a fundamental principle across biology. Many biological processes—including photosynthesis and aspects of growth—are rate-limited, making time a crucial commodity (Nobel *et al.*, 1983; Cosgrove, 2005; Hilty *et al.*, 2021). Thus, the hypothesis that longer growing seasons should increase growth is intuitive—and pervasive.

Foundational evidence comes from spatial clines across elevation and latitude, with growth decreasing alongside growing season length at higher elevations and latitudes (Fig. ??). Experimentally, this assumption is supported by small-scale field warming studies that find that phenologically advancing species also grow more with warming (Cleland et al., 2012), while observationally, ecosystem-scale studies have reported a similar positive relationship between season length and carbon fluxes across decades with global warming (Keenan et al., 2014) or in years with warm, early springs (Chen et al., 1999). However, some recent high-profile studies find no support for this relationship (Dow et al., 2022). These studies, which often focus on inter-annual correlations with metrics of standardized individual tree growth (Dow et al., 2022; Silvestro et al., 2023), have generated debate about whether future carbon storage forecasts are overestimated and which metrics of growth (Green & Keenan, 2022), or growing season length (Körner et al., 2023), are relevant (Fig., see Box).

Despite this recent debate, we found that longer seasons lead to increased growth in a slight majority of papers spanning 25 years. Though the number of papers directly addressing this topic is small, we found studies have variously found evidence for—or against—the relationship, with no clear pattern by method or year (Fig. 2 and see 'Literature review methods' in Supplement). For example, carbon assimilation studies were evenly split in finding evidence for or against the relationship (or simply not testing it, Fig. 2). Diverging results were consistently found within methods, suggesting the drivers of this variation are likely due to biological mechanisms, not solely inconsistent definitions of growth or growing season length (as some have recently suggested, e.g. Green & Keenan, 2022; Körner et al., 2023, see also Fig. ).

Most studies tested the hypothesis that longer seasons with climate change increase growth via either increased time to grow (10 of 36 papers) or because longer seasons are usually warmer (8 papers), although many also considered hypotheses that could disconnect growth from season length. Studies from dendrochronology (the study of tree rings and their dating) and physiology have readily offered explanations for findings that increased growth may not be a universal outcome of longer seasons (Fig. 1). External climatic drivers that offset the positive growth effects of longer seasons are often reported in tree ring studies (Kolář et al., 2016; de Sauvage et al., 2022; Camarero et al., 2022). In particular, the hypothesis that higher temperatures paired with lower precipitation produce negative correlations of season length with growth appeared in 58% of tree ring studies we reviewed (and was only mentioned once outside of these studies, see also Fig. 1). In contrast, 43% of lab experimental and wood phenology (xylogenesis) studies suggested fundamental internal constraints that prevent trees from responding to longer seasons (Fig. S1, Cuny et al., 2012; Michelot et al., 2012; Zohner et al., 2023). Yet we found that these hypotheses have been tested in radically different ways on different species, never together, and ignore a suite of relevant research from other disciplines.

# 85 Controllers on growth $\times$ season length relationships

Major mechanisms that could limit or disrupt the positive effects of longer growing seasons generally fall into two categories: (1) external factors, such as drought, which should impact ecosystem-level trends at regional scales, and (2) internal physiological constraints, which some research suggests are either universal across plants (e.g. Zohner et al., 2023), or species- and population-specific (e.g. Soolanayakanahally et al., 2013). While we address each in turn, these drivers clearly operate together. Further, the importance of internal versus external drivers likely varies by species, highlighting the need to integrate perspectives from community and phylogenetic ecology.

#### 94 External drivers

Temperature limits many biological processes. Temperatures that are too cool (below 5°C for temperate trees) and too warm (an area of active research, but likely between 35-45°C; Martinez-Meier et al., 2008; Cabon et al., 2022) slow down biological processes and eventually can lead to tissue death (see Fig. 3a, Larcher, 1980; Kramer, 2012). Between these upper and lower limits, biological processes underpinning growth generally accelerate such that warming can have a direct effect, by accelerating biological time, up until the maximum rate for that particular process. Assuming a common growth response curve to temperature, possible increased growth should be predictable at an ecosystem level based on the current seasonal temperatures and the amount of warming (Fig. 3b).

How much or whether growth increases at all depends on the non-linear effect of temperature on biological processes (Fig. 3a). At very cool temperatures—such as in early spring—a small increase in temperature may have limited effect (or even increase frost risk through early budburst, Fig. 1e, Chamberlain et al., 2021), while an increase at warmer temperatures—such as those more common in the summer (e.g. 16 to 18°C)—could have a larger physiological impact. However, warming that pushes plants beyond their optima, where many biological rates crash, could have large negative impacts (Nobel et al., 1983; Leuning, 2002). Thus, some studies

hypothesize that longer seasons effectively only extend the very cool early-season periods and may have no discernible effect on growth (with varying definitions of growth, see Box), while other studies—based on tree rings—suggest that any increases in growth due to longer seasons can be offset by reduced growth due to high summer temperatures (Fig. 1, Gantois, 2022; Dow et al., 2022). In contrast, other researchers argue that warmer temperatures have not yet pushed trees above their optima (Schaber & Badeck, 2002), and instead have driven increases in growth through accelerated rates, rather than longer seasons (e.g. Ren et al., 2019), or through a combination of both.

Positive effects of longer—or warmer—seasons on growth predicted from temperature responses 119 alone, however, could be counteracted by other external drivers. Moisture deficits from reduced 120 precipitation or higher evaporative demand (commonly invoked in tree ring studies, Fig. 1) 121 can slow or stall growth. Support for this hypothesis comes from negative correlations between 122 growth and precipitation (or other metrics related to plant access to water in tree ring studies, Kolář et al., 2016; Etzold et al., 2022), and is well supported by physiological observations that 124 tree water status can be a biophysical limit to growth (i.e., cells cannot expand without sufficient 125 turgor, Peters et al., 2021; Cosgrove, 2023), though we found few physiological studies on season 126 length that considered this effect (Fig. 2). External biotic factors are also shifting with longer 127 seasons—including herbivory, disease and competition (Mitton & Ferrenberg, 2012; Lange et al., 128 2006; Cleland & Wolkovich, 2024)—and can limit productivity (Sturrock et al., 2011; La Porta et al., 2008; Senf et al., 2017), though they are missing from the current debate on the impacts 130 of longer seasons on growth (we found no mention of them, Fig. 1e). 131

#### 32 Internal constraints

When and how growth is initiated and ceases is under genetic and developmental control, and thus plants' internal programming could limit growth responses to longer seasons (Howe et al., 134 2003). Research has repeatedly shown that populations vary in their growth and its responses to extended seasons (Fig. 1d), reflecting differences in genetic and developmental controls that likely evolved to limit tissue loss to rare early or late-season events (Mitton & Ferrenberg, 2012; 137 Lange et al., 2006; Cleland & Wolkovich, 2024). Populations often vary predictably in their endof-season phenology, with more poleward populations tending to stop height growth (budset) earlier using locally adapted photoperiod cues (Soolanayakanahally et al., 2013; Aitken & Bemmels, 2016). This means longer seasons are generally driven by spring phenology, which appears 141 far more flexible, and has advanced more rapidly than fall events (Aitken & Bemmels, 2016). 142 Some recent studies suggest novel roles for the summer solstice (Zohner et al., 2023) in setting a fixed universal developmental switch between when warming temperatures hasten or delay leaf senescence, and in determining when warmer temperatures trigger greater reproduction (Journé et al., 2024).

Trade-offs between vegetative and reproductive investments may produce important growth response differences across years within individuals, as well as between species. Years of high

reproductive output can reduce growth (Thomas, 2011; Hacket-Pain *et al.*, 2016a). For species that mast—producing abundant cones or fruits in only some years—high reproduction could especially impact measures of wood growth. Higher summer temperatures may trigger masting in the following year (Hacket-Pain *et al.*, 2016a,b); if true, then reduced growth in years following warm summers may not indicate temperatures too high for growth, as recent studies have suggested (e.g. Gantois, 2022; Dow *et al.*, 2022), but instead shifting investment to reproduction.

#### Species-level variation

The effects of these external and internal drivers are likely to vary across species, with species identity strongly predicting variation in growth × season length relationships (e.g. Cuny et al., 2012; Michelot et al., 2012). Though this reality was rarely acknowledged in studies we reviewed (Fig. 1c), research in dendrochronology, physiology and in phenology often mentions important differences between certain species groups that should affect how longer seasons affect growth.

The distinct strategies of deciduous versus evergreen species, including in how and when they 161 invest in leaf and shoot elongation versus cambial growth, can affect how they respond to longer 162 seasons. While evergreen species generally leaf out later than deciduous species they can more immediately photosynthesize with earlier springs, though both types of species generally invest in buds (for new leaves, shoots and flowers) in the preceding year. This means neither can rapidly change their investment in leaf area in response to an earlier spring, but both can have 166 multiple flushes of leaves (Day & Greenwood, 2011; Soolanayakanahally et al., 2013). Wood 167 growth in evergreen species is generally thought to come from current season photosynthesate, 168 while deciduous species may more often use stored carbon resources (Gordon & Larson, 1968; 169 Monson et al., 2018). These differences would suggest season length by growth relationships 170 may be most apparent via lagged effects in deciduous species, but this is rarely studied (and not clearly supported to date Coulthard et al., 2020; Klesse et al., 2023). Further, evergreen species are thought to grow more slowly and thus differences due to season length may be harder to detect (Waring & Franklin, 1979).

This division between evergreen and deciduous species hints at a larger suite of traits that predict growth by growing season length relationships among species. Species that budburst earlier and 176 more readily produce additional leaves (e.g., leaf flushes after budset, and other characteristics 177 more common to 'indeterminate' species, Kikuzawa, 1982; Lechowicz, 1984) may grow more with 178 longer seasons (though potentially with a lag, see Box, Fig.) versus those that budburst later 179 and flush new primary growth only once. Similarly, species adapted to cold, dry or high latitude conditions across their range may have different thresholds for when these external drivers limit 181 or promote growth (e.g., some *Populus* and *Quercus* species, Soolanavakanahally et al., 2013; 182 McKown et al., 2016; Delpierre et al., 2017; de Sauvage et al., 2022). Such differences could 183 easily obscure any overall relationship between growth and growing season length. Supporting this possibility, current studies finding divergent results (Fig. S2, Fig. ??) span a wide range of species (we found 57 species from 26 genera across 36 papers). While this diversity may appear

to make identifying a common relationship between growth and growing season length more difficult, it may instead offer the path to an improved framework.

# Building a new framework for growth imes season length relationships

Useful models of tree growth for climate change forecasting have to include a diversity of different species, while overcoming the challenges of uneven sampling across species and their contrasting 191 responses. Leveraging the diversity of responses observed across species is possible by integrat-192 ing ecological advances in how species traits and evolutionary history shape species responses 193 to climate change (Cornwell & Nakagawa, 2017). In particular, advances in phylogenetic comparative methods (Webb et al., 2002) have moved research away from treating species identity 195 as a simple grouping factor where each species is unique (e.g., Fagus sylvatica is different from 196 Quercus robur and Pinus sylvestris) or fits into a limited set of groups (e.g., deciduous versus 197 evergreen) and towards species as suites of correlated observations, separated by their evolu-198 tionary distance (Fagus sylviatica is much more closely related to Quercus robur compared to 199 Pinus sylvestris). New models built from these advances can fit data from all species at once and layer in species-level information. Traits can capture differences in growth strategies while 201 phylogeny can capture additional species differences, which likely capture unmeasured 'latent' 202 traits (Pearse et al., 2019; Morales-Castilla et al., 2024). 203

In step with these advances, trait ecology has documented leaf and wood economic spectra that suggest major traits to include in these models (with related databases of these traits often available, Chave et al., 2009; Diaz et al., 2016). These 'economics' define a common trade-off along an acquisitive to conservative axis, where some species grow rapidly and more flexibly to take advantage of resources, but are less defended against herbivores and compete poorly at low resource levels, whereas other species compete well at low resource levels, but at the expense of growing slower (Grime, 1977; Chave et al., 2009; Diaz et al., 2016). While these traits likely miss critical components for understanding how growing season length shapes growth, such as when different species invest in shoot and leaf versus wood growth, they provide a baseline from which to build, and a powerful approach to combine data usefully across species. This approach has already been used to identify that early-leafout species often show faster-growing more acquisitive strategies compared to later-leafout species (Cleland & Wolkovich, 2024)—differences that may also impact how they respond to longer seasons.

204

205

206

207

208

210

211

212

213

In addition to naturally organizing species differences, a trait-based phylogenetic comparative approach can help build a more testable and predictable framework. Because this approach can flexibly fit evolutionary history and traits together, it allows clades or species groupings that respond similarly to emerge from the data and models (Davies et al., 2019), versus being a priori grouped or defined. Similarly traits that co-vary with different responses can be more quickly identified (e.g., Willis et al., 2008; Davies et al., 2019, see Fig. 4). Both of these benefits could help identify which species or traits to focus additional studies on to gain the most insights, while

similarly suggesting areas that should be less studied (e.g., traits that may be too confounded with evolutionary history, Cornwell *et al.*, 2014; Westoby *et al.*, 2023) or outlier species that may not represent most species (Morales-Castilla *et al.*, 2024). This approach may thus redefine debates over which metrics of growth or growing season length are relevant into debates over which metrics are most relevant for which clades and/or traits.

Importantly, trait-mediated phylogenetic models aid the search a universal model that can be useful for global forecasts of how growing season length influences growth. By effectively assuming one model, which is then shaped by evolutionary history and trait differences to produce 231 the divergent responses observed across species today, this approach yields an overall estimated 232 growth response alongside a response for each species, which can then be variously combined to 233 scale up. For example, species-level estimates combined with data on species abundance across 234 forests (e.g. Forest Service - United States Department of Agriculture; Fischer et al., 2019) could 235 predict larger-scale metrics, such as satellite observations of phenology and productivity. This framework also provides an organizing method to re-approach the fundamental question of how 237 external and internal drivers shape growth responses to longer seasons, and a way to tackle what we are argue the three most critical questions in this area. 239

# What is the scale of variation in growing season length x growth relationships across space and time?

The idea that growing season length influences plant growth is fundamental to plant biology, but we found it is rarely tested in ways relevant to the current debate (see 'Growth × elevation relationships' in Supplement), providing a major gap that limits progress. While multiple papers report a lack of relationship between growth and growing season length (Figs. 1, 2), we have no fundamental understanding of what the effect size of this relationship should be, and thus no way to know if we have sufficient power in current studies to detect it.

Identifying the macro-scale pattern of how growth and growing season length relate should be a primary goal, and one that seems broadly tractable. Tree ring studies designed to leverage 249 latitudinal and elevational gradients in climate could quickly provide the raw data. Research 250 will then need to develop models that tease out the effects of warmer temperatures across the 251 season—likely affecting important biological rates (Figs. 3)—versus longer seasons. Disentan-252 gling these may require focused efforts to understand xylogenesis across species and climates, 253 but doing so across major climatic gradients could make differences more obvious. Wood growth provides an obvious and tractable baseline from which to set expectations of how much growth 255 can vary across space, and links to existing major datasets (Fig. ??). Research will also then 256 need to integrate beyond wood growth, including methods to better characterize changes across the leaf, shoot and wood architecture of different species (e.g., Puletti et al., 2020; Sillett et al., 258 2024) and also extending to the complexity of roots (McKown et al., 2016; Radville et al., 2016). These data can provide a baseline to compare to the scale of shifts over time, which studies of growth × growing season length to date have focused on (Fig. 2), since the same tree rings measured for understanding spatial variation will also capture inter-annual variation.

#### How prevalent are internal constraints across species and populations?

New evidence suggests inter-annual variation in growth may be limited because of internal con-264 straints that prevent plants from fully using longer seasons (Zohner et al., 2023). If true, this would have major ramifications for how much we expect growth to shift with warming. All plants are limited by internal constraints and how quickly they can build new tissues (Marchand et al., 2021; Luo et al., 2024), but selection towards different growth strategies (e.g., acquisitive 268 versus conservative) should drive variation in these constraints across species. Selection should 269 also drive local adaptation in these constraints at the population-level (McKown et al., 2016; 270 Soolanayakanahally et al., 2013), by favoring individuals that match to local environmental 271 optima (Colautti et al., 2010; McKown et al., 2014). This appears to be the case for budset— 272 which indicates the end of height growth, though we currently have data on only a few species (Aitken & Bemmels, 2016; Zeng & Wolkovich, 2024). Further, new studies could rapidly test for 274 constraints across species and populations, to work towards a predictive framework using phy-275 logeny and traits to predict these constraints. This approach has already yielded useful insights 276 in spring phenology, highlighting which environmental factors consistently drive budburst across 277 species while also showing widely-cited results may not extend beyond one well-studied species (Morales-Castilla et al., 2024).

#### How do external drivers and internal constraints act together?

The external and internal factors that affect how longer seasons impact growth are inherently interconnected (Nobel et al., 1983). While research often acknowledges this, modeling these together will require both experiments and observational studies, ideally designed to integrate into trait-mediated phylogenetic models. Studies across space could provide some inference by studying how growing seasons measured by vegetative versus wood phenology vary—and attributing variation through models that nest species within populations and include traits while also testing for how climate drives growth.

The complexity of climate change and plant growth in response to longer, warmer seasons makes experiments vital to building useful mechanistic models for forecasting. Observational data—used mainly to date to tackle this question (Figs. 2, S1)—generally confounds multiple external drivers, including season length, temperature and precipitation regimes (Ren et al., 2019; Canadell et al., 2021; Camarero et al., 2022), making it impossible to tease out actual drivers behind observed trends. Experiments, in contrast, can provide more robust tests. While juvenile stages of trees are often more flexible than their adult forms, they still usually provide predictable inference in differences across species and populations, and so should be integrated far more into studies of how season length affects growth. Using saplings and controlled environments could quickly test how much growth can shift with longer seasons—providing a

potentially standardized way to compare constraints across species and populations—and then layer on shifts in external drivers.

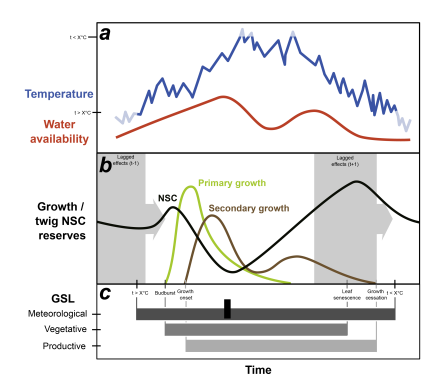
In particular, mechanistic models will need to tease out changes in season length from warming 300 that affects rates; a challenge best addressed by new experiments that decouple these two factors. 301 Such experiments could start on juvenile trees to help inform the underlying model, select 302 representative species to focus on, and develop predictions for large-scale studies. Experiments could also inform a better model of lag effects across species, with small-scale studies sampling saplings multiple years after manipulations (versus the common practice of destructive sampling 305 at the end of the treatment growing season) and large-scale studies following existing efforts to 306 test for ecological 'memory' (e.g. Flinker et al., 2021; Schweiger et al., 2022; Chin et al., 2025). 307 These efforts should help bridge across the contrasting timescales of current physiological and 308 dendrochronological studies of growing season length: we found most physiological studies of 309 growth × growing season length relationships studied 1-2 years of dynamics, usually of juvenile 310 trees, while tree ring studies focused on synthesizing across decades of adult tree growth. 311

Expanding studies across more species will be critical for useful models that can forecast at 312 relevant scales and to help design large-scale experiments. While experimenting on adult trees is difficult, previous challenges in climate change research have led to large-scale experiments to understand other complex drivers (e.g. SPRUCE, DroughtNet, Pfynwald, Norby & Zak, 315 2011; Hanson et al., 2017; Smith et al., 2016). We expect similar experiments will be critical 316 here. Preparing for these large experiments using trait-mediated phylogenetic models to under-317 stand responses across species, however, could advance new experiments well beyond past efforts (Morales-Castilla et al., 2024). By informing which species or clades to study, new experiments could span enough phylogenetic and trait diversity to forecast to species beyond the experiment 320 and maximize the information gained (Cadotte et al., 2017). 321

Conclusions: Anthropogenic climate change has often been described as an unfortunate and unplanned experiment. Like many experiments, it has highlighted important biology we do not know well. Understanding when, how and why longer seasons lead to increased tree growth requires an interdisciplinary reckoning with how temperature, time and a suite of external and internal drivers affect plant growth across species. A mechanistic understanding of how these drivers integrate over species diverse growth strategies and the imprints of evolutionary history to affect growth today is possible, but will require new approaches. Starting now to leverage data across species to inform and design new large-scale studies and experiments be critical for accurate models of future forest and related carbon dynamics, with implications for projections of carbon sequestration and carbon markets.

Acknowledgements: B. Wuu for extracting growth  $\times$  elevation data; R. Zäch for logistical support; N. Pederson for discussion, J. Davies and three reviewers for comments that improved the manuscript.

Box. Growth, growing season length and the challenge of standardized metrics



A major challenge in determining how growth responds to longer growing seasons is the complexity of each, which means that neither can have one simple definition. Here we show the simplified climate of one year (a), which drives variation (b) in primary growth (root and shoot elongation and leaf production from meristems) and secondary growth (radial wood and bark growth from cambia), both of which often depend on conditions determining NSC (non-structural carbohydrate) and storage from previous seasons. Each of these types of growth could define the growing season length (GSL, c) but similarly it can be defined meteorologically (shown here as time, t, above some minimum  $X^{\circ}C$  and below above some maximum  $X^{\circ}C$ , with sufficient soil moisture) or by large-scale measures of plant productivity (Körner et al., 2023). Lagged effects (shown in gray in b) are lasting impacts of previous time periods either in the form of NSC stores or structural legacies influencing productivity (e.g., vessel diameter)

Of studies in our literature review, the largest proportion used metrics related to secondary growth, quantifying growth by measuring radial growth (e.g., through increment cores or dendrometers, n = 28), but a number also looked at metrics related to primary growth, including C assimilation (e.g. net ecosystem productivity or gross primary productivity, n = 20). A smaller number used metrics that reflect combined primary and secondary growth, including biomass, height, or number of stems (n = 9), and root:shoot ratio (n = 1), while others used various mod-

eled estimates potentially related to growth. For growing season length, the largest number of studies used vegetative (e.g., budburst to leaf senescence in our figure above, 26 studies) or wood phenology 11 studies) as their definition, while a smaller number used a meteorological definitions or fixed dates (7 studies). We found 14 that did not directly measure GSL (e.g., Zhu et al., 2021; Dow et al., 2022; Zohner et al., 2023). Further, these definitions of GSL are simplified and could not be easily aligned, as we found 14 different metrics of start of season, 16 metrics of end of season (25 metrics of growing season length. See also 'The challenge of metrics: Measuring growth and growing season length' in the Supplement.

#### 2 1 References

363

- Aitken, S.N. & Bemmels, J.B. (2016) Time to get moving: assisted gene flow of forest trees.

  Evol Appl 9, 271–90.
- <sup>366</sup> Ávila-Lovera, E., Winter, K. & Goldsmith, G.R. (2023) Evidence for phylogenetic signal and correlated evolution in plant—water relation traits. *New Phytologist* **237**, 392–407.
- Berdanier, A.B. & Klein, J.A. (2011) Growing season length and soil moisture interactively constrain high elevation aboveground net primary production. *Ecosystems* **14**, 963–974.
- Cabon, A., Kannenberg, S.A., Arain, A., Babst, F., Baldocchi, D., Belmecheri, S., Delpierre,
   N., Guerrieri, R., Maxwell, J.T., McKenzie, S. et al. (2022) Cross-biome synthesis of source
   versus sink limits to tree growth. Science 376, 758–761.
- Cadotte, M.W., Davies, T.J. & Peres-Neto, P.R. (2017) Why phylogenies do not always predict ecological differences. *Ecological Monographs* 87, 535–551.
- Camarero, J.J., Campelo, F., Colangelo, M., Valeriano, C., Knorre, A., Solé, G. & RubioCuadrado, Á. (2022) Decoupled leaf-wood phenology in two pine species from contrasting
  climates: Longer growing seasons do not mean more radial growth. *Agricultural and Forest Meteorology* 327, 109223.
- Canadell, J., Monteiro, P., Costa, M., Cotrim da Cunha, L., Cox, P., Eliseev, A., Henson, S.,
  Ishii, M., Jaccard, S., Koven, C., Lohila, A., Patra, P., Piao, S., Rogelj, J., Syampungani, S.,
  Zaehle, S. & Zickfeld, K. (2021) Climate Change 2021: The Physical Science Basis. Contribution of Working Group I to the Sixth Assessment Report of the Intergovernmental Panel on
  Climate Change. Cambridge University Press, New York, NY.
- Chamberlain, C.J., Cook, B.I., Morales-Castilla, I. & Wolkovich, E.M. (2021) Climate change reshapes the drivers of false spring risk across european trees. *New Phytologist* **229**, 323–334.
- Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009) Towards a
   worldwide wood economics spectrum. *Ecology Letters* 12, 351–366.
- Chen, W., Black, T., Yang, P., Barr, A., Neumann, H., Nesic, Z., Blanken, P., Novak, M., Eley, J., Ketler, R. *et al.* (1999) Effects of climatic variability on the annual carbon sequestration by a boreal aspen forest. *Global Change Biology* **5**, 41–53.
- Chen, W., Chen, J., Liu, J. & Cihlar, J. (2000) Approaches for reducing uncertainties in regional forest carbon balance. *Global Biogeochemical Cycles* **14**, 827–838.

- Chin, A.R., Gessler, A., Laín, O., Østerlund, I., Schaub, M., Théroux-Rancourt, G., Voggeneder,
- K. & Lambers, J.H.R. (2025) The memory of past water abundance shapes trees 7 years later.
- American Journal of Botany p. e16452.
- <sup>396</sup> Cleland, E.E., Allen, J.M., Crimmins, T.M., Dunne, J.A., Pau, S., Travers, S.E., Zavaleta, E.S.
- & Wolkovich, E.M. (2012) Phenological tracking enables positive species responses to climate
- change. *Ecology* **93**, 1765–1771.
- Cleland, E.E. & Wolkovich, E. (2024) Effects of phenology on plant community assembly and
- structure. Annual Review of Ecology, Evolution, and Systematics 55, 471–492.
- Colautti, R.I., Eckert, C.G. & Barrett, S.C.H. (2010) Evolutionary constraints on adaptive
- evolution during range expansion in an invasive plant. Proceedings of the Royal Society B-
- Biological Sciences 277, 1799–1806, colautti, Robert I. Eckert, Christopher G. Barrett, Spencer
- 404 C. H.
- Cornwell, W. & Nakagawa, S. (2017) Phylogenetic comparative methods. Current Biology 27,
- 406 R333-R336.
- Cornwell, W.K., Westoby, M., Falster, D.S., FitzJohn, R.G., O'Meara, B.C., Pennell, M.W.,
- McGlinn, D.J., Eastman, J.M., Moles, A.T., Reich, P.B. et al. (2014) Functional distinctive-
- ness of major plant lineages. Journal of Ecology 102, 345–356.
- Cosgrove, D.J. (2005) Growth of the plant cell wall. Nature reviews molecular cell biology 6,
- 411 850-861.
- 412 Cosgrove, D.J. (2023) Structure and growth of plant cell walls. Nature Reviews Molecular Cell
- 413 *Biology* pp. 1–19.
- 414 Coulthard, B.L., George, S.S. & Meko, D.M. (2020) The limits of freely-available tree-ring
- chronologies. Quaternary Science Reviews 234, 106264.
- 416 Cuapio-Hernández, L., Reyes-Ortiz, J.L., De La Rosa, A.B., Pavón, N.P., López-Herrera, M.,
- Villanueva-Díaz, J. & Sánchez-González, A. (2022) Is there a response pattern between radial
- growth of trees and elevation gradient? Tree-Ring Research.
- 419 Cuny, H.E., Rathgeber, C.B., Lebourgeois, F., Fortin, M. & Fournier, M. (2012) Life strategies
- in intra-annual dynamics of wood formation: example of three conifer species in a temperate
- forest in north-east france. Tree physiology **32**, 612–625.
- Davies, T.J., Regetz, J., Wolkovich, E.M. & McGill, B.J. (2019) Phylogenetically weighted
- regression: A method for modelling non-stationarity on evolutionary trees. Global ecology and
- biogeography 28, 275–285.
- Day, M.E. & Greenwood, M.S. (2011) Regulation of ontogeny in temperate conifers. Size-and
- age-related changes in tree structure and function, pp. 91–119, Springer.

- de Sauvage, J.C., Vitasse, Y., Meier, M., Delzon, S. & Bigler, C. (2022) Temperature rather
- than individual growing period length determines radial growth of sessile oak in the pyrenees.
- Agricultural and Forest Meteorology 317, 108885.
- Delpierre, N., Guillemot, J., Dufrêne, E., Cecchini, S. & Nicolas, M. (2017) Tree phenological
- ranks repeat from year to year and correlate with growth in temperate deciduous forests.
- Agricultural and Forest Meteorology 234, 1–10.
- Diaz, S., Kattge, J., Cornelissen, J.H.C., Wright, I.J., Lavorel, S., Dray, S., Reu, B., Kleyer, M.,
- Wirth, C., Prentice, I.C., Garnier, E., Bonisch, G., Westoby, M., Poorter, H., Reich, P.B.,
- Moles, A.T., Dickie, J., Gillison, A.N., Zanne, A.E., Chave, J., Wright, S.J., Sheremet'ev,
- S.N., Jactel, H., Baraloto, C., Cerabolini, B., Pierce, S., Shipley, B., Kirkup, D., Casanoves,
- F., Joswig, J.S., Gunther, A., Falczuk, V., Ruger, N., Mahecha, M.D. & Gorne, L.D. (2016)
- The global spectrum of plant form and function. *Nature* **529**, 167–U73.
- Dow, C., Kim, A.Y., D'Orangeville, L., Gonzalez-Akre, E.B., Helcoski, R., Herrmann, V., Harley,
- G.L., Maxwell, J.T., McGregor, I.R., McShea, W.J. et al. (2022) Warm springs alter timing
- but not total growth of temperate deciduous trees. Nature 608, 552–557.
- Etzold, S., Sterck, F., Bose, A.K., Braun, S., Buchmann, N., Eugster, W., Gessler, A., Kahmen,
- A., Peters, R.L., Vitasse, Y. et al. (2022) Number of growth days and not length of the growth
- period determines radial stem growth of temperate trees. *Ecology Letters* **25**, 427–439.
- 445 Finzi, A.C., Giasson, M.A., Plotkin, A.A.B., Aber, J.D., Boose, E.R., Davidson, E.A., Dietze,
- M.C., Ellison, A.M., Frey, S.D., Goldman, E., Keenan, T.F., Melillo, J.M., Munger, J.W.,
- Nadelhoffer, K.J., Ollinger, V. S., Orwig, D.A., Pederson, N., Richardson, A.D., Savage, K.,
- Tang, J., Thompson, J.R., Williams, C.A., Wofsy, S.C., Zhou, Z. & Foster, D.R. (2020) Carbon
- budget of the harvard forest long-term ecological research site: pattern, process, and response
- to global change. Ecological Monographs **90**.
- Fischer, C., Traub, B. et al. (2019) Swiss National Forest Inventory-methods and models of the
- 452 fourth assessment. Springer.
- Flinker, R.H., Cardenas, M.B., Caldwell, T.G., Flerchinger, G.N., Roy, R. & Reich, P.B. (2021)
- 454 Promise and pitfalls of modeling grassland soil moisture in a free-air co2 enrichment experi-
- ment (biocon) using the shaw model. *Pedosphere* **31**, 783–795.
- Forest Service United States Department of Agriculture (????) Forest Inventory and Analysis
- 457 (FIA) database.
- Frank, D., Fang, K. & Fonti, P. (2022) Dendrochronology: Fundamentals and innovations. Stable
- 459 Isotopes in Tree Rings: Inferring Physiological, Climatic and Environmental Responses, pp.
- 460 21–59, Springer International Publishing Cham.

- Friedlingstein, P., O'sullivan, M., Jones, M.W., Andrew, R.M., Gregor, L., Hauck, J., Le Quéré,
- 462 C., Luijkx, I.T., Olsen, A., Peters, G.P. et al. (2022) Global carbon budget 2022. Earth System
- Science Data Discussions **2022**, 1–159.
- 464 Gantois, J. (2022) New tree-level temperature response curves document sensitivity of tree
- growth to high temperatures across a us-wide climatic gradient. Global Change Biology 28,
- 466 6002-6020.
- 467 Gordon, J.C. & Larson, P.R. (1968) Seasonal course of photosynthesis, respiration, and distri-
- bution of 14c in young pinus resinosa trees as related to wood formation. Plant Physiology 43,
- 469 1617-1624.
- Green, J.K. & Keenan, T.F. (2022) The limits of forest carbon sequestration. Science 376, 692–693.
- Grime, J.P. (1977) Evidence for existence of 3 primary strategies in plants and its relevance to ecological and evolutionary theory. *American Naturalist* 111, 1169–1194.
- Hacket-Pain, A., Friend, A., Lageard, J. & Thomas, P. (2016a) Tree rings and masting: con-
- sidering reproductive phenomena when interpreting tree rings? Tree Rings in Archaeology,
- climatology and Ecology 14, 78–85.
- Hacket-Pain, A.J., Cavin, L., Friend, A.D. & Jump, A. (2016b) Consistent limitation of growth
- by high temperature and low precipitation from range core to southern edge of european beech
- indicates widespread vulnerability to changing climate. European Journal of Forest Research
- **135**, 897–909.
- 481 Hanson, P.J., Riggs, J.S., Nettles, W.R., Phillips, J.R., Krassovski, M.B., Hook, L.A., Gu, L.,
- Richardson, A.D., Aubrecht, D.M., Ricciuto, D.M. et al. (2017) Attaining whole-ecosystem
- warming using air and deep-soil heating methods with an elevated co 2 atmosphere. Biogeo-
- sciences **14**, 861–883.
- Hilty, J., Muller, B., Pantin, F. & Leuzinger, S. (2021) Plant growth: The what, the how, and
- the why. New Phytologist **232**, 25–41.
- Howe, G.T., Aitken, S.N., Neale, D.B., Jermstad, K.D., Wheeler, N.C. & Chen, T.H. (2003)
- From genotype to phenotype: unraveling the complexities of cold adaptation in forest trees.
- Canadian Journal of Botany 81, 1247–1266.
- 490 Ito, G., Romanou, A., Kiang, N.Y., Faluvegi, G., Aleinov, I., Ruedy, R., Russell, G., Lerner,
- P., Kelley, M. & Lo, K. (2020) Global carbon cycle and climate feedbacks in the nasa giss
- modele 2. 1. Journal of Advances in Modeling Earth Systems 12, e2019MS002030.
- <sup>493</sup> Journé, V., Szymkowiak, J., Foest, J., Hacket-Pain, A., Kelly, D. & Bogdziewicz, M. (2024) Sum-
- mer solstice orchestrates the subcontinental-scale synchrony of mast seeding. Nature Plants
- **10**, 367–373.

- Keenan, T.F., Gray, J., Friedl, M.A., Toomey, M., Bohrer, G., Hollinger, D.Y., Munger, J.W.,
- O'Keefe, J., Schmid, H.P., Wing, I.S. et al. (2014) Net carbon uptake has increased through
- warming-induced changes in temperate forest phenology. Nature Climate Change 4, 598–604.
- <sup>499</sup> Kikuzawa, K. (1982) Leaf survival and evolution in betulaceae. Annals of Botany **50**, 345–353.
- King, G.M., Gugerli, F., Fonti, P. & Frank, D.C. (2013) Tree growth response along an elevational gradient: climate or genetics? *Oecologia* 173, 1587–1600.
- Klesse, S., Babst, F., Evans, M.E., Hurley, A., Pappas, C. & Peters, R.L. (2023) Legacy effects
- 503 in radial tree growth are rarely significant after accounting for biological memory. Journal of
- Ecology **111**, 1188–1202.
- Kolář, T., Giagli, K., Trnka, M., Bednářová, E., Vavrčík, H. & Rybníček, M. (2016) Response of
- the leaf phenology and tree-ring width of european beech to climate variability. Silva Fennica
- **50**7 **50**.
- Körner, C., Möhl, P. & Hiltbrunner, E. (2023) Four ways to define the growing season. *Ecology*
- Letters .
- 510 Kramer, P. (2012) Physiology of woody plants. Elsevier, New York.
- La Porta, N., Capretti, P., Thomsen, I., Kasanen, R., Hietala, A. & Von Weissenberg, K. (2008)
- Forest pathogens with higher damage potential due to climate change in europe. Canadian
- Journal of Plant Pathology **30**, 177–195.
- Lange, H., Økland, B. & Krokene, P. (2006) Thresholds in the life cycle of the spruce bark beetle
- under climate change. Interjournal for Complex Systems 1648, 1–10.
- Larcher, W. (1980) Plant Physiological Ecology. Springer-Verlag.
- Lechowicz, M.J. (1984) Why do temperate deciduous trees leaf out at different times adaptation
- and ecology of forest communities. American Naturalist 124, 821–842.
- Leuning, R. (2002) Temperature dependence of two parameters in a photosynthesis model. *Plant*,
- 520 Cell & Environment **25**, 1205–1210.
- Luo, Y., Zohner, C., Crowther, T.W., Feng, J., Hoch, G., Li, P., Richardson, A.D., Vitasse, Y. &
- Gessler, A. (2024) Internal physiological drivers of leaf development in trees: Understanding
- the relationship between non-structural carbohydrates and leaf phenology. Functional Ecology
- 524
- Marchand, L.J., Dox, I., Gričar, J., Prislan, P., Van den Bulcke, J., Fonti, P. & Campioli, M.
- (2021) Timing of spring xylogenesis in temperate deciduous tree species relates to tree growth
- characteristics and previous autumn phenology. Tree Physiology 41, 1161–1170.

- Martinez-Meier, A., Sanchez, L., Pastorino, M., Gallo, L. & Rozenberg, P. (2008) What is hot
- in tree rings? the wood density of surviving douglas-firs to the 2003 drought and heat wave.
- Forest Ecology and Management 256, 837–843.
- McKown, A.D., Guy, R.D., Klapste, J., Geraldes, A., Friedmann, M., Cronk, Q.C.B., El-
- Kassaby, Y.A., Mansfield, S.D. & Douglas, C.J. (2014) Geographical and environmental gra-
- dients shape phenotypic trait variation and genetic structure in populus trichocarpa. New
- Phytologist 201, 1263–1276, mcKown, Athena D. Guy, Robert D. Klapste, Jaroslav Geraldes,
- Armando Friedmann, Michael Cronk, Quentin C. B. El-Kassaby, Yousry A. Mansfield, Shawn
- D. Douglas, Carl J.
- McKown, A.D., Guy, R.D. & Quamme, L.K. (2016) Impacts of bud set and lammas phenology on
- root: shoot biomass partitioning and carbon gain physiology in poplar. Trees 30, 2131–2141.
- Michelot, A., Simard, S., Rathgeber, C., Dufrêne, E. & Damesin, C. (2012) Comparing the
- intra-annual wood formation of three European species (Fagus sylvatica, Quercus petraea and
- Pinus sylvestris) as related to leaf phenology and non-structural carbohydrate dynamics. Tree
- physiology **32**, 1033–1045.
- Mitton, J.B. & Ferrenberg, S.M. (2012) Mountain pine beetle develops an unprecedented summer
- generation in response to climate warming. The American Naturalist 179, E163–E171.
- Monson, R.K., Szejner, P., Belmecheri, S., Morino, K.A. & Wright, W.E. (2018) Finding the
- seasons in tree ring stable isotope ratios. American journal of botany 105, 819–821.
- Morales-Castilla, I., Davies, T., Legault, G., Buonaiuto, D., Chamberlain, C.J., Ettinger, A.K.,
- Garner, M., Jones, F.A., Loughnan, D., Pearse, W.D. et al. (2024) Phylogenetic estimates of
- species-level phenology improve ecological forecasting. Nature Climate Change pp. 1–7.
- Myneni, R.B., Keeling, C., Tucker, C.J., Asrar, G. & Nemani, R.R. (1997) Increased plant
- growth in the northern high latitudes from 1981 to 1991. Nature 386, 698–702.
- Nobel, P.S. et al. (1983) Biophysical plant physiology and ecology. WH Freeman and company.
- Norby, R.J. & Zak, D.R. (2011) Ecological lessons from free-air co2 enrichment (face) experi-
- ments. Annual review of ecology, evolution, and systematics 42, 181–203.
- Oleksyn, J., Modrzynski, J., Tjoelker, M., Z. ytkowiak, R., Reich, P.B. & Karolewski, P. (1998)
- Growth and physiology of picea abies populations from elevational transects: common garden
- evidence for altitudinal ecotypes and cold adaptation. Functional Ecology 12, 573–590.
- Pearse, W.D., Legendre, P., Peres-Neto, P.R. & Davies, T.J. (2019) The interaction of phylogeny
- and community structure: Linking the community composition and trait evolution of clades.
- Global Ecology and Biogeography 28, 1499–1511.

- Peters, R.L., Steppe, K., Cuny, H.E., De Pauw, D.J., Frank, D.C., Schaub, M., Rathgeber, C.B.,
- Cabon, A. & Fonti, P. (2021) Turgor—a limiting factor for radial growth in mature conifers 562
- along an elevational gradient. New Phytologist 229, 213–229. 563
- Puletti, N., Grotti, M., Ferrara, C. & Chianucci, F. (2020) Lidar-based estimates of aboveground 564
- biomass through ground, aerial, and satellite observation: a case study in a mediterranean 565
- forest. Journal of Applied Remote Sensing 14, 044501–044501. 566
- Radville, L., McCormack, M.L., Post, E. & Eissenstat, D.M. (2016) Root phenology in a chang-567 ing climate. Journal of Experimental Botany 67, 3617–3628. 568
- Ren, P., Ziaco, E., Rossi, S., Biondi, F., Prislan, P. & Liang, E. (2019) Growth rate rather than
- growing season length determines wood biomass in dry environments. Agricultural and Forest 570
- Meteorology **271**, 46–53. 571
- Rezende, E.L. & Bozinovic, F. (2019) Thermal performance across levels of biological organiza-
- tion. Philosophical Transactions of the Royal Society B 374, 20180549. 573
- Schaber, J. & Badeck, F.W. (2002) Evaluation of methods for the combination of phenological 574 time series and outlier detection. Tree Physiology 22, 973–982.
- 575
- Schweiger, A., Awdamet, F.T., Drenckhan, W. & Berauer, B.J. (2022) Transgenerational effects 576
- of elevated co 2: Downregulation of photosynthetic efficiency and stomatal sensitivity to 577
- drought. 578
- Senf, C., Seidl, R. & Hostert, P. (2017) Remote sensing of forest insect disturbances: Current 579
- state and future directions. International journal of applied earth observation and geoinfor-580
- mation 60, 49-60. 581
- Sillett, S.C., Graham, M.E., Montague, J.P., Antoine, M.E. & Koch, G.W. (2024) Ground-based 582
- calibration for remote sensing of biomass in the tallest forests. Forest Ecology and Management 583
- **561**, 121879. 584
- Silvestro, R., Zeng, Q., Buttò, V., Sylvain, J.D., Drolet, G., Mencuccini, M., Thiffault, N.,
- Yuan, S. & Rossi, S. (2023) A longer wood growing season does not lead to higher carbon 586
- sequestration. Scientific reports 13, 4059. 587
- Smith, B., Wårlind, D., Arneth, A., Hickler, T., Leadley, P., Siltberg, J. & Zaehle, S. (2014) Im-588
- plications of incorporating n cycling and n limitations on primary production in an individual-589
- based dynamic vegetation model. Biogeosciences 11, 2027–2054. 590
- Smith, M.D., Wilcox, K., Sala, O., Phillips, R., Luo, Y., Knapp, A. & Lemoine, N.P. (2016) 591
- Drought-net: A global network merging observations, experiments, and modeling to forecast 592
- terrestrial ecosystem sensitivity to drought. AGU Fall Meeting Abstracts, vol. 2016, pp. B11J-593
- 06. 594

- Soolanayakanahally, R.Y., Guy, R.D., Silim, S.N. & Song, M. (2013) Timing of photoperiodic competency causes phenological mismatch in balsam poplar (populus balsamifera l.). *Plant*, cell & environment **36**, 116–127.
- Sturrock, R., Frankel, S., Brown, A., Hennon, P., Kliejunas, J., Lewis, K., Worrall, J. & Woods,
   A. (2011) Climate change and forest diseases. *Plant pathology* 60, 133–149.
- Thomas, S.C. (2011) Age-related changes in tree growth and functional biology: The role of reproduction. Size- and Age-Related Changes in Tree Structure and Function (eds. F.C. MEINZER, B. Lachenbruch & T.E. Dawson), chap. 2, pp. 33–64, Springer.
- Wang, M., Jiang, Y., Zhang, W., Dong, M., Kang, M. & Xu, H. (2017) Climatic response of
   tracheid features of picea meyeri along altitude gradient of luyashan mountains of north china.
   Polish Journal of Ecology 65, 345–358.
- Waring, R.H. & Franklin, J.F. (1979) Evergreen coniferous forests of the pacific northwest:
   Massive long-lived conifers dominating these forests are adapted to a winter-wet, summer-dry
   environment. Science 204, 1380–1386.
- Webb, C.O., Ackerly, D.D., McPeek, M. & Donoghue, M.J. (2002) Phylogenies and community
   ecology. Annual Review of Ecology and Systematics 33, 475–505.
- Went, F.W. (1957) The experimental control of plant growth. The experimental control of plant growth. 17.
- Westoby, M., Yates, L., Holland, B. & Halliwell, B. (2023) Phylogenetically conservative trait
   correlation: quantification and interpretation. *Journal of Ecology* 111, 2105–2117.
- Willis, C.G., Ruhfel, B., Primack, R.B., Miller-Rushing, A.J. & Davis, C.C. (2008) Phylogenetic
   patterns of species loss in thoreau's woods are driven by climate change. Proceedings of the
   National Academy of Sciences 105, 17029–17033.
- Zeng, Z.A. & Wolkovich, E.M. (2024) Weak evidence of provenance effects in spring phenology
   across europe and north america. New Phytologist .
- Zhou, Y., Yi, Y., Liu, H., Song, J., Jia, W. & Zhang, S. (2022) Altitudinal trends in climate change result in radial growth variation of pinus yunnanensis at an arid-hot valley of southwest china. *Dendrochronologia* 71, 125914.
- Zhu, L., Liu, S., Arzac, A., Cooper, D.J., Jin, Y., Yuan, D., Zhu, Y., Zhang, X., Li, Z., Zhang, Y.
   et al. (2021) Different response of earlywood vessel features of fraxinus mandshurica to rapid
   warming in warm-dry and cold-wet areas. Agricultural and Forest Meteorology 307, 108523.
- Zohner, C.M., Mirzagholi, L., Renner, S.S., Mo, L., Rebindaine, D., Bucher, R., Palouš, D., Vitasse, Y., Fu, Y.H., Stocker, B.D. *et al.* (2023) Effect of climate warming on the timing of autumn leaf senescence reverses after the summer solstice. *Science* **381**, eadf5098.

# **2 Figures**

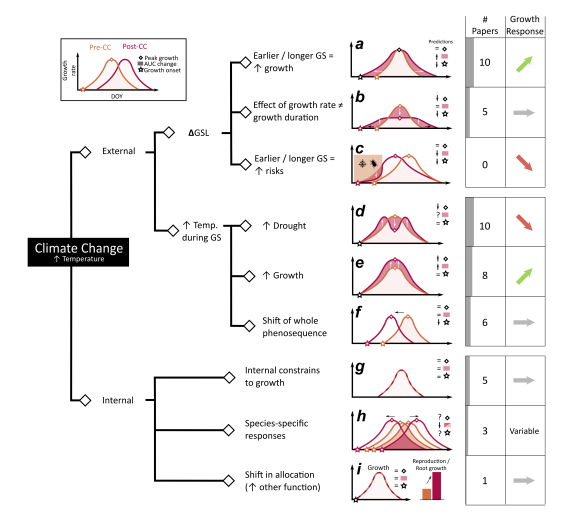


Figure 1: Climate change may alter growing season (GS) length, which can then affect growth through diverse pathways. We review hypotheses for these pathways showing the number of papers (from a review of papers studying growth × growing season length) that mentioned each hypothesis (width of the shaded areas of left column is proportional to the number of papers with the number also given, right column shows the expected growth response for each hypothesis). We group hypotheses as focused on mechanisms moderated by the environment ('external') versus those focused on internal physiological constraints, which span both source (photosynthesis-limited) and sink limitation, and could act together. For more details, see Supplement.

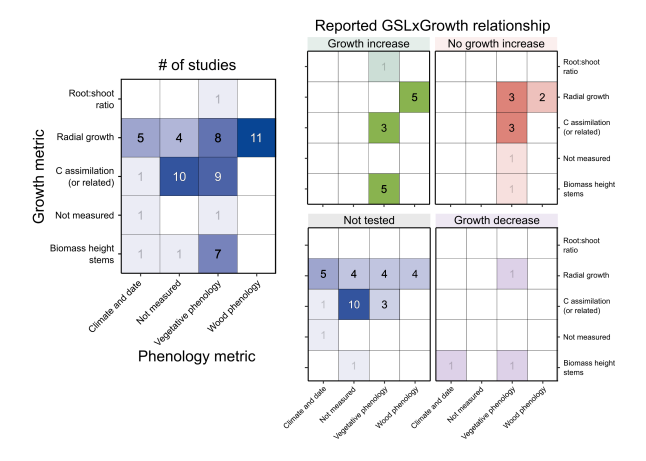


Figure 2: A review of growth  $\times$  growing season length relationship studies spanned a diversity of methods, but there was no coherency in which methods did or did not find a positive relationship. A number of studies tested relationships possibly related to growth  $\times$  growing season length (e.g., they tested how spring temperatures related to growth) but never directly growth  $\times$  growing season length, thus 'not tested' was surprisingly common across methods. See Supplement for review details.

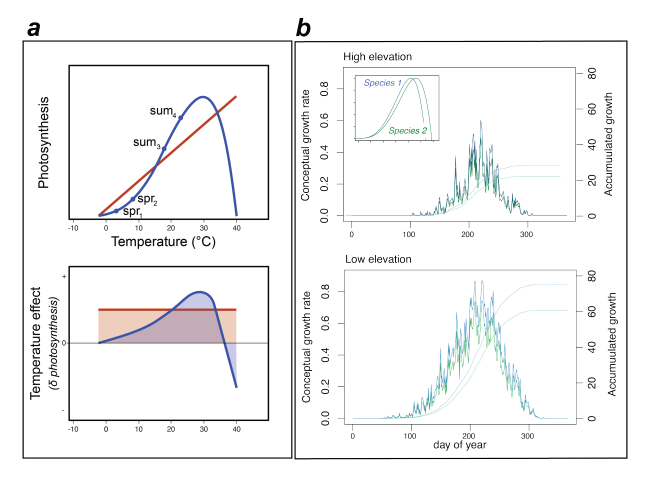


Figure 3: Understanding how longer seasons with climate change affect growth requires teasing out effects of longer seasons versus warmer seasons, which generally co-vary in observational data. Here we show two examples of this complexity. In (a) we show a general net photosynthesis response curve (top panel), which has a non-linear response to temperature (blue curve, adapted from meta-analysis of Rezende & Bozinovic, 2019), though it is often modeled as linear (red). This non-linearity means that increases in lower temperatures—such as those in the spring when much of growing season extensions may happen—have lower absolute increases in photosynthesis compared to increases in later-season (e.g., summer) warmer temperatures, while a linear response assumes a constant scale of effect across low to high temperatures (bottom). In (b) we show conceptual growth responses to temperature for two different species (top, inset) which impacts their growth across the season, leading to small differences in accumulated growth at a conceptual high elevation site (top) versus larger differences in accumulated growth low elevation site (bottom). Testing how growth varies across larger spatial gradients of growing season length, as we conceptualize here (b) could help establish a baseline expectation of the scale of temporal—especially inter-annual variation—and force a greater reckoning with drivers that shift alongside growing season length.

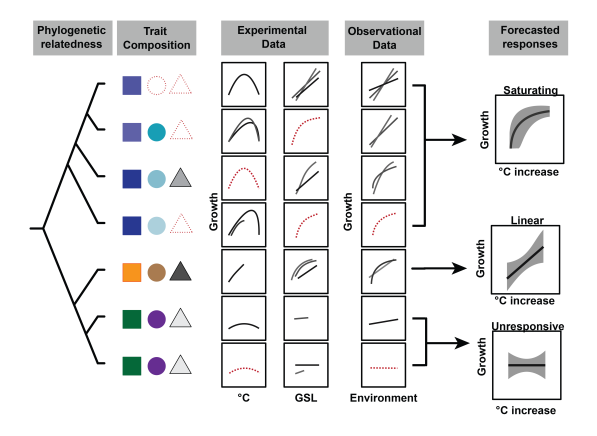


Figure 4: A trait-based phylogenetic model provides a way to naturally organize species (and, not shown, population) responses to predict how they respond to longer seasons. This approach estimates a universal model that is then shaped by species evolutionary history (shown at left via a phylogenetic tree) and traits to estimate how each species should uniquely respond. We argue this framework can organize and guide experiments that separate out changes in temperature from changes in growing season length (°C and GSL in see middle panels) to better integrate observational data and identify different responses by species that can help forecast (see 'Building a new framework for growth × season length relationships' section for more details). In this example, we show how this approach can identify one clade (top) with a common response to longer seasons that also shares a suite of similar traits, and can identify a unique response in by one species in a clade where that species also has a unique trait compared to other species with the same common ancestor (lower clade). Estimating responses across species through a phylogenetic model has the additional benefit of robustly estimating responses given uneven sampling and missing data (the dashed red lines represent that the model will predict a response for each species, informed to various degrees—with the degree determined by the model—by the traits and phylogeny).