Winter temperatures dominate spring phenological responses to warming

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1 Abstract

Decades of fundamental research on woody plant species highlight three major cues that shape spring

phenological events: chilling, forcing, and photoperiod (e.g., 1; 2; 3). Increasing research on the phenological impacts of climate change has led to debate over how common these cues are across species, and—if prevalent—whether chilling and/or photoperiod cues may be slowing phenological responses to warming in recent years (4; 5; 6). Here we use a global meta-analysis of all published controlled environment studies to test for the relative effects of these cues across 203 species. We find almost all species show strong responses to all three cues, with chilling being the strongest (1.9X greater than forcing), and photoperiod the weakest (0.7X forcing). Simple forecasts from our findings for a well-studied region (Central Europe) suggest that spring phenology will continue to advance, as stalling effects of chilling generally appear above 4°C warming for most locations. Our results unify both sides of the debate over phenological cues: while all species may respond to all cues strongly in experimental conditions, in current environmental conditions the dominant signal of climate change is from increased forcing. To improve budburst forecasts under future climate change, major progress will require greater efforts to fully separate chilling and forcing effects at the physiological-level.

Main text

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For decades, plant phenology has been one of the most reported and consistent biological imprints of climate change (7): many temperate plants are leafing and flowering days to weeks earlier with rising temperatures (8; 9). Understanding such shifts is important as phenology shapes community assembly and a suite of ecosystem services, including pollination and carbon sequestration, and scales up to impact projections of climate change itself (10).

As research interest in phenology has progressed, critical discrepancies and uncertainties in our understanding have emerged. Though responses to warming are consistent on average, they show substantial variation
among species and sites (11). Furthermore, long-term observational studies provide increasing evidence that
sensitivities of phenology to temperature are weakening in recent decades (12; 13; 14), especially in Europe, where researchers suggest that responses to multiple environmental cues underlie declining temperature
sensitivities (5).

Fundamental research in phenology outlines how three major environmental cues, chilling (cool temperatures, generally occurring in the fall and late winter), forcing (warm temperatures, generally occurring in the late winter and early spring), and photoperiod (daylength), provide multiple routes to budburst each spring, depending on the environment (15). For example, in some species a cool winter will lower the amount of

forcing required to trigger budburst, compared to a warmer winter (16). Additionally, photoperiod may
trigger budburst, given low chilling and/or forcing (6; 17; 18). Research suggests that all three cues may
affect spring phenology for many temperate woody species (3; 17; 19), which could have critical forecasting
implications—predicting delays in spring phenology as increased warming reduces chilling in some areas (20)
or where earlier budburst shortens the experienced photoperiod. However, there is strong debate, with some
research suggesting some cues—such as photoperiod—may be effectively absent in some species, but dominate
in others (6; 21; 17; 22).

Resolving this debate requires overcoming major hurdles to estimate responses to each cue. Studies attempting to estimate cues using long-term observational data (e.g., 6; 23) generally fail to overcome the 40 fundamental challenge that cues are strongly correlated in nature (e.g., during the seasonal transition from winter to spring at temperate latitudes, forcing and photoperiod usually increase in step for a given location; 42 mean estimated chilling and spring temperatures can be positively correlated in space). In contrast to observational studies and experimental field warming studies designed to test higher temperatures in natural conditions (11), controlled environment experiments can break down correlations between the cues. These experiments, which generally rely on dormant tree cuttings or dormant plants exposed to controlled temperature and light regimes in growth chambers (Fig. 1), have been shown to replicate whole-plant responses 47 in nature (24). Such experiments have been conducted for decades (though each experiment generally lasts under a year). They have produced contrasting results, however, potentially due to differences in focal species or study sites (6; 18; 25; 26; 27). Resolving these discrepancies is critical to accurate predictions of spring phenology, especially as continued climate change will yield warmer temperatures than has been experienced 51 in at least 150 years and warming will occur at a rapid pace (28; 29; 30; 31; 32).

Here, we leverage controlled environment studies through a meta-analysis to understand how chilling, forcing, and photoperiod determine budburst timing in woody species. We reviewed 201 papers from controlled environment studies, then extracted data from all experiments that reported budburst responses, yielding data from 72 studies and 203 species (Fig. S1, Tables S1, S3). The resulting Observed Spring Phenology Responses in Experimental Environments (OSPREE) database includes only studies for which we could identify forcing, photoperiod, and chilling treatments quantitatively and includes a mix of studies where dormant plant tissue (grown in greenhouses or taken directly from the field) was exposed to experimental conditions (33). Most experiments reported forcing and photoperiod treatments, while chilling occurred mainly in the field, though some studies additionally applied chilling before moving plants into forcing conditions (Fig. 1). Because chilling was rarely reported, we calculated an estimate of chilling (both in the field and in experimental conditions), using a common but approximate method (34), based on a hypothesis of how chilling accumulates (35), with no chilling accumulating below 1.4°C or above 12.4°C (throughout the main text we use the term 'chill unit,' see Supplemental Materials, especially Table S2, for details).

We estimated the effects of chilling, forcing, and photoperiod using a Bayesian hierarchical model. Our model averages over interactive effects of predictors, including only main effects that we could more robustly estimate given current study designs (see *Methods* in Supplemental Materials). Species are modeled hierarchically, producing estimates of both species-level responses (generally yielding more accurate estimates for well-studied species, such as *Fagus sylvatica* and *Betula pendula*), and the distribution from which they are drawn, yielding an estimate of the overall response across species (see *Methods* in Supplemental Materials):

$$y_i = \alpha_{sp[i]} + \beta_{forcing_{sp[i]}} + \beta_{photoperiod_{sp[i]}} + \beta_{chilling_{sp[i]}} + \epsilon_i$$

 $\epsilon_i \sim N(0, \sigma_y^2)$

The α and each of the three β coefficients were modeled at the species level, as follows:

$$lpha_{sp} \sim N(\mu_{lpha}, \sigma_{lpha})$$
 $eta_{forcing_{sp}} \sim N(\mu_{forcing}, \sigma_{forcing})$
 $eta_{photoperiod_{sp}} \sim N(\mu_{photoperiod}, \sigma_{photoperiod})$
 $eta_{chillingsp} \sim N(\mu_{chilling}, \sigma_{chilling})$

where i represents each unique observation, sp is the species or species complex grouping, α represents the intercept, β terms represent slope estimates, and y is the days to budburst since forcing conditions were applied. Some species were represented in only one dataset in the OSPREE database, making it impossible to statistically differentiate between species and dataset for these taxa. To address this, we focus on estimates

(reported as mean with 95% uncertainty intervals, unless otherwise noted) from a model of 67 taxa, which
were included in multiple datasets and treatments (generally this occurred at the species-level, but in some
cases we collapsed species found in only one study into "complexes" at the level of genera, see the *The Observed*Spring Phenology Responses in Experimental Environments (OSPREE) database section in the Supplemental
Materials for details). Estimates from this model were generally similar to estimates from a model of all 203
species (grouped into 37 species or 'species complex' groupings, Tables S4, S5). We fit several additional
models, including a model testing latitude effects on cue estimates and one testing effects of chilling study
design (see Models in the Supplemental Materials for model equations and other details).

Across studies, all cues—chilling, forcing, and photoperiod—advance budburst phenology (Fig. 2, Tables S4, S5). Chilling was the strongest cue (-8.33 days/standard unit [-11.5 to -5.3] or -2.76 days per chill unit [-3.65 to -1.89]), followed by forcing (-4.35 days/standard unit [-6.56 to -2.08] or -0.8 days per °C of warming, [-1.18 86 to -0.43]), and photoperiod (-2.96 days/standard unit [-5.43 to -0.56] or -0.53 days per hour of daylight [-0.92 to -0.15; Figs. S4, 3, 4; Tables S4, S5; see Supplemental Materials for more details). While photoperiod had the smallest effect among the three cues, our results contrast with the extensive literature suggesting photoperiod is an unimportant cue for many species (6: 36)—instead we found it was surprisingly large, even when accounting for its interaction with provenance latitude (i.e., the latitude of origin for plant material; see Supplemental Materials for details, especially Figs. S3, S10, Table S7). It was also generally consistent across species (variance = 5.18 days per hour of daylength in the standardized model), only deviating in Faqus 93 sylvatica, a species well-known for having a large response to photoperiod (which we also found, Figs. 2, S3). Species responses to chilling were slightly more variable (variance = 7.2 days per chill unit in the standardized 95 model, Fig. 2) than responses to forcing (variance = 5.72 days per forcing unit in the standardized model Fig. 2).

As temperature is fundamentally altered by climate change, our finding that different ends of the temperature spectrum—chilling and forcing—have the strongest effects on budburst suggests that understanding these two cues will be critical for forecasting phenology with climate change. Many previous studies attribute advances in budburst to increased forcing (9; 16; 17; 37). Our results, however, suggest that, across 203 species and 72 controlled environment studies, chilling has a greater effect on budburst than forcing (Figs. 2, S3, S7; Tables S4-S7). This has not been widely suggested previously, perhaps because little work has directly manipulated chilling, and the few studies that have were designed to compare chilling versus photoperiod effects (e.g., 6; 17; 19; 25), not chilling versus forcing effects. Process-based phenological models, however, that explicitly

model chilling often find it is most critical (e.g., 38).

Despite its apparent importance, chilling and its related physiological stage of endodormancy, are not well understood (15). Physiologically plants appear to accumulate forcing only after they have exited endodormancy (and entered ecodormancy, Fig. 1), which is generally thought to occur when chilling requirements have been met (15). Thus, while researchers generally define 'chilling' and 'forcing' treatments based on temperatures in controlled environments (including in the studies used here, see Fig. 1), fully separating out what plants experience as chilling versus forcing will likely require new methods to measure endo- and ecodormancy (39).

Until then, researchers must generally rely on modeled estimates of chilling, as we have used here. Models 114 of how species accumulate chilling are poorly developed for forest trees, with few relevant tests evaluating 115 the particular temperatures at which species do or do not accumulate chilling. Instead, researchers generally 116 rely on models developed for perennial fruit trees, i.e., Utah (34) and chill portions (40), both of which were 117 developed for peach species. These models are themselves hypotheses for how chilling may accumulate and 118 produce dormancy release, but are likely to be inaccurate for many species (35). We found that applying a different chilling model did not strongly affect our estimates (i.e., 95\% uncertainty intervals of estimates 120 for chilling, photoperiod, and forcing overlapped for models using Utah and chill portions, with standardized predictors, Table S4). 122

Progress on developing chilling models for wild species may be especially slow as only a small portion of studies 123 (13 of the total 72 studies) manipulate chilling directly. Instead many studies estimate chilling effects through 124 sequential removal of tissue from the field followed by exposure to 'forcing' conditions (Fig.1a,1b, 25 out of 72; the remaining studies did not appear to manipulate chilling), with the assumption that tissues collected later 126 experience more chilling (41). This method benefits from more natural chilling conditions but introduces other challenges: first, chilling duration may not always co-vary with the magnitude of total accumulated 128 chilling (35), and, second, photoperiod and other factors also change over time. Indeed, we found that 129 sequential-removal studies tended to result in later budburst, weaker effects of chilling, and stronger effects 130 of forcing compared to estimates from studies that directly manipulated chilling (Fig. S7, Table S8, (41; 42). 131 This suggests that a study's design of chilling manipulation impacts both forcing and chilling estimates and 132 further supports that an improved understanding of chilling could in turn alter our understanding of forcing. 133 Linking such short-term controlled experiments to natural conditions robustly will require more efforts to 134 understand the complex interactions between chilling, forcing, and photoperiod that we were not able to 135

quantify in this meta-analysis. Most experimental studies do not test for interactions between all three cues (Table S11). Further, many additional factors can affect phenological responses, including ontogeny (Table S10, 43)), latitude (Fig. S3), and air humidity (44).

Despite these limitations, a simple interpretation of our results does support the widespread hypotheses that chilling and/or photoperiod cues may underlie declining sensitivities to warming in long-term Central European data (5; 12; 13). Under these hypotheses, warming increases forcing and thus advances budburst, but such advances become muted if warming also causes declines in chilling and shorter photoperiods experienced near the timing of budburst (38). This basic agreement between our results and long-term observational trends integrates across experimental conditions that encompass more extreme scenarios than may be seen in nature (Fig.3,S4). A more robust comparison requires examining predictions under conditions closer to those found in nature.

Reinterpreting our model using climate and phenology data that have led to observations of declining temper-147 ature sensitivities in Central Europe suggests that chilling and photoperiod are unlikely to cause the observed 148 declines. Our model predicts such declines only at extreme warming for most sites (see Supplemental Materials). In contrast to the common hypothesis that plants experience less chilling with global warming, 150 we found that—for many sites—total estimated chilling increased with warming (Fig. S9A, C), though this varied with local climate prior to warming (Figs. S6 - ??). Portions of Central Europe have experienced more 152 dramatic warming in winter versus summer (45, though this certainly varies over time and space, (author?) 153 (46)). Yet even if warming uniquely occurs in the winter, our results suggest that delays due to decreased 154 chilling only occur at warming above at least 4°C for most sites, though responses vary by species (Fig. S8, 155 S9, S6). At high warming, predicted declines in sensitivity were due to declines in chilling—photoperiod had 156 comparatively little effect on budburst day of year, even for the photosensitive species F. sylvatica (Fig. S10). 157 Our predictions leave open the question of what underlies declining sensitivities across Europe, but one pos-158 sibility is that it may be a statistical artifact of how temperature sensitivities are calculated. Physiologically, 159 budburst is triggered by the accumulation of forcing temperatures during the spring (15; 47). However, 160 researchers today often estimate temperature sensitivities from long-term observational data using a linear 161 regression of annual budburst date versus mean spring temperature, or other aggregated temperature metrics 162 (e.g., 11). This approach has the benefit of yielding an easily interpretable metric—days change per °C—but 163 will estimate systematically lower sensitivities given warmer daily temperatures, even with no change in the underlying cues (Fig. S11). We found the declining sensitivities observed in European data are of the same 165

magnitude as those predicted from a statistical artifact (sensitivity declines of 0.8 ± 0.3 days/°C in European data versus 0.9 ± 0.5 days/°C in simulations), and the data also show a related decline in leafout variance that would not be immediately predicted from shifting cues (see *Potential statistical artifacts in declines of temperature sensitivity in observational long-term data* in the Supplemental Materials and 48, for further details). This statistical artifact is likely not confined to phenological studies; it should apply to any research using a similar days/°C metric to estimate an underlying thermal accumulation model where the thermal sum per day is non-stationary, as is the case with climate change.

Our results unify decades of experimental studies, which have shown the importance of chilling, forcing,
and daylength to determining budburst timing, with long-term observational data, where forcing appears to
dominate responses to recent warming. We do not find strong evidence for delaying budburst in the near
future, and suggest that recent observed declines in temperature sensitivity may be related to statistical
artifacts from estimating complex cues from observational data. Instead, our predictions suggest budburst
will continue to advance in many well-studied European regions in the future. The most dramatic changes
will come from regions where winter warming causes dramatic decreases in chilling, with implications for
ecosystem services related to phenology. Thus identifying processes that plants undergo when accumulating
chilling versus forcing will be critical for the most accurate forecasts (15)

82 References

- 183 [1] R. K. Campbell, A. I. Sugano, *Botanical Gazette* pp. 290–298 (1975).
- ¹⁸⁴ [2] O. M. Heide, Scientia Horticulturae **115**, 309 (2008).
- 185 [3] D. F. B. Flynn, E. M. Wolkovich, New Phytologist **219**, 1353 (2018).
- [4] O. M. Heide, Journal of Experimental Botany 62, 5397 (2011).
- ¹⁸⁷ [5] Y. S. H. Fu, et al., Nature **526**, 104 (2015).
- [6] C. M. Zohner, B. M. Benito, J. C. Svenning, S. S. Renner, Nature Climate Change 6, 1120 (2016).
- [7] IPCC, Climate Change 2014: Impacts, Adaptation, and Vulnerability (Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2014).
- ¹⁹¹ [8] A. J. Miller-Rushing, R. B. Primack, *Ecology* **89**, 332 (2008).

- ¹⁹² [9] A. Menzel, et al., Global Change Biology **12**, 1969 (2006).
- [10] E. E. Cleland, I. Chuine, A. Menzel, H. A. Mooney, M. D. Schwartz, Trends in Ecology & Evolution 22,
 357 (2007).
- 195 [11] E. M. Wolkovich, et al., Nature 485, 494 (2012).
- [12] T. Rutishauser, J. Luterbacher, C. Defila, D. Frank, H. Wanner, Geophysical Research Letters 35, L05703 (2008). Rutishauser, This Luterbacher, Juerg Defila, Claudio Frank, David Wanner, Heinz.
- [13] H. Y. Yu, E. Luedeling, J. C. Xu, Proceedings of the National Academy of Sciences of the United States
 of America 107, 22151 (2010).
- ²⁰⁰ [14] X. Wang, et al., Nature communications **10**, 2389 (2019).
- ²⁰¹ [15] I. Chuine, et al., Global Change Biology **22**, 3444 (2016).
- ²⁰² [16] C. A. Harrington, P. J. Gould, Frontiers in Plant Science 6, 120 (2015).
- ²⁰³ [17] D. Basler, C. Körner, *Tree Physiology* **34**, 377 (2014).
- ²⁰⁴ [18] A. Caffarra, A. Donnelly, I. Chuine, M. B. Jones, Climate Research 46, 147 (2011).
- ²⁰⁵ [19] A. Caffarra, A. Donnelly, I. Chuine, Climate Research 46, 159 (2011).
- ²⁰⁶ [20] H. Fraga, J. G. Pinto, J. A. Santos, Climatic Change pp. 1–15 (2019).
- ²⁰⁷ [21] O. Heide, *Physiologia Plantarum* **88**, 531 (1993).
- [22] R. K. Singh, T. Svystun, B. AlDahmash, A. M. Jönsson, R. P. Bhalerao, New Phytologist 213, 511
 (2017).
- ²¹⁰ [23] Y. Vitasse, D. Basler, European Journal of Forest Research 132, 1 (2013).
- ²¹¹ [24] Y. Vitasse, D. Basler, *Tree physiology* **34**, 174 (2014).
- ²¹² [25] J. Laube, et al., Global Change Biology **20**, 170 (2014).
- ²¹³ [26] D. Basler, C. Körner, Agricultural and Forest Meteorology 165, 73 (2012).
- ²¹⁴ [27] A. Caffarra, A. Donnelly, International Journal of Biometeorology **55**, 711 (2011).
- ²¹⁵ [28] R. Ohlemüller, E. S. Gritti, M. T. Sykes, C. D. Thomas, Global Ecology and Biogeography 15, 395 (2006).

- ²¹⁶ [29] J. W. Williams, S. T. Jackson, Frontiers in Ecology and the Environment 5, 475 (2007).
- [30] J. W. Williams, S. T. Jackson, J. E. Kutzbacht, Proceedings of the National Academy of Sciences of the
 United States of America 104, 5738 (2007).
- [31] T. Stocker, D. Qin, G. Platner, Working Group I Contribution to the Fifth Assessment Report of the
 Intergovernmental Panel on Climate Change. Summary for Policymakers (IPCC, 2013) (2013).
- 221 [32] Y. Xu, V. Ramanathan, D. G. Victor, Global warming will happen faster than we think (2018).
- [33] E. M. Wolkovich, et al., Observed Spring Phenology Responses in Experimental Environments (OS-PREE), doi:10.5063/F1QV3JQR (2019).
- ²²⁴ [34] E. Richardson, *HortScience* **9**, 331 (1974).
- ²²⁵ [35] F. Dennis, *HortScience* **38**, 347 (2003).
- ²²⁶ [36] Y. H. Fu, et al., Global change biology **25**, 1696 (2019).
- [37] N. L. Bradley, A. C. Leopold, J. Ross, W. Huffaker, Proceedings of the National Academy of Sciences
 96, 9701 (1999).
- ²²⁹ [38] J. Gauzere, C. Lucas, O. Ronce, H. Davi, I. Chuine, Ecological Modelling 441, 108805 (2019).
- 230 [39] C. van der Schoot, L. K. Paul, P. L. H. Rinne, Journal of Experimental Botany 65, 1699 (2014).
- ²³¹ [40] S. Fishman, A. Erez, G. Couvillon, Journal of Theoretical Biology 124, 473 (1987).
- [41] J. H. Weinberger, et al., Proceedings. American Society for Horticultural Science (1950), vol. 56, pp.
 122–28.
- [42] C. A. Polgar, R. B. Primack, E. H. Williams, S. Stichter, C. Hitchcock, Biological Conservation 160, 25
 (2013).
- ²³⁶ [43] Y. Vitasse, New Phytologist **198**, 149 (2013).
- ²³⁷ [44] J. Laube, T. H. Sparks, N. Estrella, A. Menzel, New Phytologist **202**, 350 (2014).
- ²²⁸ [45] R. C. J. Balling, P. J. Michaels, P. C. Knappenberger, Climate Research 9, 175 (1998).
- ²³⁹ [46] C. Li, B. Stevens, J. Marotzke, Geophysical Research Letters 42, 8131 (2015).
- ²⁴⁰ [47] H. Hänninen, Canadian Journal of Botany **73**, 183 (1995).

²⁴¹ [48] S. Güsewell, R. Furrer, R. Gehrig, B. Pietragalla, Global Change Biology 23, 5189 (2017).

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$_{251}$ Figures

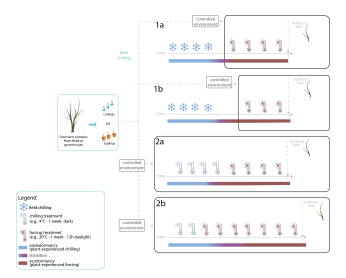


Figure 1: Controlled environment experiments that test temperature effects on woody plant phenology as we synthesize here generally manipulate photoperiod and temperature cues, which include chilling and forcing. Chilling is manipulated via either using natural chilling in the field (1A-B, in which plant material is collected after different numbers of days in the fall/winter) or experimentally (2A-B, in which plant material is place in controlled environment chambers set to different chilling temperatures and/or durations). Chilling treatments are designed to break plant endodormancy (field or experimental), after which forcing treatments are imposed by moving plant material to warmer temperatures (eco-dormancy) that allow budburst to occur (this ideal is shown in panel 1A, though it could also occur with experimentally applied chilling). A challenge with controlled environment studies is that species-specific chilling requirements are rarely known, so the experimental manipulations may not always align with what the plant experiences. Thus, in some cases, researchers' estimates of chilling may bridge across what plants experience as chilling and forcing (1B and 2A, where plants transition into ecodormancy before 'forcing' treatments are applied), or plants may end before endodormancy is fully broken (2B). In the studies synthesized here, photoperiod (not shown) is most often manipulated in forcing treatments.

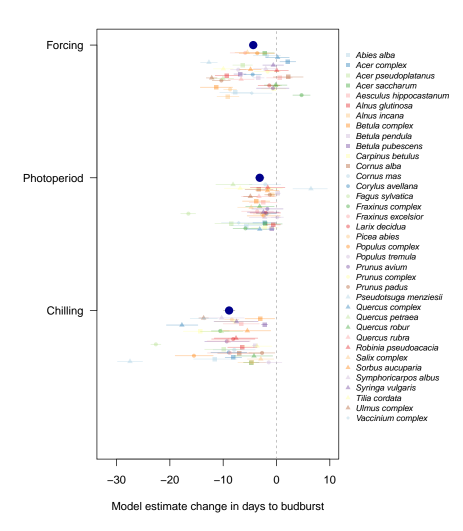


Figure 2: Estimated effects of chilling, forcing, and photoperiod on budburst timing across 67 (modeled as 37 separate taxa, see 'Models' section in the Supplemental Materials) in 42 controlled environment studies. Using standardized units, which allow comparisons across cues, we show that most species (smaller symbols) are responsive to most cues, with chilling being the strongest cue when considering overall estimates across species (larger, dark blue circles). Overall estimates shown here were generally similar to other model formulations, including using data from 203 species, and using different methods for calculating chilling (Figs. S3, S7; Tables S4-S7). Lines represent 50% uncertainty intervals (other intervals provided in Tables S4-S7)

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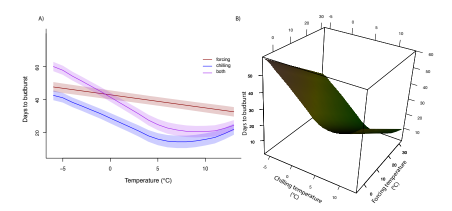


Figure 3: Estimates of budburst across a range of forcing temperatures and estimated chilling (converted to a representative mean temperature, see *Estimating chilling* in online Methods) based on overall estimates of chilling and forcing effects (Fig. 1). Note that days to budburst is relative to experimental methods and thus not comparable to day of year in the field, shading represents 50% uncertainty intervals. Panel A shows the effect of chilling temperature on budburst, with forcing kept at the mean level across all experiments (16°C); the effect of forcing temperature with chilling kept at the mean level across all experiments (1324 chilling units), and the effect of varying both chilling and forcing temperatures simultaneously. Panel B shows all possible combinations of chilling, forcing, and budburst across the experimental conditions. Maximum advances in budburst occur at intermediate chilling temperatures (*e.g.*, here at mean winter temperatures of 6-7°C) and the highest forcing (here at 32°C). We set photoperiod to eight hours, which is the most common photoperiod treatment in the database.

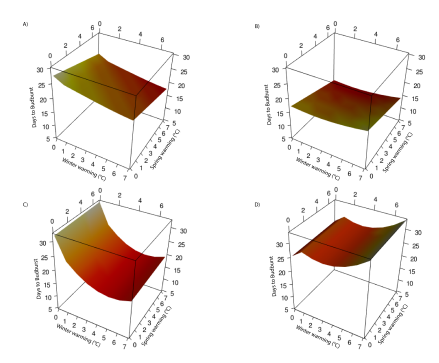


Figure 4: Implications of warming on budburst timing varies across species and sites, depending strongly on pre-warming climate conditions related to chilling for each site. Here we show species-level estimates from our model (Fig. 2) for the two most common species in the OSPREE database: Betula pendula (A, B) and Fagus sylvatica (C, D), for sites that highlight the diversity of possible budburst responses to warming (Fig. S6, which shows general trends across many sites in Central Europe). In some sites, warming increases total chilling estimates (A, C) leading to greater advances in budburst (compared to forcing alone), whereas warming decreases total chilling estimates in other sites (B, D), leading to smaller advances and, eventually, delays with substantial warming. Compare this to Fig. ?? in the Supplemental Materials, which is a simplified two-dimensional version.