Local temperatures predict breeding phenology but do not result in breeding synchrony among a community of resident cavity-nesting birds

Anna Drake & Kathy Martin

Weather and ecological factors are known to influence breeding phenology and thus individual fitness. We predicted concordance between weather conditions and annual variation in phenology within a community of eight resident, cavity-nesting bird species over a 17-year period. We show that, although clutch initiation dates for six of our eight species are correlated with local daily maximum temperatures, this common driver does not produce a high degree of breeding synchrony due to species-specific responses to conditions during different periods of the preceding winter or spring. These “critical temperature periods” were positively associated with average lay date for each species, although the interval between critical periods and clutch initiation varied from 4–78 days. The ecological factors we examined (cavity availability and a food pulse) had an additional influence on timing in only one of our eight focal species. Our results have strong implications for understanding heterogeneous wildlife responses to climate change: divergent responses would be expected within communities where species respond to local conditions within different temporal windows, due to differing warming trends between winter and spring. Our system therefore indicates that climate change could alter relative breeding phenology among sympatric species in temperate ecosystems.

In seasonal environments, favorable conditions for producing and rearing young are restricted to a brief temporal window and timing of breeding is a major determinant of individual fitness. Photoperiod and temperature are major determinants of such windows within temperate landscapes due to their impact on food abundance and metabolic costs. Correlations between local environmental factors, particularly temperature, and clutch initiation are well documented for both migrant and resident bird species. Fitness costs associated with mis-timed breeding relative to local conditions are also well documented. For example, individuals that begin nesting too early in a given year may experience adverse weather conditions that result in clutch loss, lower quality eggs, or reduced personal survival or condition. Individuals breeding too late may miss peak resource availability, lay fewer eggs, or have inadequate time to produce a second brood or to re-nest if an earlier nest is depredated. Young from late nests additionally have less time to achieve independence before conditions deteriorate, which may result in lower recruitment rates than early-initiated clutches.

The role of local ecological factors that are not weather-dependent in altering avian breeding phenology is less studied. Periodic outbreaks in forest insects may remove caloric constraints in early spring and thereby enable females to advance laying dates. In cavity-nesting vertebrate communities, nest site limitation may also delay breeding. For birds that are secondary cavity-nesting species (those relying exclusively on existing cavities for nesting), this may be driven by competition for breeding sites; for facultative excavators (species that can make cavities, but will also reuse established sites), limited cavity availability can result in delays associated with cavity excavation. Such ecological factors, combined with environmental cues, should influence timing in

1Department of Forest and Conservation Sciences, University of British Columbia, 2424 Main Mall, Vancouver, British Columbia, V6T 1Z4, Canada. 2Science & Technology Branch, Environment and Climate Change Canada, 5421 Robertson Road, R.R. 1, Delta, British Columbia, V4K 3N2, Canada. Correspondence and requests for materials should be addressed to A.D. (email: aedrake@mail.ubc.ca)
In this study, we use a 17-year dataset (1995–2011) to compare breeding phenology at the community-level for a group of eight resident, largely insectivorous, tree cavity-nesting bird species in the interior of British Columbia, Canada (four woodpeckers, three small insectivores, and one small owl species (Table 1)). Although dietary differences are likely to influence when these species initiate breeding, we predicted that annual variation in lay date would show similar directionality among our species, such that breeding would occur relatively earlier in some years and relatively later in others, across the community. We test this using pair-wise correlation analyses. We then examine whether timing of breeding is correlated with local weather cues (temperature and/or rainfall) prior to breeding and whether there is an additional effect of local ecological factors (mountain pine beetle abundance (hereafter, MPB) and cavity availability). We predicted that these resident species would show similar responses to local weather prior to breeding. Additionally, we predicted that increased MPB availability for all species (except the owl), and nest site availability among secondary and facultative cavity-nesters, would result in earlier breeding dates.

**Results**

**Breeding synchrony.** Our resident species differed in when, on average, they initiated breeding. Northern saw-whet owl and hairy woodpeckers showed the earliest mean breeding dates (multi-year average: May 4th and May 8th, respectively; Fig. 1) while red-breasted nuthatch and downy woodpecker bred the latest (May 28th and May 31st, respectively; Fig. 1). All species included in our analysis showed among-year variation in mean clutch initiation date, but only 2 of our 28 possible species dyads showed a significant correlation in timing (Table 2). Northern saw-whet owl, hairy, and pileated woodpecker showed the greatest independence in timing among-years (Table 2). In contrast, the three smallest insectivores showed some correlation in timing of breeding: mountain chickadee timing corresponded with that of black-capped chickadee (Pearson’s r = 0.84, p < 0.001, n = 16 years) and, to a lesser degree, with that of red-breasted nuthatch (Pearson’s r = 0.66, p = 0.05, n = 17 years).

**Weather cues.** The lay dates of the majority of the species we examined (all except pileated woodpecker and northern saw-whet owl) showed a negative relationship (95% CI) with annual maximum daily temperatures (i.e. breeding occurred earlier in years that were warmer (Table 3, Fig. 2)). Models using average daily minimum, and average daily mean temperatures were comparable to average daily maximum temperatures for explaining hairy woodpecker phenology. The temporal period in which temperatures best predicted lay dates differed markedly among species, as illustrated in Fig. 3: hairy woodpecker clutch initiation dates corresponded most strongly to temperatures in late winter (maximum temperature best window: Jan 7-Feb 19; minimum and mean temperatures: Jan 14-Jan 24), while higher maximum temperatures in March, April and May were associated with advanced clutch initiation for chickadee species (best windows for black-capped: March 4-April 13, mountain chickadee: March 7-May 20) and red-breasted nuthatch (March 9-April 5). American three-toed woodpecker phenology corresponded with maximum temperatures in late April and early May (April 28-May 8). Phenology of the latest initiator, the downy woodpecker, also most strongly corresponded to maximum temperatures in April and May (April 19-May 19). The interval between these critical temperature periods and clutch initiation was greater than 1 month for black-capped chickadees and red-breasted nuthatch (41 and 53 days respectively) and 2.5 months for hairy woodpecker (78 days; Fig. 3). Mountain chickadee, downy, and American three-toed woodpecker initiated nesting within 1 month of their critical temperature periods (4, 12, and 18 days respectively; Fig. 3).

There was no evidence of a temporal trend between 1995 and 2011 for either temperature (Fig. 4A; p > 0.21 for all months, n = 17 years) or for breeding phenology among seven of the eight cavity-nesters we assessed. Red-breasted nuthatch showed a trend toward later lay dates over the study period (temporal model coefficient (median (95% CI range)): 0.50 days/year (0.26,0.74)). However, this temporal model was not competitive with the daily maximum temperature model reported in Table 3. Local rainfall was not associated with lay dates for any of our species.

| Species                        | Nesting strategy          | MPB consumer | Yes/No (stage) |
|-------------------------------|---------------------------|--------------|----------------|
| Black-capped chickadee (Poecile atricapillus) | Obligate excavator | Y (L, A)     |                |
| Downy woodpecker (Picoides pubescens) | Obligate excavator | Y (L, A)     |                |
| Hairy woodpecker (Picoides villosus) | Obligate excavator | Y (L, A)     |                |
| American three-toed woodpecker (Picoides dorsalis) | Obligate excavator | Y (L, A)     |                |
| Pileated woodpecker (Dryocopus pileatus) | Obligate excavator | Y (L, A)     |                |
| Red-breasted nuthatch (Sitta Canadensis) | Facultative excavator | Y (L, A)     |                |
| Mountain chickadee (Poecile gambeli) | Secondary cavity-nester | Y (L, A)     |                |
| Northern saw-whet owl (Aegolius acadicus) | Secondary cavity-nester | N             |                |

Table 1. Interior of British Columbia cavity-nesting species monitored in the study. Nesting strategies and whether these species consume mountain pine beetle (MPB; Dendroctonus ponderosae) during its larval (L) or adult (A) stage are noted.
Ecological cues. Contrary to our predictions, we found no support for an additional direct influence of MPB abundance on clutch initiation dates (beyond the possible temperature influences). Similarly, greater cavity availability was not associated with advanced clutch initiation in red-breasted nuthatch, mountain chickadee or northern saw-whet owl. Counter-intuitively, red-breasted nuthatch instead showed weak delays associated with increasing small and medium cavity availability (Table 3).

Model performance. Marginal and conditional $R^2$ values describing the model fit to all nest records are presented in Table 3. To more intuitively present the amount of among-year variation in timing explained by average daily maximum temperatures in our species models, we additionally calculated an $R^2$ value using model-predicted versus observed mean annual clutch initiation dates. Temperature variation among years explained 78%, 65% and 54% of the variation in mean annual clutch initiation dates for American three-toed (n = 15 years), downy (n = 15), and hairy woodpeckers (n = 16), respectively. Temperature explained 77%, 48%, and 48% of the variation in mean mountain chickadee (n = 17 years), black-capped chickadee (n = 17), and red-breasted nuthatch initiation dates among years (n = 17; Fig. 2). None of our candidate explanatory variables described the variance in pileated woodpecker or northern saw-whet owl clutch initiation dates; weather models for both species did not pass our Type 1 error test (see Methods). Small sample sizes (an average of 3.3 recorded nests/assessed year for...
Table 3. The influence of the average maximum temperatures during species-specific ‘response periods’ on the annual timing of clutch initiation for resident cavity-nesting birds in the interior of British Columbia (days after January 1; n = nest records). Red-breasted nuthatch showed an additional effect of cavity availability on timing. Coefficients are the median (95% CI range) values of coefficients derived from 2000 (100 MI × 20 bootstrap) runs of the top-performing model for each species (date ~ fixed effects + a random ‘Year’ effect). The amount of variation explained by model variables is indicated by the marginal R^2 (variation explained by fixed effects only). Small conditional R^2 values (the variation explained by fixed effects + the random ‘Year’ effect) indicate large among-individual variation (i.e. low within-species synchrony) in clutch initiation, within-year.

| Model Variables | Mountain chickadee (n = 614) | Black-capped chickadee (n = 84) | Red-breasted nuthatch (n = 478) | Downy woodpecker (n = 110) | Hairy woodpecker (n = 108) | Pileated woodpecker (n = 43) | American three-toed woodpecker (n = 85) | Northern saw-whet owl (n = 36) |
|-----------------|-----------------------------|-----------------------------|-------------------------------|------------------------|---------------------------|-----------------------------|-----------------------------|-----------------------------|
| Intercept       | 169.13 (164.23, 174.83)     | 155.17 (149.31, 161.02)     | 158.68 (154.42, 162.95)       | 186.81 (176.14, 197.73) | 125.53 (123.57, 127.57)   | 165.76 (156.36, 175.82)     | —                           | —                           |
| Maximum temperature | −2.51 (-2.97, −2.03)     | −1.85 (-2.66, −1.03)        | −2.25 (-2.89, −1.81)          | −2.67 (-3.43, −1.90)     | −3.23 (-3.54, −1.26)       | −1.54 (-2.33, −0.83)         | —                           | —                           |
| Cavity availability | —                      | —                          | 0.01 (0.005, 0.02)            | —                      | —                          | —                           | —                           | —                           |
| Marginal R^2    | 0.17                       | 0.24                       | 0.18                          | 0.34                    | 0.19                       | —                           | 0.12                        | —                           |
| Conditional R^2  | 0.21                       | 0.34                       | 0.31                          | 0.52                    | 0.30                       | —                           | 0.22                        | —                           |

Discussion

We examined the underlying influences of weather factors as well as the role of other ecological resource constraints in driving annual variation in breeding phenology over a 17-year period across a suite of sympatric resident bird species within a temperate mixed forest ecosystem. To our knowledge, this is the largest community of sympatrically breeding birds where the drivers of timing of breeding have been examined. As expected, phenology was correlated with temperature within our community. Specifically, local maximum daily temperatures were correlated with clutch initiation dates for 6 of the 8 species we examined (owls and pileated woodpeckers had no detectable response to any weather variables). Contrary to our expectations, however, this commonality did not translate into similar variation in clutch initiation among years. Only the small insectivorous species (chickadees and red-breasted nuthatch) showed similar annual variation in clutch initiation. Different annual responses in breeding phenology among species corresponded to different intervals in the time period, prior to breeding, where temperature appeared to be influential. These “critical temperature periods” tended to be positively related to the average onset of clutch initiation for each species: the earliest breeder in our dataset (hairy woodpecker) adjusted clutch initiation in response to winter temperatures (January and February), while the later breeders (the congeneric American three-toed and downy woodpecker) showed the strongest correlations with April and May temperatures. None of our species responded to annual variation in precipitation.

The interval between critical temperature period and average clutch initiation date varied widely among species in our study. Downy and three-toed woodpeckers appeared to respond to local conditions within a period that was close to their average clutch initiation date, as did mountain chickadees. In contrast, black-capped chickadees and red-breasted nuthatch responded to local temperatures more than one month prior to clutch initiation (41 and 53 days, respectively) and hairy woodpeckers appeared to respond to local temperatures more than two months prior to their average lay date (78 days). These lags are longer than is generally reported for birds (less than 1 month23) and temperature may therefore be acting as a breeding cue via secondary factors such as insect abundance24.

If distinct critical periods within our community are the product of species-specific food sources that increase in abundance or availability at different threshold temperatures, we might predict a relationship between a species’ response period or average breeding onset and their foraging guild. While our two foliage-gleaning species (mountain and black-capped chickadees) both responded to March temperatures and initiated laying within the same time window, there was no shared response or lay period among the bark-gleaners. Our bark-gleaning congenersics responded to local temperatures ranging from winter (hairy woodpecker) until May (downy and American three-toed woodpecker) and exhibited notably different multi-year average lay dates. The dramatic differences we observed in breeding phenology of Picoides woodpeckers, with hairy woodpeckers initiating clutches several weeks earlier than downy woodpeckers (May 8 vs. May 31 respectively) have also been observed in the boreal mixedwood forest of northwestern Quebec where hairy woodpeckers initiated clutches more than three weeks earlier than downy woodpeckers (mean initiation of May 9 vs. May 31, respectively; Pierre Drapeau, personal communication). This indicates that our observations may be general among sympatric Picoides woodpeckers, although the underlying drivers of such divergence are unknown.

Earlier work in the interior of British Columbia found significant positive relationships between mountain pine beetle numbers and breeding densities for most of the woodpeckers, and both density and fecundity for mountain chickadee and red-breasted nuthatch25–28. Here, we found no evidence of an additional influence of beetle numbers on the timing of breeding for any of these species. Thus, the observed increased functional and numerical responses reported earlier for our study site do not appear to be related to earlier clutch initiation, but instead to a direct effect of the food subsidy on productivity. Cavity availability also did not appear to influence the breeding phenology of our secondary and facultative tree cavity-nesting species with the exception of a weak, positive relationship for red-breasted nuthatch (0.01 day delay/available cavity; Table 3). Red-breasted nuthatch showed an additional effect of cavity availability on timing.
are facultative excavators and this result may be partly explained by an increase in cavity excavation during the mountain pine beetle outbreak when nuthatch density increased and individuals shifted breeding sites to patches where beetles were abundant but cavities were not.

Our results suggest that the monitoring of a few focal species as a means to understand local responses to climate change is problematic. Warming trends associated with climate change have been heterogeneous across seasons and more pronounced in winter in both Europe and western Canada (e.g., refs 30,31). Although we did not detect a temporal trend in average maximum temperatures over our 17-year study period, the interior of British...
Columbia has experienced an average warming of 1 °C over the past 100 years. This warming has occurred disproportionately in the winter, with an increase of 1.6 °C vs. 0.8 °C in spring and summer combined. We would therefore expect that hairy woodpecker breeding dates have advanced more than other species in our community. That daily maximum temperatures, rather than minimum or means, showed the strongest correlation with phenology in our community is also noteworthy. Within our study region, maxima have increased at a slower rate than minima and, globally, maximum temperatures have increased at approximately half the rate of minimums (1950–2004). Thus resident cavity-nesting species may be somewhat buffered from climate-driven variation influencing their temporal response periods, at least initially. Overall, seasonal differences in temperature increases associated with climate change, combined with different species response periods (as observed in our study), as well as the type of temperature cue used (daily maxima, minima or means) may explain some of the heterogeneous responses of birds to climate change observed in numerous studies (e.g.34,35). These differences may therefore not represent a failure by some species to track conditions, but instead species-specific differences.
in what conditions are being tracked. Further work explicitly quantifying food availability would be needed to establish this. Examining the responses of multiple sympatrically breeding species enables us to refine predictions about the influence of climate change within communities.

Methods

Study site and breeding activity. Data on timing of breeding for our cavity-nesting community were collected at Riske Creek and Knife Creek near Williams Lake in south-central British Columbia, Canada (51°52′N, 122°21′W) from 1995–2011. The Riske Creek site is composed of trembling aspen (Populus tremuloides) within a grassland-wetland matrix. Knife Creek is largely mixed conifer with some deciduous riparian zones. Further detail about these sites can be found in Martin et al.36. Thirty-one cavity-nesting bird species are found in the community36; we restricted our analyses to the 8 resident species for which we found the highest abundance of nests (Table 1).

Active cavities were identified during systematic nest searches of the field sites that began in early May and ran until the end of July. Nesting attempts were initially identified based on adult behaviour (excavating, carrying food or entering/exiting cavities) or the vocalizations of young36. We recorded nest activity periods as the first and last days that a cavity was observed being used (containing nesting material, eggs, or nestlings). Between 1995 and 2004, active cavities within 5.2 m of the ground were accessed using ladders and mirrors to assess the stage of breeding. After 2004, a video camera mounted on a pole (TreeTop Peeper; Sandpiper Technologies, Manteca, CA, USA) was used to identify the stage of breeding in cavities up to 15 m above the ground35. The mean difference in cavity height between the period prior to the use of a pole-peeper and the period afterwards was 0.75 m. Some of this shift was due to changing excavator abundance over the monitoring period as higher cavities (>6 m above ground) produced by red-breasted sapsucker and downy woodpecker became more abundant while northern flicker cavities (an average of 3.5 m above ground) decreased in availability36,38. We do not consider this shift in cavity height to be a source of bias for our phenology data.

Nests were checked, on average, every 3–5 days during their active phase. Clutch initiation and hatch dates were determined for accessible nests using the observed final clutch size and stages of chick development combined with species-specific laying interval, duration of incubation, and nesting period. Nesting attempts found after the young had hatched could not be assigned a clutch initiation date in the field. Similarly, inaccessible cavities were recorded as active based on adult behaviour or begging chicks but could not be assigned a clutch initiation date or hatch date in the field. When not directly observed, fledging was considered to have occurred if chicks were old enough at the last nest check to survive out of the nest and if there was no evidence of predation within or around the cavity. The presence of adults foraging and feeding chicks in the immediate area was also used as an indicator of successful fledging when the event was not observed directly. All fieldwork was carried out in accordance with relevant guidelines and regulations.

Phenology. Of 1628 nesting attempts for which phenology data were recorded in the field, 352 had clutch initiation date specifically recorded. We used these dates to inform a linear mixed-effects model predicting the timing of clutch initiation for the remaining monitored nesting attempts as follows. Attempts found at the nesting stage were backdated using actual clutch size (where known) or species mean clutch size at the study site along with species-specific lay-rates (eggs/day), plus the mean incubation period from the literature39. These dates were then recorded as dates when nests were at the egg stage. Clutch initiation date was then modelled as a function of: the date and stage (pre-lay, egg, or unknown) of the earliest known/backdated nest activity, the latest date the nest was observed active, nest outcome (fledged, failed pre-hatch, failed pre-fledge, or unknown) and the number of visits to the nest by field crews; species identity was included as a random factor (intercept). With our dataset of field-recorded initiation dates (n = 352) we used a 5-fold cross-validation approach with a 20% set-aside to test the predictive capability of the model and to calculate a root mean-squared prediction error (RMSPE) for predicted clutch initiation40. The average performance of the model was high (5-run mean: r = 0.97 n = 70.4) and the RMSPE was 3.40 days. Species-specific performance ranged from r = 0.92 to r = 0.98. Our model was then trained using the entire dataset of field-recorded initiation dates (n = 352) and used to predict initiation dates for the remaining nest attempts (n = 1276). Error within predicted dates was incorporated into the final analyses by multiply imputing these values using a Monte Carlo approach41,42; predicted dates were adjusted in each imputation using values obtained from random draws of a normal distribution with a mean of zero and a standard deviation of 3.40 days (the RMSPE calculated above).

We subsequently restricted our analyses to first nesting attempts of the season. Known second nesting attempts by the same breeding pair (noted in the field) were ignored and records were additionally limited to the first occupancy of each nest cavity in each year. This removed possible missed re-nests in the same cavity by the same breeding pair as well as late nesting attempts by other species following abandonment or displacement of the first nesting pair. This reduced the final dataset from 1628 to 1570 nesting attempts.

To calculate the mean (±95% CI) annual clutch initiation dates reported in Fig. 1, we imputed 500 datasets, adjusting predicted values using repeated random draws from our normal distribution with a mean of zero and a standard deviation of the RMSPE. We then calculated mean annual initiation dates and standard deviation for each dataset. Reported values are the average of these 500 imputations. We report mean annual clutch initiation date for each species as “early” or “late” relative to the multi-year average for that species (this average is indicated by the horizontal line in Fig. 1).

Local weather variables. Daily minimum, maximum and mean temperatures (°C) and rainfall data (mm) were obtained from the Environment and Climate Change Canada weather station Williams Lake A (WMO ID 71104; 52°10′48″N, 122°03′00″W; elevation 939.7 m; http://climate.weather.gc.ca). These showed a
strong correspondence with incomplete data from a BC Wildfire Service station at the Riske Creek study site (51°57′37″N, 122°30′00″W; elevation 929 m; Station 210) (Pearson’s r = 0.95, and 0.97 for minimum and mean temperature respectively). Monthly average maximum temperatures are shown in Fig. 4A. Precipitation between the two stations also corresponded, albeit less strongly (Pearson’s r = 0.80).

Ecological variables. Beyond weather, two ecological variables that could impose additional influences on timing of breeding at our field site showed annual variation (Fig. 4B). A mountain pine beetle (MPB) outbreak impacted our study area during the monitoring period, with peak larval availability occurring in 2000, and 2003–2004. This represented a major food pulse year-round for the majority of species in our study and resulted in the increased abundance of resident cavity-nesting species at our field site21,22,26,43. We considered MPB a possible direct driver of phenology (as opposed to the indirect effect of local temperatures on the beetle outbreak) and quantified the availability of this resource as the number of live, MPB infected pines/ha on the study site in each year (for further detail see43). The number of pre-existing cavities also changed over the study period. These cavities were used by secondary cavity-nesters and facultative excavators: mountain chickadee, red-breasted nuthatch, and northern saw-whet owl. We quantified cavity availability for these species at the beginning of the breeding period as the number of cavities in each year that were not newly excavated. We tested small and medium-sized cavity (<5 cm diameter) availability for mountain chickadee and nuthatch and large cavity availability (≥5 cm, <20 cm diameter) for the owl: all types of cavities were least numerous in 1995, and most numerous in 200526.

Analysis. We tested the degree of co-variation in annual mean clutch initiation dates across our community using pair-wise correlation analyses. Comparisons were restricted to years where species had multiple nests monitored and we corrected for multiple comparisons using Benjamini and Hochberg’s54 approach to limit false discoveries. Sample sizes varied from 10 years to 17 years (Table 2).

We used a linear mixed-effects modelling approach to describe individual clutch initiation date for each species as a function of our explanatory weather and ecological variables. As above, only years where a species had multiple nests monitored were included. ‘Year’ was included as a random factor to account for the non-independence of data collected within the same year (‘Ime4’). Plausible models were competed to determine which of our candidate variables best explained phenology within our community. To limit the final model candidate set, we used 2-step hierarchical approach to model testing. The initial analyses examined the performance of our weather variables using a sliding window approach ('climwin'43). Average maximum, minimum and mean temperatures and average total daily rainfall were calculated for a moving window between November 1 of the previous year and June 1 of the breeding year. Minimum window size was restricted to 10 days (providing time for a physiological reproductive response to conditions) and the maximum encompassed the entire time period24,23. The best time window for each variable was determined using AIC. The predictive performance of all time periods examined, along with the best single window (boxed), is presented in Fig. 3 using AIC model weights.

Given the large number of time widows examined, we calculated the probability that the AIC scores we obtained could have been achieved by chance (i.e. Type I error) by using a response-data randomization program included for this purpose within ‘climwin’. Specifically, we considered our results to be spurious when our model AIC values did not differ significantly (P > 0.05) from those generated from sliding window analyses of 100 randomizations of the response data (for details see43). All weather variables that passed the ‘Type I error’ test and that fell within ΔAICc ≤ 2 of the top performing variable were subsequently run against a null (random effect of ‘Year’ only) and linear temporal model, models incorporating our ecological variables, and, finally, biologically plausible additive models that included weather and ecological factors together. For model testing, we imputed 100 datasets using repeated random draws (see Phenology, above) and bootstrapped each of these (n = 20 per dataset) to obtain our model estimates25. Reported coefficients are the medians, and reported 95% confidence intervals are the middle 95% range (or 0.025 and 0.975 quantiles), of these 2000 model runs (Table 3).

The performance of the "best" final model for each species is presented in two ways. First, we used the function sem.model.fits ('piecewiseSEM'43) on models containing ‘Year’ as a random intercept and fixed effects whose beta values did not cross zero within their 95% CI. Marginal and conditional R² values are the median values obtained from 2000 imputation-bootstrap runs, as above (Table 3). To more intuitively present the amount of among-year variation in timing explained by fixed effects in our models we additionally calculated an R² value for predicted versus observed mean annual clutch initiation dates. These values reflect the fit of regression lines presented in Fig. 2. All analyses were run in R (version 3.2.1, R Foundation for Statistical Computing 2015).

Data availability. All datasets used in this study are readily available from the corresponding author upon request.

References
1. Carey, C. The impacts of climate change on the annual cycles of birds. Philos. Trans. R. Soc. B 364, 3321–3330 (2009).
2. Lane, J. E., Kruuk, L. E. B., Charmantier, A., Murie, J. O. & Dobson, F. S. Delayed phenology and reduced fitness associated with climate change in a wild hibernator. Nature 489, 554–557 (2012).
3. Anderson, J. T., Inouye, D. W., McKinney, A. M., Colautti, R. I. & Mitchell-Olds, T. Phenotypic plasticity and adaptive evolution contribute to advancing flowering phenology in response to climate change. Proc. R. Soc. Lond. B Biol. Sci. https://doi.org/10.1098/rspb.2012.1051 (2012).
4. Gilbert, N. & Raworth, D. A. Insects and temperature: A general theory. Can. Entomol. 128, 1–13 (1996).
5. Stevenson, I. R. & Bryant, D. M. Avian phenology: Climate change and constraints on breeding. Nature 406, 366–367 (2000).
6. Moyes, K. et al. Advancing breeding phenology in response to environmental change in a wild red deer population. Glob. Change Biol. 17, 2455–2469 (2011).
47. Lefcheck, J. S. piecewiseSEM: Piecewise structural equation modelling in R for ecology, evolution, and systematics.

45. van de Pol, M.

44. Benjamini, Y. & Hochberg, Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing.

41. Schafer, J. L. & Graham, J. W. Missing data: Our view of the state of the art.

39. The Birds of North America Online (A. Poole Ed.) Ithaca: Cornell Lab of Ornithology. Available at: http://bna.birds.cornell.edu/bna/.

38. Cockle, K. L. & Martin, K. Temporal dynamics of a commensal network of cavity-nesting vertebrates: increased diversity during an insect outbreak.

37. Edworthy, A. B., Wiebe, K. L. & Martin, K. Survival analysis of a critical resource for cavity-nesting communities: patterns of tree cavity longevity.

35. Visser, M. E., Both, C. & Lambrechts, M. M. Global climate change leads to mistimed avian reproduction.

34. Crick, H. Q. P. & Sparks, T. Climate change related to egg-laying trends in migrant birds.

33. Vose, R. S., Easterling, D. R. & Gleason, B. Maximum and minimum temperature trends for the globe: An update through 2004.

32. Newton, A. R. & Martin, K. The perils of plasticity: dual resource pulses increase facilitation but destabilize populations of small-bodied cavity-nesters.

31. Martin, K., Norris, A. & Drever, M. Effects of bark beetle outbreaks on avian biodiversity in the British Columbia interior: implications for critical habitat management.

30. Williams, T. D. et al. Mid-winter temperatures, not spring temperatures, predict breeding phenology in the European starling Sturnus vulgaris. Open Sci. 2, 140301 (2015).

29. Norris, A. R. & Martin, K. Red-breasted nuthatches (Sitta canadensis) increase cavity excavation in response to a mountain pine beetle outbreak.

28. Williams, T. D.

27. Norris, A. R. & Martin, K. The perils of plasticity: dual resource pulses increase facilitation but destabilize populations of small-bodied cavity-nesters. Oikos 119, 1126–1135 (2010).

26. Martin, K., Norris, A. & Drever, M. Effects of bark beetle outbreaks on avian biodiversity in the British Columbia interior: implications for critical habitat management. J. Ecosyst. Manag. 7, 201–211 (2011).

25. Edworthy, A. B., Drever, M. C. & Martin, K. Woodpeckers increase cavity excavation but maintain fecundity in response to an outbreak of mountain pine bark beetles. For. Ecol. Manag. 261, 209–217 (2007).

24. Williams, T. D.

23. Williams, T. D. et al. Mid-winter temperatures, not spring temperatures, predict breeding phenology in the European starling Sturnus vulgaris. Open Sci. 2, 140301 (2015).

22. Wiebe, K. L., Koenig, W. D. & Martin, K. Costs and benefits of nest reuse versus excavation in cavity-nesting birds. Ann. Zool. Fenn. 44, 209–217 (2007).

21. Steward, J. S., Round, P. D. & Milne, J. R. Food availability fails to explain asynchronous breeding of two syntopic oriental trogons.

20. Smith, R. J. & Moore, F. R. Arrival timing and seasonal reproductive performance in a long-distance migratory landbird. Behav. Ecol. Sociobiol. 57, 231–239 (2005).

19. Brown, W. P. & Roth, R. R. Temporal patterns of fitness and survival in the wood thrush.

18. Smith, R. J. & Moore, F. R. Arrival timing and seasonal reproductive performance in a long-distance migratory landbird. Behav. Ecol. Sociobiol. 57, 231–239 (2005).

17. Brown, W. P. & Roth, R. R. Temporal patterns of fitness and survival in the wood thrush.

16. Rowe, L., Ludwig, D. & Schluter, D. Time, condition, and the seasonal decline of avian clutch size. Am. Nat. 143, 698–722 (1994).

15. van de Pol, M.

14. Visser, M., Holleman, L. & Gienapp, P. Shifts in caterpillar biomass phenology due to climate change and its impact on the breeding biology of an insectivorous bird. Oecologia 147, 164–172 (2006).

13. Siikamäki, P. Limitation of reproductive success by food availability and breeding time in pied flycatchers.

12. Visser, M. E.

11. Drake, A., Martin, M. & Green, D. J. Winter habitat use does not influence spring arrival dates or the reproductive success of yellow warblers breeding in the arctic. Polar Biol. 37, 181–191 (2014).

10. Townsend, A. K. et al. Warm springs, early lay dates, and double brooding in a North American migratory songbird, the black-throated blue warbler. PLOS ONE 8, e59467 (2013).

9. Rowe, L., Ludwig, D. & Schluter, D. Time, condition, and the seasonal decline of avian clutch size. Am. Nat. 143, 698–722 (1994).

8. Wilson, S. & Arcese, P. El Niño drives timing of breeding but not population growth in the song sparrow (Melospiza melodia). Proc. Natl. Acad. Sci. USA 100, 11139–11142 (2003).

7. Dunn, P. O. & Winkler, D. W. Climate change has affected the breeding date of tree swallows throughout North America. Proc. R. Soc. Lond. B Biol. Sci. 266, 2487–2490 (1999).

6. Row, L., Ludwig, D. & Schluter, D. Time, condition, and the seasonal decline of avian clutch size. Am. Nat. 143, 698–722 (1994).

5. van de Pol, M. et al. Identifying the best climatic predictors in ecology and evolution. Methods Ecol. Evol. 7, 1246–1257 (2016).

4. Scher, J. L. & Graham, J. W. Missing data: Our view of the state of the art. Psychol. Methods 7, 147–177 (2002).

3. Schomaker, M. & Heumann, C. Model selection and model averaging after multiple imputation. Comput. Stat. Data Anal. 71, 758–770 (2014).

2. Drever, M. C., Goheen, J. R. & Martin, K. Species-energy theory, pulsed resources, and regulation of avian richness during a mountain pine beetle outbreak. Ecology 90, 1095–1105 (2009).

1. Benjamini, Y. & Hochberg, Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. Ser. B Methodol. 57, 289–300 (1995).
Author Contributions
K.M. conceived and designed the fieldwork. A.D. conducted the analyses and wrote the manuscript. K.M. reviewed and edited the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018