Version dated: July 19, 2020

Running title: Tracking & climate change

How temporal tracking shapes species and communities in stationary and non-stationary environments

E. M. Wolkovich 1 & M. J. Donahue 2

Corresponding author: see¹ above; Ph: 604.827.5246 (no fax).

Authorship statement: EMW and MJD both conceived of the paper, performed modeling work and edited the paper, EMW wrote the paper and did the literature review, while MJD wrote the supplementary information on the model.

Data statement: Review, so no new primary data, but data from a comprehensive literature review will be archived in an appropriate public repository and the data DOI will be included at the end of the article.

Keywords: community assembly, global change, climate change, phenology, environmental variability

Article type: Reviews and Syntheses

Article information: Abstract: 190 words; Main text: 5,272; Figures: 4; Boxes: 4 (text in Box 1: 343; Box 2: 856; Box 3: 264, Box 4: 727); 115 references (max of 100)

¹ Forest & Conservation Sciences, Faculty of Forestry, University of British Columbia, 2424 Main Mall, Vancouver, BC V6T 1Z4 (e.wolkovich@ubc.ca)

² Hawai'i Institute of Marine Biology, University of Hawai'i at Mānoa, Kān'eohe, HI 96744 (donahuem@hawaii.edu)

1 Abstract

Climate change reshapes the environments of all species. Predicting species responses requires understanding how well species can track environmental change, and how such tracking shapes communities. Growing empirical evidence suggests that environmental temporal tracking—how an organism shifts the timing of major biological events in response to proximate environmental cues—is linked to species performance and is a structuring force of species and communities today. Here, we review the concept of tracking in empirical studies and through the lens of community ecology theory. We argue that much current theory for tracking ignores the importance of a multi-species context beyond trophic interactions. Yet community assembly theory predicts competition should drive variation in tracking and trade-offs with other traits. Existing community assembly theory can help us understand tracking in stationary and non-stationary systems and, thus, predict the species-and community-level consequences of climate change. But it will require greater efforts to integrate priority effects into modern coexistence theory and improved empirical estimates of tracking and the underlying environmental cues.

16 1 Main text

Anthropogenic climate change is causing widespread changes in species distributions, with many species shifting in both time and space (IPCC, 2014; Hoegh-Guldberg et al., 2018). Species are 18 moving to higher elevations and poleward (Chen et al., 2011), shifting their recurring life his-19 tory events (phenology) earlier (Wolkovich et al., 2012; Cohen et al., 2018), or both as climate 20 warms (Amano et al., 2014; Socolar et al., 2017). These general trends, however, hide high vari-21 ability across species. A large proportion of species are not shifting (Cook et al., 2012; Amano 22 et al., 2014), which has raised concerns about whether these species may be more vulnerable 23 to population declines with continued warming. Such concerns come in part from increasing 24 research that links how well species track climate change—especially through temporal shifts— 25 to changes in biomass, growth and other metrics related to performance (Cleland et al., 2012). Tracking—a form of phenotypic flexibility—may then be a major component to understanding 27 and predicting the fitness consequences of climate change, including population declines, with 28 cascading effects on community and ecosystem structure (Menzel et al., 2006; Parmesan, 2006). The hypothesis that tracking improves fitness outcomes with climate change has gained significant traction in the ecological literature (e.g., Cleland et al., 2012) and several areas of theory 31 support it. Evolutionary models predict species that track will be favored in novel environmental conditions (Chevin et al., 2010). Niche models of community assembly suggest that a 33 warming climate should open up new temporal niche space and favor species that can exploit 34 that space (Gotelli & Graves, 1996; Wolkovich & Cleland, 2011; Zettlemoyer et al., 2019). How-35 ever, empirical data on the costs (and benefits) of tracking are mixed (e.g., Block et al., 2019). 36 Further, there has been comparatively little work connecting tracking to community assembly theory, which shows temporal sequencing and environmental variability can alter the relative fitness and niche differences between species that determine coexistence, suggesting important ecological constraints to tracking. This disconnect could be because most ecological theory was constructed for stationary systems (e.g., Chesson & Huntly, 1997). While major arenas of research such as 'modern coexistence 42 theory' or population ecology now embrace environmental stochasticity, they generally still assume stationarity, where the underlying distribution of the environment is unchanged across time (i.e., constant mean and variance, Barabas et al., 2018). Climate change upends the assumption of stationarity. By causing increases in temperature, larger pulses of precipitation, shifts in snowpack, increased drought, and more storms (Stocker 47 et al., 2013), climate change has fundamentally shifted major attributes of the environment from stationary to non-stationary regimes (see Fig. 1 and Box: Environmental variability & change). This transition is reshaping ecological systems. New work has aimed to adapt coexistence theory to non-stationary environments (Chesson, 2017; Rudolf, 2019). Yet there is still little 51

Here, we review the concept of tracking as commonly used in the empirical climate change impacts literature and in related ecological theory. We provide definitions that distinguish between fundamental and environmental tracking, highlighting the distinction between measuring

theoretical work on what such a transition may mean for communities and the species within them, including how community-level processes might feedback to modify species responses. tracking and its fitness outcomes in empirical systems. Then, after briefly reviewing evolutionary theory that predicts variation in tracking across species and environments, we examine how well community assembly theory—especially priority effects and modern coexistence theory can be extended to test the current paradigm that climate change should favor species that track.

$_{62}$ 1.1 Defining & measuring tracking

63 Defining tracking

Tracking is a commonly used word in the climate change literature, especially relating to phenology (e.g., Menzel et al., 2006; Parmesan, 2006; Cleland et al., 2012; Deacy et al., 2018). Yet
there are few, if any, definitions of it. Conceptual and theoretical treatments of tracking often
relate how well an organism matches the timing of a life history event to the ideal timing for
that event, what we refer to as 'fundamental tracking'. In contrast, empirical studies of tracking
often focus on estimating a change in the timing of an event relative to a measured environmental variable, with the aim of measuring what we refer to as 'environmental tracking' (Fig.
2)—the change in timing of a major biological event due to an organism's cue system given
change in the environment (though most studies lack the required knowledge of the underlying
cue system, Chmura et al., 2019). Both these definitions are readily applied to phenology—the
timing of recurring life history events—though they can also apply to non-recurring life history
events (e.g., seed germination), or events not normally defined as part of life history.

Fundamental tracking rests on an assumption that there is a timing (an 'ideal timing') that yields maximum fitness, and event timings moving away from this ideal result in reduced fitness. 77 This is a foundational concept of the trophic mis-match literature (Visser & Gienapp, 2019), 78 which often assumes the peak timing of a resource defines the ideal timing for phenological events dependent on that resource (e.g. egg laying dates dependent on caterpillar abundance, Visser & Both, 2005). For most events, however, fitness outcomes are likely dependent on a suite 81 of interacting forces (e.g., Reed et al., 2013)—for example, egg laying dates for migratory birds may depend both on the timing of peak prey abundance and the need to leave nesting grounds before winter. Additionally, the timing of events is often connected to the investment process of an event (e.g., number of eggs in a clutch, Inouve et al., 2019) in current, and sometimes past and future, years. Whatever their full underlying causes, fitness consequences of suboptimal timing should drive the evolution of organismal cues to predict and match the optimal timing 87 (the degree of this match defines cue reliability, Fig. 2). Environmental tracking combines the outcome (timing) of these cues with environmental variation.

Environmental tracking depends on the intersection of the environment's variability—which aspects of the environment vary, how (e.g., temporally each year, spatially at x scale) and how much—and an organism's response to the environment via its proximate cues. If the varying components of the environment are not in the organism's set of cues, then the organism does not 'track' per this definition (although covariation with other environmental components might give the appearance of tracking). Environmental tracking at the individual-level is a purely plastic response to environmental variation (in line with current findings on most climate change

responses, Bonamour et al., 2019), with the plasticity itself an outcome of selection (Chevin et al., 2010) through its connection to fundamental tracking (Fig. 2). At the population-level, tracking may also incorporate evolutionary change—change in genotype frequencies—depending on both the timescales of study and the species' generation time (this evolutionary response can be predicted as the difference between the environmental sensitivity of phenotypic selection and an organism's plasticity, |B - b| in Chevin et al., 2010).

103 Measuring tracking

Measuring 'tracking' and comparing variation in it across species, space and time is a rapidly 104 growing area of ecological research (e.g., Cook et al., 2012; Fu et al., 2015; Thackeray et al., 105 2016; Cohen et al., 2018). Studies that directly quantify fundamental tracking are uncommon 106 (but see Visser et al., 2006; Charmantier et al., 2008), given in part the difficulty of estimating 107 fitness, though many studies in the synchrony literature attempt to link consumer change to 108 resource change, with an assumption that the measured resource is the dominant determinant 109 of ideal timing for the consumer (though this may rarely be true, see Singer & Parmesan, 110 2010: Johansson & Jonzen, 2012: Reed et al., 2013). Instead, most studies focus on estimates 111 closer to environmental tracking. Some studies estimate simply change in days over time (e.g., 112 Parmesan, 2007; Kharouba et al., 2018), though most studies now estimate shifts as responses 113 per unit temperature (for example, multiple meta-analyses show plants' spring phenology shifts 114 with spring or annual temperatures 4-6 days/°C on average across species, Richardson et al., 2006; Wolkovich et al., 2012; Thackeray et al., 2016) or precipitation (Inouye et al., 2002; Craine 116 et al., 2012). 117

All species-rich studies of phenology-climate relationships find high variation (Cook et al., 118 2012; Thackeray et al., 2016), including some species that do not track or track poorly (i.e., 119 high noise surrounding observed statistical relationships). Researchers have worked to link such 120 variation to the underlying cues (e.g., Cook et al., 2012), species traits (e.g., Cohen et al., 2018) 121 and trophic level (e.g., Thackeray et al., 2016). These approaches hint at the three majors 122 explanations for why some species do not appear to track climate or appear to track poorly: 123 (1) environmental tracking is either not possible or optimal for all species or in all environments 124 (Simons, 2011), (2) researchers have measured the wrong environmental variable (i.e., a variable 125 species do not track, Chmura et al., 2019), and (3) statistical artifacts that make it difficult to measure tracking robustly (see Box 'Challenges & opportunities in measuring tracking'). 127

These confounding factors may make many current estimates of interspecific variation in tracking less reliable than they appear. This in turn makes robust quantitative analyses across species difficult (Brown et al., 2016; Kharouba et al., 2018), yielding a muddy picture of which species, when, and where, do and do not track. Given this difficulty, we believe clear testable predictions from ecological theory are especially critical to guide the field forward (Smaldino & McElreath, 2016).

4 1.2 Tracking in single-species environments

Predicting variation in environmental tracking in stationary systems

Evolutionary models predict strong selection for tracking in heterogeneous environments where

there are predictable cues for the ideal timing of events and the underlying genetics to develop 137 a heritable cue system (Piersma & Drent, 2003; Reed et al., 2010). Tracking is likely strongly 138 heritable, given that many cue systems are themselves heritable (e.g., van Asch et al., 2007; 139 Wilczek et al., 2010). The strength of selection is determined by the costs and benefits of 140 cues (Donahue et al., 2015). The costs include the machinery an organism uses to monitor its 141 environment (e.g., accumulated temperature or daylength), while the benefits are the increases 142 in fitness gained from better timing (e.g., how much tissue is saved by avoiding a coldsnap). 143 Selection, however, can be lower than expected from reaction norms predicted by simple evolu-144 tionary models for many reasons, including trade-offs with tracking (Singer & Parmesan, 2010; 145 Johansson & Jonzen, 2012), gene flow from other environments that may continually push a 146 population away from its local optimum (Lenormand, 2002), limits due to standing genetic 147 variation (Franks et al., 2007; Ghalambor et al., 2015), or deeper evolutionary history that may 148 produce co-evolved traits making it difficult for selection to act solely on tracking (Ackerly, 149 2009). 150

Apparently poor cues may occur for organisms in environments where there is both a low cost 151 and low benefit to the cue(s). Whereas expensive cues, such as complex multivariate systems, 152 are possible given a high pay-off. Most in-depth studies of species' phenological cues find 153 evidence for complex multivariate systems that appear adapted to handle unusual—though not 154 completely uncommon—years (Chuine & Regniere, 2017). This suggests that multivariate cues 155 may better couple environmental tracking to fundamental tracking, while simple cues are more 156 likely to trigger growth or reproduction at a suboptimal time. 157

Tracking should generally not be favored in unpredictable environments (e.g. when early season 158 climate cannot be used to predict later season climate), or environments where species otherwise 159 face high uncertainty in the timing of investment decisions. Instead theory suggests the optimal 160 strategy may often be to bet-hedge (Venable, 2007; Donaldson-Matasci et al., 2012; de Casas 161 et al., 2015) via a high diversity of timings or a conservative timing. Because bet-hedging, 162 by definition, maximizes geometric-mean fitness in the long-run, its short-term outcomes can 163 appear maladaptive. How often observed 'maladaptations,' which may easily include species 164 that do not track or appear to track poorly, are actually the outcome of bet-hedging is difficult 165 to estimate, as robustly assessing bet-hedging requires studies of fitness over longer timescales 166 than many current field experiments (Simons, 2011). Environmental variation often includes 167 both predictable and less predictable aspects. In such cases theory predicts organisms may 168 evolve tracking that is a mixed strategy between bet-hedging and plasticity (Wong & Ackerly, 2005). 170

Evolutionary theory, which integrates over the sometimes hidden costs and benefits of particular 171 cue systems and considers environmental predictability, thus provide multiple reasons species 172 may not track or track weakly. This suggests that—at least in stationary systems—we should 173 expect a number of species that do not track. 174

Predicting variation in environmental tracking in non-stationary systems 176

175

177

A major open area of research is adapting life history theory to the non-stationary environments introduced by climate change. Critical to predictions is whether cue systems maintain 178

their reliability with change; i.e., whether they continue to yield high fundamental tracking 179 (Bonamour et al., 2019). Consider a simple case in which an organism's cues evolved based 180 on a correlation between peak prey abundance and daylength: in a stationary environment 181 the daylength cue may be fairly reliable (generally predicting peak prey abundance based on 182 daylength, with some interannual variation), but would become unreliable, and lead to fitness 183 declines, if warming continually advances peak prey abundance. Multivariate cues are often 184 argued to be more reliable because they can capture multiple attributes of the environment 185 (Dore et al., 2018; Bonamour et al., 2019), but they may be equally vulnerable to failure if non-stationarity decouples the cues from fundamental tracking (Bonamour et al., 2019) and 187 thus optimal fitness is no longer associated with the cue system. Predicting the outcome of 188 non-stationarity from the stationary environment requires that researchers know: (1) the full 180 cue system of an organism, (2) how it relates to fundamental tracking, and (3) how both that 190 cue system and the underlying fundamental model shift with non-stationarity. 191

In recent years, plasticity theory has developed to provide insights on non-stationarity (or 192 'sustained environmental change,' see Chevin et al., 2010). Models of the role of plasticity 193 in novel environments provide an important bridge to understanding the outcomes of non-194 stationarity, generally predicting non-stationarity should favor highly plastic species. At the 195 individual level, environmental tracking is a plastic response, and thus we might predict greater 196 individual tracking in non-stationary environments. This outcome, however, assumes there are 197 no costs related to plasticity (Ghalambor et al., 2007; Tufto, 2015). If there are costs associated 198 with tracking, as discussed above in stationary systems, then species may evolve lower tracking 199 (Auld et al., 2010). 200

1.3 Tracking in multi-species environments

201

Life history theory often ignores other (non-focal) species or abstracts them as an aspect of the environment. While the trophic mis-match literature has addressed this gap for trophic interactions (Visser & Both, 2005; Visser & Gienapp, 2019), these is little consideration of competitive coexistence, yet this perspective is critical to understanding environmental tracking (Metcalf et al., 2015). Considering selection in multi-species environments structured by competition highlights that tracking cannot be considered as a singular trait, but must be evaluated as part of a trait syndrome (or mosaic of traits, Ghalambor et al., 2007) and should ultimately produce communities of species where tracking trades-off with other traits.

As tracking often relates to the timing of a resource pulse, traits related to resource acquisition 210 are likely contenders for a trade-off. Species with traits that make them poor resource com-211 petitors may need to track the environment closely to take advantage of transient periods of 212 available resources, but will risk tissue loss to harsh environmental conditions more prevalent 213 early in the season (e.g., frost or snow) and thus may need to be more stress tolerant. In 214 contrast, species with traits that make them superior resource competitors may perform well 215 even if they track environments less closely, because their resource acquisition is not strongly 216 constrained by competitors. Examples include under-canopy species leafing out earlier to gain 217 access to light (Heberling et al., 2019) or species with shallow roots starting growth sooner in an 218

alpine meadow system, while species with deeper roots begin growth later (Zhu et al., 2016). In 219 such cases, tracking is akin to a competition-colonization trade-off (Amarasekare, 2003), where 220 species that track well gain priority access to resources and, thus, may co-exist with superior 221 competitors. Research to date supports this, with several studies linking higher tracking to 222 traits associated with being poor competitors (Dorji et al., 2013; Lasky et al., 2016; Zhu et al., 223 2016). Further, many studies have found a correlation between higher tracking and 'early-224 ness' each season, which has been linked to resource acquisition traits associated with lower 225 competitive abilities (Wolkovich & Cleland, 2014, see Box 'Trait trade-offs with tracking'). 226

Understanding these trade-offs is clearly critical, but the short-term dynamics of a changing 227 environment with plastic species is additionally important and highlights how little ecological 228 theory we have for tracking. While evolutionary theory sometimes predicts the fitness out-229 comes of a new environment, non-stationarity in the climate today means understanding the 230 trajectory may be most relevant—and bridges across evolutionary and ecological timescales. 231 Evolutionary models show how plasticity may limit standing variation and thus reduce fitness 232 in novel environments (Ghalambor et al., 2007; Fournier-Level et al., 2016; Fox et al., 2019). 233 But such findings may not hold if ecological dynamics reshape the environment as systems 234 transition from stationary to non-stationary. 235

236 Including tracking in multi-species community assembly models

246

247

249

250

252

253

254

255

256

257

258

250

Predicting how tracking may determine which species are effectively winners and losers with climate change requires integrating non-stationary environments into models of community 238 assembly. Recent advances in coexistence models, sometimes called 'modern coexistence theory,' 230 recognize that both mechanisms independent of fluctuations in the environment (e.g., R* and 240 other classical niche differences) and mechanisms dependent on fluctuations in the environment 241 (relative non-linearity and storage effect) can lead to coexistence (Chesson & Huntly, 1997; 242 Chesson, 2000). These models, which underlie much of current community ecology research 243 (Mayfield & Levine, 2010; Barabas et al., 2018; Ellner et al., 2019), provide a framework to 244 begin to integrate tracking and non-stationarity into community ecology theory. 245

How the environment is defined in most community models falls into two broad categories. In some models the environment is expressed as variation in parameters related to species. For example, in many lottery models the environment appears, effectively, as variation in birth and death rates; in storage effect models the environment appears as filtered through species low density per capita growth rates (E_i) . Building a changing environment into such models thus requires knowing how environmental shifts filter through to species-level parameters (Tuljapurkar et al., 2009) to impact fundamental tracking. For example, storage effect models predict shifts in communities when environmental change impacts low density per capita growth rates in such a way that the environmental change decreases the covariance between the E_i and competition (i.e., changing $cov(E_i, C_i)$ that leads to a decrease in the storage effect as a means of competitive coexistence). In another example, Rudolf (2019) added the temporal environment to competition models by defining interaction strength as dependent on the temporal distance between species. This is somewhat similar to models that include the environment effectively through different levels of asynchrony (e.g., Nakazawa & Doi, 2012; Revilla et al., 2014).

In other models the environment is more specifically defined as a resource (e.g., many seed

Please fix this WHOLE rest of this para-

Please check this reference to fundamental tracking!

Please check this WHOLE paragraph and the next!

germination models that begin with a resource pulse each year), and thus again generally 261 model something close to fundamental tracking. Models that explicitly include the environment 262 provide a major opportunity to predict how tracking and non-stationarity determine future 263 communities. As an example, we modeled a shift to earlier growing seasons using a common 264 coexistence model where the environment is defined as a limiting resource that determines the 265 start of growth each year. The timing of the resource also determines the offspring species 266 produce each year, allowing us to include fundamental tracking. The shift to earlier seasons 267 favored species that could track, increasing their prevalence by shifting a modelled trade-off 268 between tracking and competitive ability (via R^* , see Fig. 3 and Box: 'Adding tracking and 269 non-stationarity to a common coexistence model'). Like all models, it rests on a number of 270 assumptions, including that species' cues remain as reliable in the non-stationary environment, 271 but shows how non-stationarity could benefit trackers. 272

Most current models (including the previous example) examine the environment from only one 273 of two relevant perspectives: they represent the environment as directly affecting fitness (e.g., 274 the storage effect model), or they represent the environment as used for species' cues (e.g., many 275 models of plasticity). Combining these two perspectives, which connect to fundamental and 276 environmental tracking (respectively), may be especially critical to understanding the costs, 277 benefits and community outcomes of tracking in non-stationary environments. 278

Layered onto these different views of the environment is how species responses are defined. In general, species responses to the environment can be broadly grouped into models that explicitly 280 define when species start an event (e.g., spawning or germination) versus those that model the magnitude of response (e.g., the number of propagules or seeds). Models that explicitly include 282 when a species starts an event are often focused on situations where order of arrival is critical. 283 For example, models of priority effects through niche pre-emption highlight the advantage tracking may provide when it allows species to be early: early arrivals receive a head-start 285 advantage, by gaining priority access to resources (the environment) they can draw down the 286 resources available to later arrivals (Fukami, 2015). Such models predict early-arriving species to out-compete other species, unless there is a cost to being too early or there are trade-offs 288 with other species' traits (see Fig. 4).

281

284

287

280

Other models canalize species' responses to the environment into production and investment. Most models of inter-annual competition (most explicit examples of 'modern coexistence the-291 ory,' e.g., Chesson et al., 2004; Angert et al., 2009) fall into this camp. Species produce (via 292 investment in offspring, tissue etc.) differentially depending on the environment each year and 293 outcomes are mediated through density. While these models superficially may seem discon-294 nected from timing, they highlight how event timing often relates to production and, thus, 295 investment across years. Further, they almost always model the environment as a distribu-296 tion (see Fig. 4), which provides the opportunity for the environment to alter the competitive 297 environment each year and, thus, structure coexistence. 298

A model where species vary both when they start an event and how much they invest would 299 capture the important attributes of fundamental tracking—combining head-start advantages 300 from being early with production variation based on the fitness of the environment. To our 301 knowledge, however, most models approach these questions separately, though models of bethedging come closest (Gourbiere & Menu, 2009; Tufto, 2015). A model that explicitly includes
the linked decisions of when to time an event and how much offspring/tissue to produce during
the event could provide insights on the relative importance of each aspect of this process.
Such a model could be adapted to address multiple questions of tracking, including how these
decisions ('when' and 'how much') may trade-off and which other traits may be most strongly
linked to tracking, as well as explicitly modeling the costs and benefits of tracking in stationary
systems—a critical precursor to extending it to non-stationary systems.

Extending models to non-stationary systems is crucial to testing how environmental tracking relates to fundamental tracking and species persistence with climate change, and research has 311 already begun to tackle this non-trivial challenge (Chesson, 2017; Legault & Melbourne, 2019; 312 Rudolf, 2019). Most work to date, however, focuses on conclusions from systems that are 313 initialized as non-stationary, ignoring the transition between stationary and non-stationary 314 environments. Yet we expect this transition may be critical because communities formed in 315 stationary environments (or periods with lower non-stationarity) are effectively filtered and 316 assembled by that environmental regime and thus produce the baseline of variation and assembly 317 dynamics for a shifting environment. While analytical solutions for systems transitioning from 318 stationary to non-stationary may take time to develop (Chesson, 2017), simulation work can 319 provide an immediate intuition and framework to address this challenge (for an example, see 320 Box: Adding tracking and non-stationarity to a common coexistence model). 321

Outcomes for such community assembly models also depend on how effectively closed commu-322 nities are (i.e., without dispersal or evolution). Dispersal of species or individuals with traits 323 that make them better matched to the non-stationary environment would lead to new com-324 munities that may persist or be continually re-assembled as long as the environment remains 325 non-stationary. Indeed, this logic underlies the argument that invasive species may be superior 326 trackers benefiting from how climate change has altered growing seasons (Willis et al., 2010; 327 Wolkovich & Cleland, 2011). Evolutionary responses could also rescue species with low plastic-328 ity. Long-term population (e.g., Colautti et al., 2017) and resurrection studies (Wilczek et al., 329 2014: Yousev et al., 2018), as well as field experiments (Colautti et al., 2017; Exposito-Alonso 330 et al., 2019), have repeatedly shown species can shift to earlier flowering times, higher thermal 331 tolerances or related genetically-controlled traits that confer higher fitness in warmer climates. Yet these studies also highlight that responses can be lagged (e.g., Wilczek et al., 2014), associ-333 ated with reduced population viability (e.g., Colautti et al., 2017), and that other factors may 334 constrain adaptive responses. 335

336 1.4 Future directions

Growing empirical research highlights that environmental tracking is linked to species performance and, thus, may be critical to understanding the forces that assemble communities and determine species persistence, especially as anthropogenic climate change is reshaping the environment. We have outlined above how current community ecology theory could make advances through models that combine effects of variation in timing and production amounts and models that include environment as impacting species' cues, as well as species' fitness. Such models

would explicitly allow the potential costs and benefits of tracking depending on how closely environmental tracking matches fundamental tracking. But to best test and develop such models we need a greater understanding of how the environment is changing, more robust estimates of environmental tracking and how it fits within a mosaic of other correlated traits that determine individual fitness.

348 How is an organism's environment changing?

Currently, much research has focused on one major shift in the climate system (earlier growing 349 seasons), but research on multivariate environmental shifts is growing and will be critical to 350 understanding how climate change affects an organism's whole environment. We can help guide 351 these efforts by identifying environmental shifts that are often linked. For example, warming 352 temperatures may drive earlier seasons and higher evaporative water loss. Researchers can also 353 aim to more consistently and fully characterize the environmental distributions of their systems 354 that appear to drive species performance and interactions: the environment of the years of 355 study should be clearly reported and compared against long-term and recent climate for each 356 system. 357

More interdisciplinary research with climate science could also speed a fuller understanding of what shifts are and are not expected with climate change, and what climate variables are inherently correlated. Such correlations make estimating cues and other biological parameters from long-term data especially precarious (Tansey et al., 2017). But these correlations are equally critical in considering how species may view their environment and whether environmental change will couple or uncouple links between proximate cues and fundamental tracking (Bonamour et al., 2019).

Understanding and measuring 'tracking'

359

360

362

363

364 365

366

Understanding how the environment is changing represents just one step towards robust mea-367 sures of environmental tracking. Shifting environmental regimes must then be filtered through 368 species cues, and impacts on growth and survival. Studies should clarify their definition of tracking, how the environment is defined, how an event relates to fitness, and how well—or 370 not—the underlying cue system is understood (see Box: 'Challenges & opportunities in mea-371 suring tracking'). Currently, many studies examine fundamental and environmental tracking simultaneously(e.g., Visser et al., 2006; Charmantier et al., 2008; Cleland et al., 2012; Yang & 373 Cenzer, 2020), which is clearly helpful in advancing the field. However, the more researchers can 374 clarify when and how they are addressing fundamental tracking versus environmental tracking, the more easily we can compare results across studies. We expect progress will come from a 376 balance between measures of fundamental tracking (measuring both event date variation and 377 fitness), estimating an organism's system of cues (generally through controlled experiments fol-378 lowed by tests in the field), and measuring the change in an event date relative to environmental 379 variation that is due to cues (environmental tracking). Clear statements of what is known, 380 not known and what is measured will help. 381

What major traits trade-off with tracking?

Basic theory of plasticity and competition suggest that environmental tracking must trade-off with other traits to allow multi-species communities. Yet to date empirical work has mainly

documented tracking, linked it to performance, or focused on how it varies between native and non-native species (Willis et al., 2010; Wolkovich et al., 2013; Zettlemoyer et al., 2019).

Research has highlighted some traits that co-vary with tracking (e.g., Kharouba et al., 2014;

Lasky et al., 2016; Zhu et al., 2016), but to tie this work to models requires more research on traits that link clearly to theory, and a fuller understanding of how tracking and other traits jointly contribute to performance under varying environments.

Traits that link to resource competition, as we focus on here (as others have as well, see Rudolf, 2019), may be especially fruitful for greater research, but should not be the only ones considered. For example, traits related to predator tolerance or avoidance may also play a role, but have been effectively unstudied. As empirical research in this area grows, models can aid progress in understanding the outcomes of these trade-offs for community assembly.

397 1.5 Stationarity in the future

396

While most environments today are climatically non-stationary and have been for decades, the 398 climate will return to a more stationarity form in the future. There are many possible pathways 399 to climatic stabilization, but almost all require first the stabilization of greenhouse gases—the 400 subject of much policy and political debate. Once greenhouse gas emissions stabilize climate 401 will not quickly snap back to a new stationary phase. Instead systems will slowly approach a 402 new climatic stationarity depending on how they are effected by the earth's multiple thermal 403 reservoirs, and, in turn, how quickly those reservoirs stabilize. The timescale of this approach 404 is generally expected to be on the scale of centuries, but could be much longer in certain 405 oceanic systems (Collins et al., 2013). Thus, ecologists are—and will remain for the foreseeable 406 future—in a research area structured by climatic non-stationarity. 407

As paleobiologists and evolutionary biologists often point out, climatic nonstationarity is a 408 common part of the earth's history (Jansson & Dynesius, 2002)—even if stationary periods— 409 be they cold or warm (glacial and interglacial periods), or dry or wet (e.g., megadroughts or 410 pluvials)—are more common. Indeed, while much of this work has examined how species survive 411 for millions of years given large oscillations in climate (Provan & Bennett, 2008), the periods that 412 provide the most dramatic community reshuffling are periods shifting from stationary to non-413 stationary climate regimes (Vrba, 1980, 1985). Such stories of the past are now fundamentally 414 happening today, and ecology is challenged to understand how transitions between stationary 415 and non-stationary environments are reshaping the species and communities we have today and 416 will in the altered climates of our future.

418 2 Acknowledgments

We thank I. Breckheimer, D. Buonaiuto, E. Cleland, J. Davies, G. Legault, A. Phillimore and three anonymous reviewers for helpful comments that improved the manuscript, K. Slimon for help with trade-offs review, and the NSERC Discovery Award (RGPIN-05038 to EMW) and Canada Research Chair in Temporal Ecology (EMW) programs that provided funding.

423 Boxes

424

3.1 Box: Environmental variability & change

Decades of ecological research highlight how temporally variable environments shape species and 425 their communities at multiple scales (Sale, 1977; Chesson & Huntly, 1997). In seasonal land-426 scapes, the environment limits periods for growth each year (e.g., by temperature or drought); 427 within-year variability in the environment (e.g., daily, hourly or finer resolution temperatures 428 or rainfall amounts) compounds into inter-annual variability that shapes the distribution of 429 the start and end of growing seasons. For long stretches of history this variability has been 430 effectively stationary; that is, the underlying probability distribution that describes the start 431 (or end) of the season (e.g., the date of the last major frost) does not change, even though the 432 date may be dramatically different from one year to the next. 433

In other time periods, variability has been non-stationary in one or multiple dimensions. For example, climate in the northern hemisphere includes long warming and then cooling periods (i.e., increasing then decreasing means of the probability distribution) at the start of the Holocene, when the earth was coming out of the last glacial maximum. Anthropogenic climate change is a similar non-stationary process, with warming evident around the globe and knock-on effects for other climate metrics, such as heat extremes and the size of precipitation events.

Understanding non-stationarity in ecological systems requires first identifying which aspects of 440 the environment have shifted—and how they have shifted with respect to one another—as the 441 underlying distributions transition from stationary to non-stationary (Fig. 1). For example, 442 with climate change, warming has increased mean temperatures over time, with minimum tem-443 peratures generally increasing more than maximum—this results in an underlying distribution 444 for daily temperature where the mean is increasing through time while the within-day vari-445 ance is decreasing (Stocker et al., 2013; Screen, 2014). Understanding the impacts of climate change further requires recognizing that most systems can be considered stationary or non-447 stationary depending on the timescale and period of study. Thus, predicting the consequences 448 of current non-stationarity in ecological systems benefits from identifying the type and scale of 449 non-stationarity, relative to long-term trends. 450

451 3.2 Box: Challenges & opportunities in measuring tracking

Increasingly, research has outlined statistical difficulties in measuring tracking, which may underlie many observations of species that do not track. These issues mostly relate to the challenges of observing phenological distributions (Steer *et al.*, 2019; Carter *et al.*, 2018) using discrete (calendar) time, low power, and the complexity of climate data.

Our units of observation may be non-stationary—estimates of mean days shifted per decade depend on the climate of the decade(s) studied, which is not consistent in many systems (McCabe et al., 2012). Estimates based on a relevant climate variable can sometimes ameliorate this problem, but may be equally vulnerable to non-stationarity in units (e.g., Sagarin, 2001).
For example, processes that depend on thermal sums reported as days/°C will generally appear

to decline with warming, as the thermal sum of an average day has increased in most regions with climate change. Relatedly, estimates of long-term change using simple linear regression depend on the climate at the start of the time-series (with greater changes seen from time-series that started in unusually cold decades, such as the 1950s for much of North America).

Even 'long' time-series may be too short for robust analyses of trends (Bolmgren *et al.*, 2013).

Authors should be especially cautious if they find only large effects appear significant (e.g.,

CaraDonna *et al.*, 2014), which is a well-known statistical bias associated with p-values (Loken

& Gelman, 2017). Additionally, effect sizes that are higher when climate variability is higher

(for example, in temperate habitats temperature is highly variable in the spring and autumn

compared to summer) may be more related to variation in statistical power than to biology.

Many statistical issues can be addressed by improved statistical approaches (e.g., Gienapp et al., 471 2005; Pearse et al., 2017), though such approaches may uncomfortably highlight how uncer-472 tain many current estimates are (Brown et al., 2016) or reveal lower effect sizes. Impacts of 473 start-years for long-term time-series can be muted by applying change-point or hinge models 474 (e.g., Kharouba et al., 2018). We suggest mixed models should be used more widely alongside 475 randomization and/or data-simulation approaches (e.g., Bolmgren et al., 2013), and we need 476 models that can discriminate among confounding factors. For example, growing evidence sug-477 gests a potential fundamental trade-off where early species track grow fast and die young, while 478 later species track less, grow slowly and live longer—this might suggest later species bet-hedge 479 more given their longer investment window. Or it could be an artifact where early species use 480 simpler cues, and, thus, their tracking is measured more accurately given current methods. 481

Even without statistical issues, translating phenological and climate data into estimates of 482 tracking requires a firm biological understanding of an organism's cues that we rarely have 483 (Chmura et al., 2019). Currently, 'tracking' is often measured as the relationship between the 484 dates of the phenological event and a simple abiotic metric. Such measures, however, are almost 485 always proxies for a more complicated underlying physiology where simple cues—such as warm 486 temperatures or snowpack—can be modified by other cues, such as photoperiod, drought or light spectra (Bagnall, 1993; Stinchcombe et al., 2004). Teasing out these other cues requires 488 nuanced approaches to observational data with explicit assumptions (Tansey et al., 2017) or 489 controlled experiments (Wilczek et al., 2009; Caffarra et al., 2011). 490

Modeling multivariate cues is inherently difficult (Chuine et al., 2016), especially since one cue 491 may dominate in many conditions. For example, woody plant leafout responds to warm spring 492 temperatures, but also to cool winter temperatures to prevent leafout in mid-winter warm snaps. 493 Often this cool-temperature effect may be masked by sufficiently cool conditions. With warming 494 from climate change, however, this additional trigger may become critical (and potentially lead 495 many phenological models to fail spectacularly in the future, see Chuine et al., 2016). Tracking 496 in species with longer generation times may be especially complicated, as species may track low 497 frequency climate signals and make investment choices on far longer timescales than species 498 with shorter lifespans (Morris et al., 2008). In some semi-arid systems, species time growth to 499 pulses of rain that occur with cooler temperatures, indicating the start of the rainy season, and 500 not a rare summer rainfall event in the middle of months of drought (Wainwright & Cleland, 2013). 502

Addressing these issues is possible if we embrace our inner physiologists, or collaborate with one, to develop models that explicitly include species' cues, and consider the framework under which we expect cue systems have evolved (e.g., bet-hedging). We then must interrogate these models to understand when they work and where they fail (see Johansson & Bolmgren, 2019, for an example). This approach can help embrace the contradictory pulls of conducting experiments to identify mechanistic cues and understanding how they are filtered through the multivariate climate of the real world (see Wilczek et al., 2010, 2009).

3.3 Box: Trait trade-offs with tracking

510

529

530

531

532

533

534

535

536

537

538

539

540

Research on phenological tracking and traits has increased greatly in recent years, with a 511 major uptick in studies after 2010 (see SI Fig. ??). Most papers examining tracking and 512 other traits across species have focused on plants (20/30), followed by birds and Lepidoptera 513 (both 4/30), plankton and aphids (both 1/30). The most studied trait was how early or late a 514 phenophase occurred (e.g., date of flowering of start of migration for a species, termed 'earlyness' 515 by some authors), with earlier species tending to track more (studies included both birds and 516 Lepidotera, Diamond et al., 2011; Ishioka et al., 2013; Kharouba et al., 2014; Jing et al., 2016; 517 Du et al., 2017). While this is an important link, it is vulnerable to statistical challenges (see 518 Box 'Statistical challenges in measuring tracking'). Few studies examined whether tracking 519 correlates with resource acquisition traits; those that did generally found species with higher tracking also had traits associated with lower competitive abilities under low resources (e.g., 521 being shallower or lacking a taproot rooted Dorji et al., 2013; Lasky et al., 2016; Zhu et al., 522 2016). These species were often also early (e.g., Dorji et al., 2013; Zhu et al., 2016), suggesting 523 tracking may relate to a syndrome of traits that allow species to be rapid colonizers each season, 524 but poor competitors for resources. Indeed, previous work has documented that species with 525 earlier phenophases tend to have resource acquisition traits associated with lower competitive abilities (e.g., they tend to be of lower height, have shallower roots, narrower diameter vessels, 527 thinner leaves, and grow faster, reviewed in Wolkovich & Cleland, 2014). 528

3.4 Box: Adding tracking and non-stationarity to a common coexistence model

To understand the role of environmental tracking by species in variable environments we use a simple model that allows within- and between-year dynamics to contribute to coexistence. As the model is akin to many commonly used seed germination models (Chesson $et\ al.$, 2004), we follow a similar terminology for ease; however the basic structure of our model could apply to other systems with one dominant (non-renewing) pulse of a limiting resource each season (e.g., water from rain or snowpack), or a multivariate resource pulse that acts effectively as one resource (e.g., nitrogen and light drawn down together over the season). In this model the environment is included between-years via variable germination, and within-years the environment is explicitly included as a resource pulse at the start of the season. We adjust the biological start time of species (τ_i for species i) to also allow species to respond to the environment dynamically through what we refer to as tracking. Here, tracking effectively moves a

species intrinsic start time closer to the environmental start time in that year, resulting in a higher germination fraction (see SI for complete description and equations).

Species can co-occur via equalizing mechanisms, but they require stabilizing mechanisms to 544 coexist. Species that is, cannot coexist given only variation in tracking—coexistence requires 545 variation in another trait axis. As theory and empirical work suggest this trade-off may involve 546 traits related closely to resource competition, we varied species' R^* . With variation in tracking 547 and in R^* species can persist together as long as those species with a temporal niche advantage 548 are also the inferior competitors (Fig. 3). That is, species that can draw resources down to a 540 lower level and are thus the superior within-season resource competitors (lower R^*) can persist 550 with species with that are inferior competitors but have realized biological start times closer 551 to the environmental start time—a finding inline with currently observed empirical trade-offs (see Box 'Trait trade-offs with tracking'). These trade-offs, however, are all environmentally 553 dependent. They hold only so long as the environment is stationary. 554

We examined how trade-offs may be transformed by a non-stationary environment, by transitioning a stationary environment—in which two-species communities had persisted for 500 years—to non-stationary, via an earlier start of season (earlier timing of the resource pulse, τ_p , Fig. 3; see SI for more details). By changing a fundamental niche axis (the distribution of the environment, an axis along which these communities were structured), we shifted one major part of the trade-off: the new non-stationary environment favored an earlier start time than the previous stationary environment. This, in turn, reshaped our two-species communities, which depended on this trade-off for persistence.

While the non-stationary environment favored higher trackers (who in turn drove the extinction of species with lower tracking values from many two-species communities) some two-species communities persisted (257 out of 1698 two-species communities persisting after end of stationary, or 15.1%, Fig. 3). These two-species communities persisted because the same fundamental trade-off between biological start time and within-season competitive ability, while narrowed, was not fully lost. Taken together, these simple simulations show how non-stationarity can drive local species extinction and reshape the underlying assembly mechanisms of communities.

Our simulations support growing work that tracking should not be considered alone (Diamond 570 et al., 2011; Dorji et al., 2013; Ishioka et al., 2013; Kharouba et al., 2014; Du et al., 2017), but 571 may be part of a larger trait syndrome. Indeed, this model trivially shows that multi-species 572 communities cannot form given only variation in the temporal niche—a trade-off is required. 573 Our results thus support empirical work showing a trade-off where trackers are also inferior 574 resource competitors (Lasky et al., 2016; Zhu et al., 2016)—we show this must be the case for 575 multi-species persistence; otherwise, the species best matched to the environment would drive 576 the other extinct. 577

Finally, our results highlight that non-stationarity may reshape the balance of equalizing versus stabilizing mechanisms. As environments shifted from stationarity to non-stationarity, species that co-occurred via equalizing mechanisms persisted longer. While the outcome that equalized species will be more similarly affected by environmental shifts is rather obvious, it has several important implications. First, it may make identifying which traits climate change promotes through stabilizing mechanisms more difficult. Second, it suggests climate change—or other factors that cause an environment to shift from stationary to non-stationary—may cause a

 $_{585}$ fundamental shift away from assembly via stabilizing mechanisms.

586 4 Figures

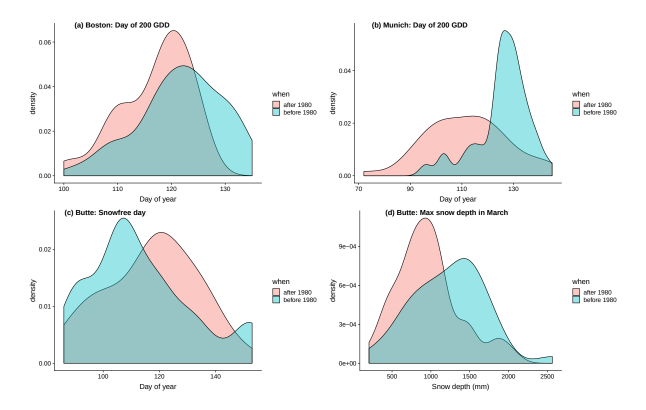


Figure 1: Examples of non-stationarity in climate variables linked to environmental tracking: shifts before and after 1980 (a major change-point in climate for many regions) in several metrics related to the start of growing seasons (a-c) or resource pulse connected to growing season length (d). Density plots of day of 200 growing degree day units (a metric of thermal sum, here based on 0 degree base temperature using daily minima in °C) in Boston, MA, USA (a), and Munich, Germany (b), first snowfree day (followed by at least 9 snowfree days) in Crested Butte, CO, USA (c) and maximum snowdepth (mm) in March (often the month before the first snowfree day) in Crested Butte, CO, USA (d). Note that (c) and (d) are likely related, with lower snowpacks leading to an earlier first snowfree day. We selected sites that have been studied for plant phenological data and included at least 80 years of daily climate data from a Global Historical Climatology Network site; we subsetted data so that there were 40 years before and after 1980 for all sites.

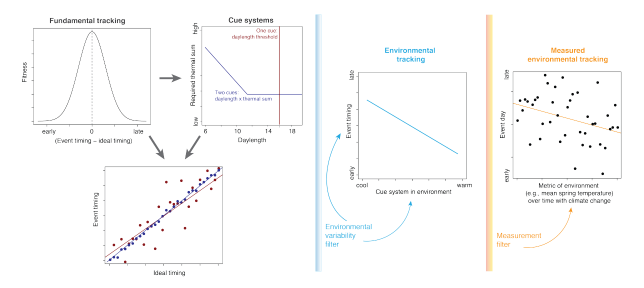


Figure 2: Defining tracking: Fundamental tracking (top left) represents how an organism matches the actual timing of a life history event to the timing that maximizes its fitness (i.e., ideal timing). Here, we show one conceptualization, where fitness declines as event timing moves away from the ideal timing, though realizations in nature may take diverse forms. The cue system is the physiological machinery that an organism uses to predict the ideal timing; we show one dependent only on daylength (red line: the event occurs when the organism's environment exceeds a certain daylength) and one that depends on a combination of daylength and thermal sums (navy line: the event occurs when the organism accumulates enough temperature for the current daylength). The skill of a cue system in predicting the ideal timing represents cue quality (bottom left). The combination of environmental variability and an organism's cue system results in different event timings depending on the environment, what we define as environmental tracking (middle). Measured environmental tracking (right) depends both on environmental tracking and the metric(s) of the environment that researchers measure. MAKE SURE WE GET ACROSS: Why we have such a picky definition of environmental tracking (versus measured environmental tracking)—because covariation between timing and the measured environment is not so useful to model and could lead to wildly incorrect predictions!



Figure 3: Example of how non-stationarity can reshape communities in a simple model. We shifted the environment (top panels) by changing the timing of the resource pulse from a stationary period ($\tau_p \sim \beta(10,10)$ for the 500 years) to a nonstationary period ($\tau_p \sim \beta(5,15)$ over the 500 years), then examined outcomes for two-species communities (bottom panels) where tracking (X axis: species 1/species 2) trades off with R^* (Y axis: species 1/species 2): each point represents one two-species community color-coded by whether both species persisted or one or more species was extirpated through 500 years of a stationary environment (bottom-left), followed by an additional 500 years of non-stationary environment (bottom-right, only two-species communities that persisted through the stationary period are shown).

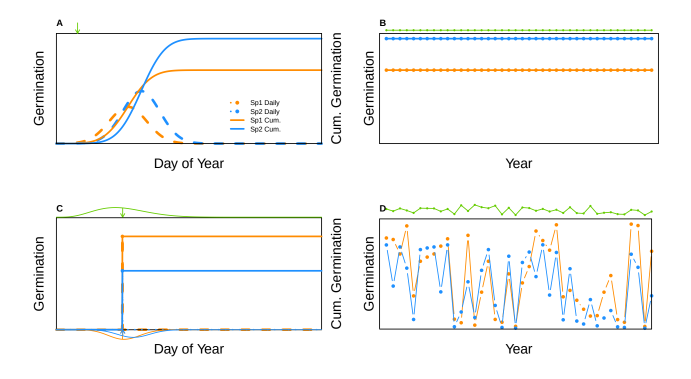


Figure 4: Tracking can be conceptualized as changes in priority effects or changes in storage effects. In a priority effect model (A,B), the coexistence mechanism is a within-year tradeoff between, an early-germinating species that pre-empts resources (sp 1) and late-germinating species that is a superior resource competitor (sp 2) (A, where green arrow indicates the start of season); no between-year variation is required to maintain coexistence. In a storage effect model (C,D), variation in the timing of the start of season (indicated by the distribution in green, top of C) results results in differential species-response to the environment (illustrated by species-specific germination curves, bottom of C); this interannual variation in species-response to the environment (D)—along with a seedbank or other interannual storage mechanism—can maintain coexistence through reduced interspecific competition.

References

- Ackerly, D. (2009). Conservatism and diversification of plant functional traits: Evolutionary rates versus phylogenetic signal. *Proceedings of the National Academy of Sciences of the* United States of America, 106, 19699–19706.
- Amano, T., Freckleton, R.P., Queenborough, S.A., Doxford, S.W., Smithers, R.J., Sparks, T.H. & Sutherland, W.J. (2014). Links between plant species' spatial and temporal responses to a warming climate. *Proceedings of the Royal Society B-Biological Sciences*, 281.
- Amarasekare, P. (2003). Competitive coexistence in spatially structured environments: a synthesis. *Ecology Letters*, 6.
- Angert, A.L., Huxman, T.E., Chesson, P. & Venable, D.L. (2009). Functional tradeoffs determine species coexistence via the storage effect. *Proceedings of the National Academy of Sciences of the United States of America*, 106, 11641–11645.
- van Asch, M., Tienderen, P.H., Holleman, L.J.M. & Visser, M.E. (2007). Predicting adaptation of phenology in response to climate change, an insect herbivore example. *Global Change Biology*, 13, 1596–1604.
- Auld, J.R., Agrawal, A.A. & Relyea, R.A. (2010). Re-evaluating the costs and limits of adaptive phenotypic plasticity. *Proceedings of the Royal Society B-Biological Sciences*, 277, 503–511.
- Bagnall, D.J. (1993). Light quality and vernalization interact in controlling late flowering
 Arabidopsis ecotypes and mutants. Annals of Botany, 71, 75–83.
- Barabas, G., D'Andrea, R. & Stump, S.M. (2018). Chesson's coexistence theory. Ecological
 Monographs, 88, 277–303.
- Block, S., Alexander, J. & Levine, J.M. (2019). Phenological plasticity is a poor predictor of subalpine plant population performance following experimental climate change. *Oikos*, p. 10.1111/oik.06667.
- Bolmgren, K., Vanhoenacker, D. & Miller-Rushing, A.J. (2013). One man, 73 years, and 25 species. evaluating phenological responses using a lifelong study of first flowering dates. *International Journal of Biometeorology*, 57, 367–375.
- Bonamour, S., Chevin, L.M., Charmantier, A. & Teplitsky, C. (2019). Phenotypic plasticity in response to climate change: the importance of cue variation. *Philosophical Transactions of the Royal Society B-Biological Sciences*, 374.
- Brown, C.J., O'Connor, M.I., Poloczanska, E.S., Schoeman, D.S., Buckley, L.B., Burrows,
 M.T., Duarte, C.M., Halpern, B.S., Pandolfi, J.M., Parmesan, C. & Richardson, A.J. (2016).
 Ecological and methodological drivers of species' distribution and phenology responses to
- climate change. Global Change Biology, 22, 1548–1560.
- Caffarra, A., Donnelly, A. & Chuine, I. (2011). Modelling the timing of Betula pubescens
 budburst. II. Integrating complex effects of photoperiod into process-based models. Climate
 Research, 46, 159–170.

- CaraDonna, P.J., Iler, A.M. & Inouye, D.W. (2014). Shifts in flowering phenology reshape a
 subalpine plant community. Proceedings of the National Academy of Sciences, 111, 4916–4921.
- Carter, S.K., Saenz, D. & Rudolf, V.H.W. (2018). Shifts in phenological distributions reshape
 interaction potential in natural communities. *Ecology Letters*, 21, 1143–1151.
- de Casas, R.R., Donohue, K., Venable, D.L. & Cheptou, P.O. (2015). Gene-flow through space and time: dispersal, dormancy and adaptation to changing environments. *Evolutionary Ecology*, 29, 813–831.
- Charmantier, A., McCleery, R.H., Cole, L.R., Perrins, C., Kruuk, L.E.B. & Sheldon, B.C.
 (2008). Adaptive phenotypic plasticity in response to climate change in a wild bird population.
 Science, 320, 800–803.
- Chen, I.C., Hill, J.K., Ohlemuller, R., Roy, D.B. & Thomas, C.D. (2011). Rapid range shifts
 of species associated with high levels of climate warming. Science, 333, 1024–1026.
- Chesson, P. (2000). Mechanisms of maintenance of species diversity. Annual Review of Ecology
 and Systematics, 31, 343–366.
- Chesson, P. (2017). Aedt: A new concept for ecological dynamics in the ever-changing world.
 Plos Biology, 15.
- Chesson, P., Gebauer, R.L.E., Schwinning, S., Huntly, N., Wiegand, K., Ernest, M.S.K., Sher,
 A., Novoplansky, A. & Weltzin, J.F. (2004). Resource pulses, species interactions, and diversity maintenance in arid and semi-arid environments. *Oecologia*, 141, 236–253.
- Chesson, P. & Huntly, N. (1997). The roles of harsh and fluctuating conditions in the dynamics
 of ecological communities. American Naturalist, 150, 519–553.
- Chevin, L.M., Lande, R. & Mace, G.M. (2010). Adaptation, plasticity, and extinction in a
 changing environment: Towards a predictive theory. *Plos Biology*, 8.
- Chmura, H.E., Kharouba, H.M., Ashander, J., Ehlman, S.M., Rivest, E.B. & Yang, L.H. (2019).
 The mechanisms of phenology: the patterns and processes of phenological shifts. *Ecological Monographs*, 89.
- Chuine, I., Bonhomme, M., Legave, J.M., de Cortazar-Atauri, I.G., Charrier, G., Lacointe, A. &
 Ameglio, T. (2016). Can phenological models predict tree phenology accurately in the future?
 the unrevealed hurdle of endodormancy break. Global Change Biology, 22, 3444–3460.
- Chuine, I. & Regniere, J. (2017). Process-based models of phenology for plants and animals.
 Annual Review of Ecology, Evolution, and Systematics, 48, 159–182.
- 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.
- Cohen, J.M., Lajeunesse, M.J. & Rohr, J.R. (2018). A global synthesis of animal phenological
 responses to climate change. Nature Climate Change, 8, 224-+.

- Colautti, R.I., Aring;gren, J. & Anderson, J.T. (2017). Phenological shifts of native and invasive
 species under climate change: insights from the boechera lythrum model. *Philosophical Transactions of the Royal Society B-Biological Sciences*, 372.
- Collins, M., Knutti, R., Arblaster, J.L., Dufresne, T., Fichefet, P., Friedlingstein, X., Gao,
 W., Gutowski, T., Johns, G., Krinner, M., Shongwe, C., Tebaldi, A., Weaver & Wehner, M.
 (2013). Climate Change 2013: The Physical Science Basis Working Group I Contribution
 to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, Cam-
- bridge University Press, chap. Long-term Climate Change: Projections, Commitments and Irreversibility, pp. 1029–1119.
- Cook, B.I., Wolkovich, E.M. & Parmesan, C. (2012). Divergent responses to spring and winter
 warming drive community level flowering trends. Proceedings of the National Academy of
 Sciences of the United States of America, 109, 9000–9005.
- Craine, J.M., Wolkovich, E.M., Towne, E.G. & Kembel, S.W. (2012). Flowering phenology as
 a functional trait in a tallgrass prairie. New Phytologist, 193, 673–682.
- Deacy, W.W., Erlenbach, J.A., Leacock, W.B., Stanford, J.A., Robbins, C.T. & Armstrong,
 J.B. (2018). Phenological tracking associated with increased salmon consumption by brown
 bears. Scientific Reports, 8.
- Diamond, S.E., Frame, A.M., Martin, R.A. & Buckley, L.B. (2011). Species' traits predict phenological responses to climate change in butterflies. *Ecology*, 92, 1005–1012.
- Donahue, M.J., Karnauskas, M., Toews, C. & Paris, C.B. (2015). Location isn't everything:
 Timing of spawning aggregations optimizes larval replenishment. *Plos One*, 10.
- Donaldson-Matasci, M.C., Bergstrom, C.T. & Lachmann, M. (2012). When unreliable cues are good enough. *American Naturalist*, 182, 313–327.
- Dore, A.A., McDowall, L., Rouse, J., Bretman, A., Gage, M.J.G. & Chapman, T. (2018). The
 role of complex cues in social and reproductive plasticity. Behavioral Ecology and Sociobiology,
 72.
- Dorji, T., Totland, O., Moe, S.R., Hopping, K.A., Pan, J. & Klein, J.A. (2013). Plant functional
 traits mediate reproductive phenology and success in response to experimental warming and
 snow addition in Tibet. Global Change Biology, 19, 459–472.
- Du, Y.J., Chen, J.R., Willis, C.G., Zhou, Z.Q., Liu, T., Dai, W.J., Zhao, Y. & Ma, K.P. (2017).
 Phylogenetic conservatism and trait correlates of spring phenological responses to climate
 change in northeast china. *Ecology and Evolution*, 7, 6747–6757.
- Ellner, S.P., Snyder, R.E., Adler, P.B. & Hooker, G. (2019). An expanded modern coexistence theory for empirical applications. *Ecology Letters*, 22, 3–18.
- Exposito-Alonso, M., Burbano, H.A., Bossdorf, O., Nielsen, R., Weigel, D., Exposito-Alonso,
 M., Rodriguez, R.G., Barragan, C., Capovilla, G., Chae, E., Devos, J., Dogan, E.S., Friedemann, C., Gross, C., Lang, P., Lundberg, D., Middendorf, V., Kageyama, J., Karasov,

- T., Kersten, S., Petersen, S., Rabbani, L., Regalado, J., Reinelt, L., Rowan, B., Seymour,
- D.K., Symeonidi, E., Schwab, R., Tran, D.T.N., Venkataramani, K., Van de Weyer, A.L.,
- Vasseur, F., Wang, G., Wedegartner, R., Weiss, F., Wu, R., Xi, W.Y., Zaidem, M., Zhu,
- W.S., Garcia-Arenal, F., Burbano, H.A., Bossdorf, O., Weigel, D. & Genomes Field Expt, T.
- 701 (2019). Natural selection on the arabidopsis thaliana genome in present and future climates.
- 702 Nature, 573, 126-+.
- Fournier-Level, A., Perry, E.O., Wang, J.A., Braun, P.T., Migneault, A., Cooper, M.D., Met-
- calf, C.J.E. & Schmitt, J. (2016). Predicting the evolutionary dynamics of seasonal adaptation
- to novel climates in arabidopsis thaliana. Proceedings of the National Academy of Sciences
- of the United States of America, 113, E2812–E2821.
- Fox, R.J., Donelson, J.M., Schunter, C., Ravasi, T. & Gaitan-Espitia, J.D. (2019). Beyond
- buying time: the role of plasticity in phenotypic adaptation to rapid environmental change.
- 709 Philosophical Transactions of the Royal Society B-Biological Sciences, 374.
- Franks, S.J., Sim, S. & Weis, A.E. (2007). Rapid evolution of flowering time by an annual plant
- in response to a climate fluctuation. Proceedings of the National Academy of Sciences of the
- United States of America, 104, 1278–1282.
- Fu, Y.S.H., Zhao, H.F., Piao, S.L., Peaucelle, M., Peng, S.S., Zhou, G.Y., Ciais, P., Huang,
- M.T., Menzel, A., Uelas, J.P., Song, Y., Vitasse, Y., Zeng, Z.Z. & Janssens, I.A. (2015).
- Declining global warming effects on the phenology of spring leaf unfolding. Nature, 526,
- 716 104–107.
- Fukami, T. (2015). Historical Contingency in Community Assembly: Integrating Niches, Species
- Pools, and Priority Effects, vol. 46 of Annual Review of Ecology Evolution and Systematics,
- 719 pp. 1–23.
- Ghalambor, C.K., Hoke, K.L., Ruell, E.W., Fischer, E.K., Reznick, D.N. & Hughes, K.A.
- (2015). Non-adaptive plasticity potentiates rapid adaptive evolution of gene expression in
- nature. Nature, 525, 372-+.
- Ghalambor, C.K., McKay, J.K., Carroll, S.P. & Reznick, D.N. (2007). Adaptive versus non-
- adaptive phenotypic plasticity and the potential for contemporary adaptation in new envi-
- ronments. Functional Ecology, 21, 394–407.
- Gienapp, P., Hemerik, L. & Visser, M.E. (2005). A new statistical tool to predict phenology
- under climate change scenarios. Global Change Biology, 11, 600–606.
- Gotelli, N.J. & Graves, G.R. (1996). The temporal niche. In: Null Models in Ecology (eds.
- Gotelli, N.J. & Graves, G.R.). Smithsonian Institution, Washington, DC, pp. 95–111.
- Gourbiere, S. & Menu, F. (2009). Adaptive dynamics of dormancy duration variability: evolu-
- tionary trade-off and priority effect lead to suboptimal adaptation. Evolution, 63, 1879–1892.
- Heberling, J.M., Cassidy, S.T., Fridley, J.D. & Kalisz, S. (2019). Carbon gain phenologies
- of spring-flowering perennials in a deciduous forest indicate a novel niche for a widespread
- invader. New Phytologist, 221, 778–788.

- Hoegh-Guldberg, O., Jacob, D., Taylor, M., Bindi, M., Brown, S., Camilloni, I., Diedhiou,
- A., Djalante, R., Ebi, K., Engelbrecht, F., Guiot, J., Hijioka, Y., Mehrotra, S., Payne, A.,
- Seneviratne, S., Thomas, A., R. Warren, R. & Zhou, G. (2018). Impacts of 1.5C Global
- Warming on Natural and Human Systems.
- Inouye, B.D., Ehrlen, J. & Underwood, N. (2019). Phenology as a process rather than an event:
- from individual reaction norms to community metrics. Ecological Monographs, 89.
- Inouye, D.W., Morales, M.A. & Dodge, G.J. (2002). Variation in timing and abundance of
- flowering by *Delphinium barbeyi* Huth (Ranunculaceae): the roles of snowpack, frost, and La
- Niña, in the context of climate change. Oecologia, 130, 543–550.
- ⁷⁴⁴ IPCC (2014). Climate Change 2014: Impacts, Adaptation, and Vulnerability. Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA.
- Ishioka, R., Muller, O., Hiura, T. & Kudo, G. (2013). Responses of leafing phenology and
- photosynthesis to soil warming in forest-floor plants. Acta Oecologica-International Journal
- of Ecology, 51, 34–41.
- Jansson, R. & Dynesius, M. (2002). The fate of clades in a world of recurrent climatic change:
- Milankovitch oscillations and evolution. Annual Review of Ecology and Systematics, 33, 741–
- 751 777.
- Jing, J., Li, K. & Liu, Z.G. (2016). Effects of varying temperature on leaf phenology and
- herbivory of dominant tree species in subtropical evergreen broad-leaves forest in eastern
- china. Polish Journal of Ecology, 64, 53–69.
- Johansson, J. & Bolmgren, K. (2019). Is timing of reproduction according to temperature sums
- an optimal strategy? Ecology and Evolution, 9, 11598–11605.
- Johansson, J. & Jonzen, N. (2012). Game theory sheds new light on ecological responses to
- current climate change when phenology is historically mismatched. Ecol Lett, 15, 881–8.
- Kharouba, H.M., Ehrlen, J., Gelman, A., Bolmgren, K., Allen, J.M., Travers, S.E. & Wolkovich,
- E.M. (2018). Global shifts in the phenological synchrony of species interactions over recent
- decades. Proceedings of the National Academy of Sciences of the United States of America,
- 762 115, 5211–5216.
- Kharouba, H.M., Paquette, S.R., Kerr, J.T. & Vellend, M. (2014). Predicting the sensitivity
- of butterfly phenology to temperature over the past century. Global Change Biology, 20,
- 765 504–514.
- Lasky, J.R., Uriarte, M. & Muscarella, R. (2016). Synchrony, compensatory dynamics, and the
- functional trait basis of phenological diversity in a tropical dry forest tree community: effects
- of rainfall seasonality. Environmental Research Letters, 11.
- Legault, G. & Melbourne, B.A. (2019). Accounting for environmental change in continuous-time
- stochastic population models. Theoretical Ecology, 12, 31–48.

- Lenormand, T. (2002). Gene flow and the limits to natural selection. Trends in Ecology & Evolution, 17, 183–189.
- Loken, E. & Gelman, A. (2017). Measurement error and the replication crisis. *Science*, 355, 584–585.
- Mayfield, M.M. & Levine, J.M. (2010). Opposing effects of competitive exclusion on the phylogenetic structure of communities. *Ecology Letters*, 13, 1085–1093.
- McCabe, G.J., Ault, T.R., Cook, B.I., Betancourt, J.L. & Schwartz, M.D. (2012). Influences of the el nino southern oscillation and the pacific decadal oscillation on the timing of the north american spring. *International Journal of Climatology*, 32, 2301–2310.
- Menzel, A., Von Vopelius, J., Estrella, N., Schleip, C. & Dose, V. (2006). Farmers' annual activities are not tracking the speed of climate change. *Climate Research*, 32, 201–207.
- Metcalf, C.J.E., Burghardt, L.T. & Koons, D.N. (2015). Avoiding the crowds: the evolution of plastic responses to seasonal cues in a density-dependent world. *Journal of Ecology*, 103, 819–828.
- Morris, W.F., Pfister, C.A., Tuljapurkar, S., Haridas, C.V., Boggs, C.L., Boyce, M.S., Bruna, E.M., Church, D.R., Coulson, T., Doak, D.F., Forsyth, S., Gaillard, J.M., Horvitz, C.C., Kalisz, S., Kendall, B.E., Knight, T.M., Lee, C.T. & Menges, E.S. (2008). Longevity can buffer plant and animal populations against changing climatic variability. *Ecology*, 89, 19–25.
- Nakazawa, T. & Doi, H. (2012). A perspective on match/mismatch of phenology in community contexts. *Oikos*, 121, 489–495.
- Parmesan, C. (2006). Ecological and evolutionary responses to recent climate change, vol. 37 of Annual Review of Ecology Evolution and Systematics, pp. 637–669.
- Parmesan, C. (2007). Influences of species, latitudes and methodologies on estimates of phenological response to global warming. *Global Change Biology*, 13, 1860–1872.
- Pearse, W.D., Davis, C.C., Inouye, D.W., Primack, R.B. & Davies, T.J. (2017). A statistical
 estimator for determining the limits of contemporary and historic phenology. Nature Ecology
 Evolution, 1, 1876-+.
- Piersma, T. & Drent, J. (2003). Phenotypic flexibility and the evolution of organismal design.
 Trends in Ecology & Evolution, 18, 228–233.
- Provan, J. & Bennett, K.D. (2008). Phylogeographic insights into cryptic glacial refugia. Trends
 in Ecology & Evolution, 23, 564-571.
- Reed, T.E., Grotan, V., Jenouvrier, S., Saether, B.E. & Visser, M.E. (2013). Population growth in a wild bird is buffered against phenological mismatch. *Science*, 340, 488–491.
- Reed, T.E., Waples, R.S., Schindler, D.E., Hard, J.J. & Kinnison, M.T. (2010). Phenotypic plasticity and population viability: the importance of environmental predictability. *Proceedings of the Royal Society B-Biological Sciences*, 277, 3391–3400.

- Revilla, T.A., Encinas-Viso, F. & Loreau, M. (2014). (a bit) earlier or later is always better:
 Phenological shifts in consumer-resource interactions. *Theoretical Ecology*, 7, 149–162.
- Richardson, A.D., Bailey, A.S., Denny, E.G., Martin, C.W. & O'Keefe, J. (2006). Phenology of a northern hardwood forest canopy. *Global Change Biology*, 12, 1174–1188.
- Rudolf, V.H.W. (2019). The role of seasonal timing and phenological shifts for species coexistence. *Ecology Letters*.
- Sagarin, R. (2001). Phenology false estimates of the advance of spring. Nature, 414, 600–600.
- Sale, P.F. (1977). Maintenance of high diversity in coral reef fish communities. *American*Naturalist, 111, 337–359.
- Screen, J.A. (2014). Arctic amplification decreases temperature variance in northern mid- to high-latitudes. *Nature Climate Change*, 4, 577–582.
- Simons, A.M. (2011). Modes of response to environmental change and the elusive empirical evidence for bet hedging. *Proceedings of the Royal Society B-Biological Sciences*, 278, 1601–1609.
- Singer, M.C. & Parmesan, C. (2010). Phenological asynchrony between herbivorous insects and their hosts: signal of climate change or pre-existing adaptive strategy? *Philosophical Transactions of the Royal Society B: Biological Sciences*, 365, 3161–3176.
- Smaldino, P.E. & McElreath, R. (2016). The natural selection of bad science. Royal Society
 Open Science, 3.
- Socolar, J.B., Epanchin, P.N., Beissinger, S.R. & Tingley, M.W. (2017). Phenological shifts conserve thermal niches in north american birds and reshape expectations for climate-driven range shifts. *Proceedings of the National Academy of Sciences of the United States of America*, 114, 12976–12981.
- Steer, N.C., Ramsay, P.M. & Franco, M. (2019). nlstimedist: An r package for the biologically meaningful quantification of unimodal phenology distributions. *Methods in Ecology and Evolution*, 10, 1934–1940.
- Stinchcombe, J.R., Weinig, C., Ungerer, M., Olsen, K.M., Mays, C., Halldorsdottir, S.S., Purugganan, M.D. & Schmitt, J. (2004). A latitudinal cline in flowering time in *Arabidopsis thaliana* modulated by the flowering time gene FRIGIDA. *Proceedings of the National Academy of Sciences of the United States of America*, 101, 4712–4717.
- Stocker, T., Qin, D. & Platner, G. (2013). Climate change 2013: The physical science basis.

 Working Group I Contribution to the Fifth Assessment Report of the Intergovernmental Panel
 on Climate Change. Summary for Policymakers (IPCC, 2013).
- Tansey, C.J., Hadfield, J.D. & Phillimore, A.B. (2017). Estimating the ability of plants to plastically track temperature-mediated shifts in the spring phenological optimum. *Global Change Biology*, 23, 3321–3334.

- Thackeray, S.J., Henrys, P.A., Hemming, D., Bell, J.R., Botham, M.S., Burthe, S., Helaouet,
- P., Johns, D.G., Jones, I.D., Leech, D.I., Mackay, E.B., Massimino, D., Atkinson, S., Bacon,
- P.J., Brereton, T.M., Carvalho, L., Clutton-Brock, T.H., Duck, C., Edwards, M., Elliott,
- J.M., Hall, S.J.G., Harrington, R., Pearce-Higgins, J.W., Høye, T.T., Kruuk, L.E.B., Pem-
- berton, J.M., Sparks, T.H., Thompson, P.M., White, I., Winfield, I.J. & Wanless, S. (2016).
- Phenological sensitivity to climate across taxa and trophic levels. *Nature*, 535, 241.
- Tufto, J. (2015). Genetic evolution, plasticity, and bet-hedging as adaptive responses to tem-
- porally autocorrelated fluctuating selection: A quantitative genetic model. *Evolution*, 69,
- 851 2034-2049.
- Tuljapurkar, S., Gaillard, J.M. & Coulson, T. (2009). From stochastic environments to life
- histories and back. Philosophical Transactions of the Royal Society B-Biological Sciences,
- 364, 1499–1509.
- Venable, D.L. (2007). Bet hedging in a guild of desert annuals. Ecology, 88, 1086–1090.
- Visser, M.E. & Both, C. (2005). Shifts in phenology due to global climate change: the need for
- a yardstick. Proceedings of the Royal Society B-Biological Sciences, 272, 2561–2569.
- Visser, M.E. & Gienapp, P. (2019). Evolutionary and demographic consequences of phenological
- mismatches. Nature Ecology & Evolution, 3, 879–885.
- Visser, M.E., Holleman, L.J.M. & Gienapp, P. (2006). Shifts in caterpillar biomass phenol-
- ogy due to climate change and its impact on the breeding biology of an insectivorous bird.
- 862 Oecologia, 147, 164–172.
- Vrba, E.S. (1980). Evolution, species and fossils how does life evolve. South African Journal
- of Science, 76, 61–84.
- Vrba, E.S. (1985). Environment and evolution alternative causes of the temporal distribution
- of evolutionary events. South African Journal of Science, 81, 229–236.
- Wainwright, C.E. & Cleland, E.E. (2013). Exotic species display greater germination plasticity
- and higher germination rates than native species across multiple cues. Biological Invasions,
- 869 15, 2253–2264.
- Wilczek, A.M., Burghardt, L.T., Cobb, A.R., Cooper, M.D., Welch, S.M. & Schmitt, J. (2010).
- Genetic and physiological bases for phenological responses to current and predicted climates.
- Philosophical Transactions of the Royal Society B: Biological Sciences, 365, 3129–3147.
- Wilczek, A.M., Cooper, M.D., Korves, T.M. & Schmitt, J. (2014). Lagging adaptation to
- warming climate in arabidopsis thaliana. Proceedings of the National Academy of Sciences
- of the United States of America, 111, 7906–7913.
- Wilczek, A.M., Roe, J.L., Knapp, M.C., Cooper, M.D., Lopez-Gallego, C., Martin, L.J., Muir,
- 877 C.D., Sim, S., Walker, A., Anderson, J., Egan, J.F., Moyers, B.T., Petipas, R., Giakountis, A.,
- Charbit, E., Coupland, G., Welch, S.M. & Schmitt, J. (2009). Effects of genetic perturbation
- on seasonal life history plasticity. Science, 323, 930–934.

- Willis, C.G., Ruhfel, B.R., Primack, R.B., Miller-Rushing, A.J., Losos, J.B. & Davis, C.C.
 (2010). Favorable climate change response explains non-native species' success in Thoreau's
 woods. *PLoS ONE*, 5, e8878.
- 882 WOOds. 1 Los ONE, 5, 60010.
- Wolkovich, E.M. & Cleland, E.E. (2011). The phenology of plant invasions: A community ecology perspective. Frontiers in Ecology and the Environment, 9, 287–294.
- Wolkovich, E.M. & Cleland, E.E. (2014). Phenological niches and the future of invaded ecosystems with climate change. *AoB Plants*, 6, plu013.
- Wolkovich, E.M., Cook, B.I., Allen, J.M., Crimmins, T.M., Betancourt, J.L., Travers, S.E.,
- Pau, S., Regetz, J., Davies, T.J., Kraft, N.J.B., Ault, T.R., Bolmgren, K., Mazer, S.J.,
- McCabe, G.J., McGill, B.J., Parmesan, C., Salamin, N., Schwartz, M.D. & Cleland, E.E.
- 890 (2012). Warming experiments underpredict plant phenological responses to climate change.
- Nature, 485, 494–497.
- Wolkovich, E.M., Davies, T.J., Schaefer, H., Cleland, E.E., Cook, B.I., Travers, S.E., Willis,
- 893 C.G. & Davis, C. (2013). Temperature-dependent shifts in phenology contribute to the success
- of exotic species with climate change. American Journal of Botany, 100, 1407–1421.
- Wong, T.G. & Ackerly, D.D. (2005). Optimal reproductive allocation in annuals and an informational constraint on plasticity. *New Phytologist*, 166, 159–172.
- Yang, L.H. & Cenzer, M.L. (2020). Seasonal windows of opportunity in milkweed-monarch interactions. *Ecology*, 101.
- Yousey, A.M., Chowdhury, P.R., Biddinger, N., Shaw, J.H., Jeyasingh, P.D. & Weider, L.J.
- 900 (2018). Resurrected 'ancient' daphnia genotypes show reduced thermal stress tolerance com-
- pared to modern descendants. Royal Society Open Science, 5.
- Zettlemoyer, M.A., Schultheis, E.H. & Lau, J.A. (2019). Phenology in a warming world: differences between native and nonnative plant species. *Ecology Letters*.
- ⁹⁰⁴ Zhu, J.T., Zhang, Y.J. & Wang, W.F. (2016). Interactions between warming and soil moisture
- 905 increase overlap in reproductive phenology among species in an alpine meadow. Biology
- 906 Letters, 12, 4.