

1 Thesis Proposal

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4 **1 Introduction**

5 **1.1 Climate change impacts on tree phenology**

6 An important body of research from the past decades have shown convincing evidence that human activi-  
7 ty are increasingly affecting many worldwide environmental processes (Parmesan2003Towards; Intergov-  
8 ernmental2023Climate; Laurance2006Have;Ceballos2017Biological). This can be through land use change  
9 and destruction, pollution, invasive species, ressource overexploitation and climate change (Wu2013Key;  
10 Driscoll2018A biodiversity; Parmesan1999Poleward). That alone raises major concern and actions have  
11 been deployed to mitigate these impacts, with varying success (e.g. Cambell2014Producing). Even though  
12 immediate actions can have positive impacts and potentially reduce some threats to biodiversity, reversing  
13 150 years of human induced greenhouse gas emissions is harder. These emissions have already affected Earth's  
14 climate and they are projected to continue affecting Earth's climate centuries. While there is a scientific con-  
15 sensus that observed climate change is human-caused (IPCCDetection, Lynas2021Greater, Oreskes2004The  
16 consensus), the magnitude and the extent of the consequences that a warming climate will have on biological  
17 processes are still debatable (Huey2012Predicting). Historically, the first case of attribution of a biological  
18 change to climate change was about poleward shifts of european butterflies in Europe in response to regional  
19 warming (Parmesan1999Poleward).

20 **1.1.2. Trends of spring and autumn phenological events and their drivers** The most frequently  
21 observed biological impact of climate change over the past decades are major changes on spring and au-  
22 tumn phenology — the timing of recurring life history events ? ? ? ? . Understanding the consequences  
23 of these shifts on ecosystems requires understanding how much the growing season has changed ? . Spring  
24 phenological events (e.g. budburst and leafout) have been advancing from 0.5 ? to 4.2 days/decade ? ? and  
25 are mainly driven by temperature ? ? ? . In contrast, autumn phenology (e.g. budset and leaf colouring)  
26 is delayed, though to a much lesser extent than spring ? ? . The drivers regulating autumn phenology are  
27 far less understood than those of spring for many reasons. First, autumn phenology has attracted much  
28 less attention compared to spring ? . Second, the data is often much noisier, since meteorological condi-  
29 tions in the fall can drastically influence phenological phenomena. To illustrate this, trees going through  
30 leaf senescence are subjected to a gradual leaf abscission that follows nutrient reabsorption, and the leaves  
31 within the same individual might be at different senescence stage, but a strong wind spell may trigger leaf  
32 drop for all leaves, thus affecting the temporal resolution of these data (Wu2024Canopy). However, there  
33 is a general belief that autumn phenophases are driven by shortening photoperiod and colder tempera-  
34 tures ? ? ? ? . Different hypothesis are proposed to explain delayed autumn phenophases. First, warmer  
35 autumn temperature may extend the activity of photosynthetic enzymes which could be maintained at a  
36 a higher level. Thus, the degradation rate of chlorophyll would decrease and timing of senescence would  
37 be delayed (Yan2021Divergent). Second, summer droughts could make trees pause their activity schedule  
38 and delay senescence to increase carbon assimilation (Dox2022Severe). Third, there could be an antago-  
39 nistic effect of warming and brightening—caused by reductions in atmospheric pollution and cloud cover  
40 (Sanchez2015Reassessment)—on leaf senescence (Wu2021Atmospheric). Brightening accelerates the leaf  
41 senescence processes and reduces the temperature sensitivity during that period, counteracting the expected  
42 warming-induced delays in leaf senescence. The photo-protection and sink limitation hypothesis provide  
43 plausible explanations for the negative effect of radiation on leaf senescence and the declining effect of tem-

44 perature sensitivity of leaf senescence in response to brightening (Wu2021Atmospheric; Zani2020Increased).

45  
46     **1.1.3. Mis-leading evidence of declining sensitivity to warming** While we have convincing proof  
47 that spring events advanced in the past decades, there is evidence that this might decelerate because of  
48 declining sensitivity to warming (Fu2015Declining; Meng2020Urban). The proposed mechanism is through  
49 the action of warmer winters on tree dormancy. In the fall, trees in boreal and temperate forests slowly  
50 enter dormancy which is initiated with budset. During this phases, cold hardiness increases which prepares  
51 the trees for the upcoming cold temperatures and prevents tissue damage. Then, the trees enter dor-  
52 mancy, during which a certain duration of chilling temperatures—with some interaction with photoperiod  
53 for some species—is necessary for the trees to be ready to accumulate forcing (Vitasse2014TheInteraction).  
54 In the late winter and early spring, they go through two forms of deacclimation before budburst (Vi-  
55 tasse2014TheInteraction). When deacclimation is reached, a certain amount of heat (forcing) is required  
56 to initiate budburst (Fu2015Declining). The argument of declining sensitivity appears here: heat require-  
57 ment is met sooner in warm springs, but it's also negatively correlated with chilling (Yongshuo2015Declining).  
58 However, it is this interaction between chilling and forcing requirements that determines the timing of leaf  
59 unfolding. In other words, a decrease in chilling accumulation is proposed to explain the observed weaker  
60 spring temperature sensitivities, where spring forcing loses of its relative importance (Fu2015Declining;  
61 Meng2020Urban; Wolkovich2021ASimple). However, a meta-analysis compiling 72 studies of 203 species  
62 suggests that declining sensitivities observed in Europe may be a statistical artifact of how these responses  
63 are calculated, thus casting doubt on this proposed trend (Ettinger2020Winter). This statistical artifact  
64 may be caused by using linear models for calculating non-linear processes (Wolkovich2021ASimple).

65     **1.1.2. Mechanisms that could limit growth despite having a longer growing season** Plants  
66 seasonal activity has internal and external controls, both determined by environmental conditions. Internal  
67 controls operate via autonomous clocks, activating genes and releasing hormones which often rely on chilling  
68 and/or photoperiod. The external controls, often referred to as forcing, act directly on the developmental  
69 rate, meristem activity, tissue differentiation and metabolism (Korner2023Four). These controls have often  
70 been overlooked, but are nevertheless critical to our understanding of plant's capacity to adjust their activity  
71 schedule in response to changing conditions (REF). In light of this, I hypothesize two possible drivers that  
72 could explain why a longer growing season might not lead to increased growth: external (environmental)  
73 and internal (via physiological constraints)<sup>?</sup> constraints to growth.

74  
75     *1.1.2.1. External constraints* The complex nature of climate change makes predicting the external limits  
76 to growth hard to quantify at the individual level, as these drivers affect communities as a whole. However,  
77 drought, spring frost and heat waves are commonly mentioned as the main extreme events that could limit  
78 tree growth under climate change <sup>?</sup> <sup>?</sup> <sup>?</sup> <sup>?</sup> <sup>?</sup> <sup>?</sup> <sup>?</sup> <sup>?</sup> . Their respective mechanisms, global trend of occurrence,  
79 consequences and difference among species are described in Tables 3.1. (Spring frosts), 3.2. (Droughts) and  
80 3.3. (Heat waves).

81  
82     *1.1.2.2. Internal constraints* As for the internal constraints to growth, recent hypothesis propose that  
83 broadleaf deciduous tree species may be sink-saturated, such that longer growing season with more carbon  
84 fixation do not necessarily increase growth (Dow2022). This one pathway is directly linked to the internal  
85 controls of plant growth, which are under strong genetic control. Many studies showed, that in addition to  
86 height and radial growth varying across species, these growth responses also vary by site (REF). Populations  
87 from higher altitude or latitude grow less under the same conditions than individuals from lower altitude  
88 or latitude. This is further supported by phenological studies showing that growth end arrives earlier from  
89 populations of higher latitude, demonstrating local adaptation to potentially avoid fall frost, before nutrient  
90 uptake has finished. These trees rely on photoperiod cues for setting buds (stopping height growth)  
91 (WolkovichUnpublishedWhy).

92     Growth seasonality has huge consequences on overall annual growth and so are the environmental conditions  
93 during these periods. For instance, warming spring temperatures seem to positively affect growth, but  
94 recent evidence suggest a shift in this net positive effect near the summer solstice. Indeed Zohner 2023 found  
95 that warmer temperatures before the solstice advanced the onset of senescence, but it slowed the progression  
96 of leaf colouring when temperatures postsolstice were warmer. This further reinforces the importance of  
97 understanding how the timing of warmer conditions affect tree species.

**99 1.1.3. How these shifts translate into effects on trees/forests are not clear - Pros and cons**  
**100 of early/late start/end of season (figure?)** Shifts in spring and autumn phenology support a long-lasting  
**101 and intuitive assumption that earlier spring and delayed autumn events lead to longer seasons—and thus**  
**102 increased growth?** (Stridbeck2022Partly). However, research from the past three years has cast doubt on  
**103 this hypothesis?** . For instance, Dow *DOWet al.* (2022) showed that despite an earlier growth onset,  
**104 neither growth rate nor overall annual increment was increased by longer seasons. This could substantially**  
**105 affect carbon-cycle model projections and thus feedbacks to future climate?**.

**106 Understanding these findings requires answering why trees do not grow more despite longer growing sea-**  
**107 sons. While carbon allocation to above ground biomass is one of the largest carbon sink, how this carbon is**  
**108 allocated into wood is poorly understood. Indeed, the linear relationship assumption between wood growth**  
**109 and carbon assimilation is not well supported mechanistically and represents an important limitation of veg-**  
**110 etation models (Cabon2022). Net primary production represents the difference between photosynthesis and**  
**111 plant respiration, but this commonly used metric completely omits the representation of growth processes.**  
**112 This is perhaps because of a long-lasting paradigm of source-limited photosynthesis. This has now since**  
**113 then been debunked with carbon fertilization experiments, supporting evidence that direct environmental**  
**114 constraints on growth may be stronger than those on photosynthesis (Friend2019On; Parent2010Modelling).**  
**115 This opens up a new, poorly tested, temperature sensitivity between cambial activity and photosynthesis.**  
**116 Growing evidence suggest that cambial activity may be more sensitive than photosynthesis to a range of**  
**117 environmental conditions such as: water and nutrient availability, and temperature (Cabon2022; Muller2011,**  
**118 Peters2021, Cabon2020). This demonstrates that carbon projection models that heavily rely on vegetation**  
**119 alone may mislead the amount of carbon sequestered in our forests.**

**120 1.1.4. Growing season shifts and consequences on forest ecosystems and services** Spring  
**121 and fall phenological events are shifting with debatable consequences on tree growth. The sensitivity of**  
**122 cambial activity to water, temperature and nutrients have the potential to have far-reaching consequences**  
**123 given the hard-to-predict nature of future climate change, where any of these variables could vary from low**  
**124 to high amplitude (Cabon2022; Almagro2025Long-term). This expected asymmetry of future environmental**  
**125 changes makes understanding the internal and external limits to growth critical. Especially, the capacity to**  
**126 tease apart different biomes—as for example boreal vs tropical forests are expected to react differently—is**  
**127 critical and empirical data coming from experiments, but also from observations are paramount to predict**  
**128 the changes of forest carbon offset from human GHG emissions.**

## **129 1.2 Nature of the problem, and how to address it**

**130 1.2.1. Past phenological trends can help (or not) predict future phenological changes** Observed  
**131 phenological trends in the last decades cannot be used directly to extrapolate future phenological changes**  
**132 because: (1) the mechanisms guiding them aren't clear and (2) phenological responses of trees to warming are**  
**133 very likely to not be linear (Fu2013Sensitivity; Ettinger2020Winter). Indeed, accurate predictions require**  
**134 an in-depth accurate mechanistic understanding of phenophases and their sensitivities to environmental**  
**135 drivers, especially to temperature and photoperiod (Fu2013Sensitivity). Therefore, the very foundation of**  
**136 the assumption that longer seasons increase growth may shift with future climate change. The well observed**  
**137 advance in spring phenology may slow down, and delayed fall phenology may shift towards earlier leaf**  
**138 senescence (through summer drought induced growth cessation).**

**139 1.2.2. The assumption that longer seasons lead to increased growth is called into question**  
**140 Recent work shows an absence of increased growth despite apparent better environmental conditions, thus**  
**141 casting doubt on a simple and intuitive positive relationship between growing season length and growth. This**  
**142 paradox further emphasizes the need to understand better the drivers regulating growth across biomes, but**  
**143 also how these drivers vary across the species within these biomes. Phenology vary greatly across species (e.g.**  
**144 closely related species tend to budburst at similar time under similar conditions) (Wolkovich2014Progress),**  
**145 but so does the relationship between growth and season length which may explain the wide variation of**  
**146 this relationship among communities (Buckley2012Functional). This highlights a weakness of certain carbon**  
**147 sequestration models that pool species together, which likely overpass important that nuances in the growth**  
**148 responses that could be explained by species differences. This may mislead the future of forest carbon**  
**149 source-sink ratios?**. To better understand how different species respond to warming, different strategies

150 can be used. Experiments are paramount to robustly tease apart the external vs internal drivers (e.g. warmer  
151 springs from severe drought later in the season—a common co-occurring reality in natural environments)? ? .  
152 However, experiments are most often performed on juvenile trees, which are critical for their role in forest  
153 regeneration projections, but their responses can hardly be translated to mature trees, which hold the  
154 overwhelming carbon biomass proportion of forests? ? ? . Leaf phenology through ground-based observations  
155 can provide valuable insights into the growth onset and end of trees that are not suitable for experimental  
156 trials since cambial and leaf phenology are closely linked to another. It is to say that knowing when leaves  
157 elongate and color, can guide when trees start and stop to grow—fundamental metrics to determine the  
158 growing season length. Ground observations has the advantage of providing accurate data of phenological  
159 events for specific sites and species. Recently, the widespread use of smart-phones has opened a whole  
160 new world of possible phenological through citizen scientists records of data over much larger areas and  
161 for a wider range of species (Piao2019, Dickinson2012, Hufkens2019). While there are drawbacks of these  
162 observations (e.g. non-standard protocols, highly uneven spatial-temporal distribution of these observations),  
163 these methods have a huge potential to diversify the phenology data.

#### 164 **1.2.3. Impacts on carbon source-sink projections**

165 **1.2.4. Goals of my thesis** Using citizen science data, a common garden trial and a large-scale experi-  
166 ment, I aim to better understand how different tree species, at different lifespan stages vary in their growth  
167 responses to different season length.

### 168 **1.3 Complexity of measuring growth and defining growing season length**

#### 169 *What is a growing season?*

170 To understand how trees respond to growing season conditions, it is important to define the growing season  
171 and growth. First, a problem that arises when one tries to quantify how shifting growing seasons affect  
172 growth comes from the definition of the growing season itself. Recently, Korner 2023 proposed four definitions  
173 addressing this issue: 1. true growing season, based on measurable growth; 2. phenological season, based on  
174 visible phenological markers; 3. the productive season, based on primary production and 4. meteorological  
175 season, based on environmental conditions.

176 Here, I will focus on how definition 2., incorporating 4. affects definition 1. as the data collected for this  
177 thesis can't address 3. I will use definition 2. to infer a "window of opportunity" during which meteorological  
178 conditions (4.) will be used to calculate growing degree days (GDD). I am using the meteorological season  
179 within a constrained window, instead of simply using it irrespective of phenology because of the illusion  
180 that an absolute increase in GDD over the last decades—is irrespective of the timing of phenophases—also  
181 increases growth. Springs are warmer, falls are also sometimes warmer, and summers are warmer which  
182 together, increase the number of GDD which may appear to be a reliable proxy for better environmental  
183 conditions. However, a fundamental distinction is that there is a "theoretical" and a "real" period at which  
184 trees can grow, which is usually defined through the period between budburst and leaf senescence, highlighting  
185 the importance of accurate phenology data (REF).

186 Models using degree-days are increasing even though they have been used for decades in agriculture  
187 (e.g. McMaster1997Growing, Gilmore and Rogers1958Heat). These rely on developmental patterns that are  
188 based on temperature dependence to estimate a particular ecological process, in my case, tree-ring width.  
189 These models describe a particular response variable as a composite of time and temperature as opposed  
190 to time alone. This is a parsimonious method that requires three variables: daily minimum and maximum  
191 temperatures and the base temperature at which the process of interest cannot occur (cambial activation in  
192 this case) (McMaster1997Growing-degree; Moore2014Developmental). However, this simplicity comes with a  
193 drawback of over-simplifying potentially complex developmental processes in response to varying environ-  
194 mental conditions within a season (Bonhomme2000Bases).

#### 195 *What is growth?*

196 What is growth? Wood formation (xylogenesis) is the major biological process by which carbon is allocated  
197 and long-term stored in woody plants. Radial growth is determined by the production of xylem and phloem  
198 cells that begins with cambial activation and cell production by cambial initials, following by cell differentia-  
199 tion through the following events: 1. Cell enlargement 2. Secondary-wall formation and lignification and  
200 3. programmed cell death (Silvestro2025From; Etzold2021Number). The rate and duration of these phases  
201 lead irreversible radial growth increment usually represented through tree-rings. In these, secondary xylem

202 cells account disproportionately to the number of cells produced because they divide more than phloem cells  
203 (Rathgeber2016Biological; Plomion2001Wood).

### 205 **1.3.1. Traditional diameter measurements miss the resolution of annual growth increment**

206 Foresters have measured tree diameter and height for decades to infer allometries that could give them a  
207 good estimate of how much wood they could collect in a forest (e.g. Meyer1940A Mathematical; Saun-  
208 ders2008Height). The widely used method in forestry is to measure diameter at breast height at punctual  
209 time intervals (REF). However these measurements don't provide short-term indicators of growth—especially  
210 if taken at multiple years intervals. Therefore, extreme events affecting growth are likely to be missed. In  
211 addition, many forest inventories only report diameter measurements and exclude height, because of logisti-  
212 cal constraints (Saunders2007Height), which reduces biomass estimation quality. This growth data lacks the  
213 temporal resolution necessary to properly infer a relationship between growth and environmental conditions.

214 Alongside diameter-height allometric relationship, dendrochronology and its related fields also have long  
215 standing traditions in forestry and silviculture. Indeed, tree-ring research was developed in the early 1900s  
216 to perform archaeological dating and climate reconstruction (Pearl2020New and e.g. Douglass1928). Since  
217 then, different dendrochronology methods have developed for different purposes, such as the calibration  
218 of the terrestrial radiocarbon curve (Helama2023Distinguishing; Reimer2020The) and modelling Earth's past  
219 climate and ecological change (Pearl2020New). More recently, dendroecology—applications of dendrochrono-  
220 logical techniques to problems in ecology (Fritts1989Dendroecology)—emerged to answer ecological prob-  
221 lems as well as to hindcast (e.g. Bergeron2004Fire) and forecast ecological processes both at the regional  
222 (Gazol2018Forest) and global scale (Manzanedo2019Towards; Büntgen2018). Now, these methods can be  
223 used to understand more precise growth patterns and their relationship with different environmental factors.

### 225 **1.3.2. Growth increment needs to incorporate wood density in order to evaluate how much**

226 **structural carbohydrates were stored within a single year.** While tree-ring width are reliable proxies  
227 to how much trees grow in each year, at relatively low cost and time, the inclusion of wood density in the  
228 analysis may provide data hidden within the tree rings. Indeed Cufar 2008 (Cufar2008Tree-ring) showed  
229 that intra- and inter- annual density variations in beech provided more information and at a higher temporal  
230 resolution than tree-rings widths alone.

231 In addition to densitometry, increasing number of studies are going beyond traditional ring width by  
232 performing analysis using wood anatomy data. For instance, low cost techniques (e.g. "blue intensity" proxy  
233 for latewood density (Babst2016Blue, Campbell2007Blue) and high resolution imaging (Levanič2007Atrics;  
234 Griffin2021Gigapixel; von Arx G 2014Roxas; SkippyWSL; García2022Capturing) give rise to a whole new  
235 world of possibility regarding the microscopic components of wood anatomical features. Micro-anatomical  
236 analyses within tree-rings could be applied to a wide range of applications (Pearl2020New). The character  
237 of annual rings, cell structure, timing of growth and markers for trauma can assist in answering a variety  
238 ecological and physiological questions previously unanswered with ring width or density alone, such as how  
239 growth is affected by growing season length under anthropogenic forcing.

### 240 **1.3.3. Asynchrony between primary and secondary growth (internal growth control?)** I argue

241 that internal growth control in trees may shape their growth responses to growing season length and these  
242 can be split into two categories: primary and secondary growth synchronicity and growth determinancy.

243 Primary (shoot) and secondary (xylem and phloem formation) growth both contribute to how much  
244 carbon is stored in plant tissue, but how much they differ in their responses to environmental drivers is poorly  
245 understood. After a dormancy period, trees will start growing, both vertically (primary) and horizontally  
246 (secondary), but there is extreme variation among species as to when each growth starts, for how long it lasts  
247 and when it stops. For instance, ring-porous species initiate primary growth early in the season, sometimes,  
248 even before budburst (e.g. oaks) (Stridbeck2022Partly), whereas xylogenesis in diffuse-porous species are  
249 usually more synchronized with budburst. These two examples highlight how more complicated it might be  
250 to infer general conclusions as to how growing season shifts may also shift growth, where some species may  
251 extend their primary growth, but restrict their secondary growth and vice versa.

252 In addition to differences in primary and secondary growth synchronicity, the role of internal growth  
253 control—often overlooked—may reshape our understanding of growth responses to growing season length  
254 (BaumgartenUnpublished). In perennial plants, two dichotomous growth strategies are commonly mentioned:  
255 determinate and indeterminate growth, though it may appear as species may exist along a gradient of

256 these (BaumgartenUnpublished; REF). Indeterminate growth is usually associated with short-lived and fast  
257 growth species where these life-history strategies may give them a competing advantage as tissue growth  
258 can be produced quickly in response to changing environmental conditions, but this comes with higher risk  
259 of late spring and early fall frost as well as late droughts (Brienen2020Forest; BaumgartenUnpublished).  
260 At the opposite side of the spectrum, determinate species are usually long-lived and slow-growth and are  
261 mainly constrained by conditions during bud formation, this may increase bud survival at the detriment of  
262 opportunistic growth in face of better-than-expected conditions (BaumgartenUnpublished; REF).

263 Thus primary versus secondary growth allocation, coupled with growth determinancy are variables often  
264 overlooked, but which greatly vary across species and that are likely to affect how they species respond to  
265 shifting season length.

266 **1.3.4. Getting growth temporal resolution is labor-intensive and expensive (e.g. dendrometer costs)** To know when trees start and stop to grow within a single growing season (data not extractable  
267 through tree-rings), there are two methods, one being labor intensive (1) and the other being expensive (2).

268 1. Trehor is a tool that allows recurrent sampling of mature trees where a 2mm microcore is extracted  
269 from the tree a different times during the growing season. By sampling multiple times a single tree, growth  
270 temporality can be inferred by counting the ring cell increment between each sample. While this non-  
271 destructive tool can be extremely valuable, getting large sample size can hardly be feasible (Rossi2006Trehor).

272 2. Dendrometers allow to monitor stem radius variation, measuring irreversible secondary growth, but  
273 also stem water fluctuations and thermal expansion, often leading to biased estimates of growth increment  
274 temporality (Camarero2010Plastic).

## 276 **1.4 Objectives**

277 My objectives for Fuelinex are to assess tree species' potential to prolong or stretch their activity schedule by  
278 artificially manipulating growing season length and analyse how this translates (or nor) into growth, during  
279 the current year (2024) and in the following year (2025). I will also conduct a secondary experiment to  
280 examine whether trees can absorb nutrients late in the season and if that translates into growth during the  
281 following season. For the CookieSpotters projects, I will investigate how the timing of phenological events  
282 affects growth across years for juvenile and mature trees, using observational phenology data and tree ring  
283 images.

284 Together, my two chapters will allow me to address the paradox of the absence of increased growth  
285 despite longer growing seasons.

## 286 **1.5 Research questions**

287 *necessary after the objectives?* Fuelinex: How do extended growing seasons affect tree growth across different  
288 species, both immediately (in the same year as the extended season) and in subsequent years? CookieSpot-  
289 ters: How does phenology regulate tree growth in urban ecosystems?

# 290 **2 Methods**

## 291 **2.1 Fuelinex**

## 292 **2.2 CookieSpotters**

### 293 **Wildchrokie**

- 294 1. Common garden from 2015 to 2023
- 295 2. Four species within the Betulaceae family (Table 2)
- 296 3. Data: phenology, height, tree rings
- 297 4. Analysis: Hierarchical model to understand how tree ring width relates to GDD

### 298 **Treespotters**

- 300      1. Citizen science project from 2015 to today (Table 3)
- 301      2. Tree coring
- 302      3. Data: phenology, tree rings
- 303      4. Analysis: Hierarchical model to understand how tree ring width relates to GDD

304      **3.1. Spring frosts**

<b>Mechanisms</b>	Early warm spells → early leaf out → hard frost (<-2Celsius) → tissue death = loss of photosynthetic capacity? ; Response: second cohort of leaves are more efficient and mitigate carbon sequestration loss?
<b>Global trend of occurrence</b>	Most vulnerable regions are the ones with no past risk of occurrence (); ↑ in Europe and East Asia, but ↓ North America; Global trend is controversial?
<b>Consequences (Individual and Ecosystem level consequences)</b>	Loss of vegetative tissue = ↓ photosynthesis = ↓ and remobilization of NSC to repair damaged tissues = ↓ secondary growth (Meyer24); Loss of reproductive tissue (higher flower mortality) (REF); Costs for orchards and stuff?
<b>Differences across species/provenance</b>	

Table 1: Fuelinex species grouped by tree type, life history, and wood anatomy.

Deciduous Trees			
Common Name (Latin)	Life History Strategy	Wood Anatomy	n (ap- prox)
Bur oak ( <i>Quercus macrocarpa</i> )	Slow-growth, long life	Ring-porous	87
Bitter cherry ( <i>Prunus virginiana</i> )	Fast-growth, short life	Diffuse-porous	78
Box elder ( <i>Acer negundo</i> )	Fast-growth, short life	Diffuse-porous	90
Balsam poplar ( <i>Populus balsamifera</i> )	Fast-growth, short life	Diffuse-porous	84
Paper birch ( <i>Betula papyrifera</i> )	Fast-growth, short life	Diffuse-porous	90
Evergreen Trees			
White pine ( <i>Pinus strobus</i> )	Slow-growth, long life	Non-porous	89
Giant Sequoia ( <i>Sequoiadendron giganteum</i> )	Slow-growth, long life	Non-porous	54

Table 2: Wilchrokic species grouped by tree type, life history, and wood anatomy.

Deciduous Trees			
Common Name (Latin)	Life History Strategy	Wood Anatomy	n
Paper birch ( <i>Betula papyrifera</i> )	Fast-growth, short life	Diffuse-porous	8
Yellow birch ( <i>Betula alleghaniensis</i> )	Moderate-growth, moderate life	Diffuse-porous	21
Grey birch ( <i>Betula populifolia</i> )	Fast-growth, short life	Diffuse-porous	29
Grey alder ( <i>Alnus incana</i> )	Fast-growth, short life	Diffuse-porous	31

Table 3: Treespotters species grouped by tree type, life history, and wood anatomy.

Deciduous Trees			
Common Name (Latin)	Life History Strategy	Wood Anatomy	n
American basswood ( <i>Tilia americana</i> )	Fast-growth, moderate life	Diffuse-porous	5
Eastern cottonwood ( <i>Populus deltoides</i> )	Fast-growth, short life	Diffuse-porous	4
Northern red oak ( <i>Quercus rubra</i> )	Moderate-growth, long life	Ring-porous	4
White oak ( <i>Quercus alba</i> )	Slow-growth, long life	Ring-porous	5
Pignut hickory ( <i>Carya glabra</i> )	Slow-growth, long life	Ring-porous	4
Shagbark hickory ( <i>Carya ovata</i> )	Slow-growth, long life	Ring-porous	4
River birch ( <i>Betula nigra</i> )	Fast-growth, short life	Diffuse-porous	5
Yellow birch ( <i>Betula alleghaniensis</i> )	Moderate-growth, moderate life	Diffuse-porous	4
Sugar maple ( <i>Acer saccharum</i> )	Slow-growth, long life	Diffuse-porous	5
Red maple ( <i>Acer rubrum</i> )	Slow-growth, long life	Diffuse-porous	4
Yellow buckeye ( <i>Aesculus flava</i> )	Moderate-growth, moderate life	Diffuse-porous	5

### 3.2. Drought

	<ul style="list-style-type: none"> <li>— Hot temperature + low precipitation (aka global-change-type drought?) = ↑ evapotranspiration → less water in soil → cavitation → embolism → hydraulic failure? = tissue death? ;</li> <li>— Earlier spring phenology = longer GS → increases vegetative growth → increases evapotranspiration → increases drawdown of soil moisture = progressive water stress?</li> <li>— Long-term vs short-term stomatal responses and consequences on tissue death? ;</li> <li>— Recovery and its determinants? ?</li> </ul>
307	<ul style="list-style-type: none"> <li>— ↑ precipitation anomalies since 1990? ;</li> <li>— Models often exclude PDO/ENSO which limit the capacity to attribute increasing droughts to CC? ;</li> <li>— Weak evidence of detection and attribution of changes in meteorological drought since the mid-20th century? ;</li> <li>— Using a spacial, model-based perspective, anthropogenic forcing increased the frequency, duration and intensity of SPI-based droughts for Americas, Mediterranean, W/S Africa and E Asia? (Marvel2019Twentieth; Hidalgo2009Detection)</li> </ul>
308	<ul style="list-style-type: none"> <li>— Recurring droughts may limit trees' ability to recover from other types of stress.</li> <li>— Tree mortality (e.g. Texas and California extreme droughts are estimated to have killed 300 and 102 million trees? )</li> </ul>
309	

### 3.3. Heat waves

	<b>Definition:</b> Heat wave is a period of excessively hot weather (5 or more consecutive days of prolonged heat in which the daily maximum temperature is higher than the average maximum temperature by 5 °C ), which may be accompanied by high humidity (Marx2021Heat)
310	<b>Mechanisms</b> ↑ atmospheric CO <sub>2</sub> = ↑ temperature → ↑ heat waves... More specifically: A mechanism for the increase occurrence of heat waves is a weakening of the polar jet stream (important weather factor for middle latitude regions of North America, Europe and Asia) caused by global warming which increases the occurrence of stationary weather, resulting in heavy rain falls or heat waves(Marx2021Heat). Extreme heat → growth either through (1) Directly via cell processes disruption or (2) indirectly via effects of rising leaf-to-air vapor deficit (VPD) (Gagne2020Limited). Increased temperature leads to reduced photosynthesis which can be attributed to: 1. Damage to photosynthetic machinery 2. Inactivation of RUBISCO 3. Reduction to RuBP regeneration 4. Membrane stability 5. Increased mitochondrial respiration and photorespiration (Hauck2025Heat)
	<b>Global trend of occurrence</b> Heat waves have increased (Meehl2004More;Gagne2020Limited; Teskey2015Responses) and are expected to increase under future climate change (Teskey2015Responses; Dosio2018Extreme;IPCC2014). Summertime extreme temperatures associated with prolonged heat waves, lasting for several weeks, now impact approximately 10% of land surfaces, up from only 1% in the 1960s. (Teskey2015Responses). The more intense and more frequently occurring heat waves cannot be explained solely by natural climate variations and without human-made climate change (Marx2021Heat).
311	<b>Consequences (Individual and Ecosystem level consequences)</b> - Reduced photosynthesis - Increased mortality - Photosynthetic tissue loss (Gagne2020Limited)
	<b>Differences across species/provenance</b> Some species have thermal photosynthetic/respiratory acclimation while others don't. Growth and survival will change depending on species to thermally acclimate to both photosynthesis and respiration - This is explained by growth strategies of gymnosperms vs angiosperms (which are usually better)

### 3 References