- Inconsistent shifts in warming and temperature variability negatively impact avian fitness.
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9 ABSTRACT

- As the climate has warmed many birds have advanced their breeding timing. However, climate change also changes temperature distributions; thus breeding earlier in a hotter world might 11 increase nestling exposure to either extreme heat or cold. We combined >300,000 breeding 12 records from 24 North American birds with historical temperature data to understand 13 how exposure to extreme temperatures has changed. Average spring temperature increased 14 since 1950, but change in timing of extremes was inconsistent in direction and magnitude; 15 thus, populations could not track both average and extreme temperatures. Relative fitness 16 was reduced following heat-waves and cold-snaps in 11 and 16 of 24 species, respectively. 17 Latitudinal variation in sensitivity in three widespread species suggests that vulnerability to extremes at range limits may contribute to range shifts. Our results add to evidence 19 demonstrating that understanding individual sensitivity and its links to population level processes is critical for predicting vulnerability to changing climates.
- 22 Keywords: climate change, cold-snaps, heat-waves, reproductive success, phenology, extreme
 23 temperature. community science

1 INTRODUCTION

- Human induced climate change has resulted in an accelerating increase in the average
 temperatures that animals experience around the globe. A rich literature now documents
 the consequences that span from the scale of individuals to entire populations and species.
 The most universal observed responses to warming are reductions in body size, shifts in
 species ranges, and changes in the timing of annual events (i.e., phenological shifts), which have the potential to impact fitness via increased interspecific competition or
- phenological mismatches between predators and prey. 10-13 However, mean temperatures
- alone fail to capture the complexity of climate change, as the rate and magnitude of warming

- often differs between day and night as well as seasonally. Another component of climate
- change is increasing temperature variability, which is predicted to have greater impacts on
- species survival than increases in mean temperature. ¹⁴ As the pace, magnitude, and
- variability of climate change determines the risk of exposure to conditions outside those
- 37 historically experienced, there is a critical need to understand how and why the magnitude
- of these consequences differs between species to predict and mitigate the effects of climate
- 39 change for wild animals. 15
- ⁴⁰ Recently, Rosenberg et al. ¹⁶ estimated that the total population of North American birds has
- declined by almost 3 billion since 1970. While these declines arise from a combination of
- several factors (e.g., introduced predators; ¹⁷ noise and light pollution; ¹⁸ land use change ¹⁹;
- disease²⁰), climate change is generally considered to be one of the primary threats and
- drivers. 21,22 However, the declines in overall abundance are far from uniform, with some
- 45 groups, such as aerial insectivores, showing steep declines while others, such as many
- wetland birds, have increased in numbers. 16 These differences suggest that life history and
- 47 habitat requirements many play an important role in determining the sensitivity of different
- 48 species to climate change.
- 49 One of the major goals of many climate studies has been to assess whether wild birds are
- 50 shifting their ranges or advancing breeding phenology fast enough to track changes in mean
- temperatures^{23–25}, with several recent studies suggesting that observed changes are generally
- inadequate to keep pace. 26-28 However, it is also increasingly apparent that changes in mean
- temperatures are insufficient to understand the response to climate change, as the mean only
- 54 provides a coarse approximation of changes in minimum (nighttime) and maximum
- 55 (daytime) temperatures which can change independently in pace, magnitude, and variability.
- 56 Different organismal traits such as timing of reproduction or when a species is active likely
- 57 shape the response because animals may need to balance changing temperatures
- simultaneously with other abiotic gradients, such as day length, UV radiation, or water

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availability.<sup>29,30</sup> Even when only considering mean temperatures, perfectly tracking average
   daily temperatures may result in a more variable thermal environment when breeding
   activities are occurring earlier in the year<sup>31</sup> or when climate warming is associated with
   increased climate variability.<sup>32–34</sup> When predictable conditions are linked to survival,
   increases in either the magnitude or the frequency of conditions outside historical norms can
   result in reduced fitness and declines at the population or species level.
   The relative importance of different abiotic gradients for each species and their link to
   fitness, coupled with the pace of plasticity or evolutionary change, likely dictates which
   species persist and which will perish.<sup>22</sup> The strength of this relationship is determined by
   how efficiently organisms transfer heat and energy in different environmental conditions and
   how these traits influence individual performance as well as the ability to survive, develop,
   grow, and reproduce.<sup>35</sup> For example, several studies demonstrate that life history traits
   predict different aspects of the response to climate change. 19,36,37 Among birds, foraging
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   mode is one key life history characteristic that might influence the relative challenge posed
   by average temperature versus variation in temperature. The vast majority of birds are
   reliant on insects to provide energy and nutrients to their rapidly developing offspring. For
   example, obligate aerial insectivores depend on flying insects, even modest decreases in
   daytime temperatures can reduce insect activity and thereby overall foraging success^{38}
   (hereafter "cold-snaps"). If these reductions in insect activity occur during critical periods
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   when nestlings are most vulnerable, they can trigger mass mortality events.<sup>31</sup> Accordingly,
   species that do not rely on insect activity while foraging are likely to be less sensitive to
   cold-snaps. In contrast, heat-waves would be expected to impact a broader range of species
   as they may approach physiological upper temperature limits. 39,40 A more thorough
   understanding of relative sensitivity to average temperature, cold-snaps, and heat-waves
   during breeding attempts requires data that spans multiple species across a wide range of
   conditions paired with detailed records of reproductive performance.
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The most detailed studies of climate change on populations tend to focus on single species with a long time series of detailed breeding data.^{3,31,32,41} These studies provide evidence for the mechanisms by which climate change influences populations, but they typically do not address variation in responses between species or different populations. Another approach is to use large scale observations to model spatial, temporal, and cross species variation in 89 population phenology, 7,8 abundance, 16 body size, 4,42 or ranges 43,44 in response to climate change. However, these studies typically cannot link the patterns observed at coarse scales to specific processes and mechanisms that occur at the level of individual animals.²⁵ Linking individual fitness to population level processes with large scale and spatially variable outcomes is needed to accurately predict variation in sensitivity to climate change across species. 95 In this study, we used a database of community-contributed observations on the timing of breeding and reproductive success in wild North American birds compiled by Project 97 MartinWatch, by the NestWatch program at the Cornell Lab of Ornithology, and by Project Nestwatch from Birds Canada between 1995 and 2020. We identified >300,000 breeding records from 24 common species with nesting records spanning most of the United States 100 and Canada. By documenting the reproductive performance of individuals, this database 101 allows for detailed exploration of spatial, temporal, and interspecific variation in temperature 102 exposure and vulnerability at near continental scales. 103 We first asked how the change in timing of cold-snaps and heat-waves compares with changes 104 in overall average spring temperature across the range of locations from which breeding data 105 were available. If these abiotic gradients have advanced at different paces or in different 106 directions, then populations will be unable to track the same breeding conditions regardless 107 of phenological advancement. Next, we determined the temperatures experienced during 108 every nesting attempt and asked whether each species showed evidence of performance declines during cold-snaps or heat-waves during the breeding attempt (i.e., whether these 110

events qualify as extreme climatic events 'ECEs' with biological consequences^{45,46}). We
predicted that performance declines would be most pronounced for cold-snaps experienced
by aerial insectivores because of the direct impact on flying insect availability. In contrast,
we predicted that all species in the dataset would be vulnerable to heat-waves.

We combined breeding records from three different databases. First, we requested raw

5 2 MATERIALS & METHODS

¹⁶ 2.1 Breeding records

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breeding records from Nestwatch, a project managed by the Cornell Lab of Ornithology 118 (www.nestwatch.org). Second, we requested raw breeding records from Project Nestwatch, a 119 program managed by Birds Canada (www.birdscanada.org/bird-science/project-nestwatch/). 120 Finally, we obtained publicly archived records of breeding purple martins (*Progne subis*) 121 from Project MartinWatch (www.purplemartin.org⁴⁷). We initially manipulated each data source into a common format to allow us to merge records together. Beginning with the 123 combined dataset, we then proceeded through a series of filtering and quality control steps to 124 arrive at a final dataset for analysis. 125 While the Nestwatch database includes some older observations, most nest records are from 1995 or later (>98%); we therefore removed older records to focus on the period from 1995 to 2020. We also removed species that had fewer than 300 records. This minimum sample size was somewhat arbitrary, but given the complexity of models that we planned to fit, the 129 desire for spatial and temporal sampling, and the need to model ECEs which, by definition, 130 only occur in a small percentage of nesting attempts, we chose a 300 nest minimum to ensure 131 that we could reliably fit models for each species included. For a species with exactly 300 132 nests, we would only expect to observe an average of 7.5 nestling attempts that experienced 133 a 2 SD cold-snap or heat-wave. We also removed European starling (Sturnus vulgaris) and

house sparrow (Passer domesticus) records from the dataset, because these two invasive species are often considered nuisance pests and it was unclear whether a high reporting rate 136 of failed nests reflected human eviction from nest boxes. This resulted in 24 remaining 137 species that we considered further (Table S1). 138 Next, we removed any records that had incomplete location data, that were outside of the 139 study extent, were flagged by the data providers as having errors, or that had impossible values reported (e.g., more young than eggs or negative clutch sizes). We also removed nests 141 that did not include information on the date of laying or hatching. In cases where only a laying date was provided, we inferred approximate hatching dates for successful nests by 143 using the typical incubation length as reported in species accounts from the Birds of the 144 World online. 48 While fledging success was reported, exact fledging date was generally 145 unknown and we estimated fledging dates using the typical fledging age for a species from 146 Birds of the World online.⁴⁸ 147 At this point, we also roughly categorized the included species into four foraging modes 148 based on the primary way that food is obtained during the nestling provisioning period using 149 the species accounts in Birds of the World Online. These included aerial capture (both 150 fly-catching and aerial insectivores), sally foraging (flying to ground from a perch after 151 visually detecting arthropod movement), gleaning (a broad category of strategies involving 152 active search for arthropods), and seed eating. We did not have enough total species or 153 phylogenetic variation in foraging mode to formally analyze diet, but we discuss the possible 154 role that these foraging styles may have had on susceptibility to cold-snaps and heat-waves. 155 Finally, we plotted distributions of clutch sizes and hatching dates for each of the 24 species 156 and removed records that had values that were likely due to data entry error (i.e., outside of 157 the possible range for each species). This filtering step was somewhat subjective, but we 158 were conservative in setting limits so that only records well outside the expected ranges were 159 excluded. The end result of these filtering steps was a final dataset that included 301,514 160

breeding records from 24 species. The spatial extent of the dataset is illustrated in Figure S1 and the number of nests included for each species is shown in Table S1. Using the final set of filtered breeding records, we determined the elevation of each nest by accessing a digital elevation model raster spanning the records using the elevatr package in R.⁴⁹ Temperature data were added to each nest record as described below.

56 2.2 Spatial and temperature data

Spatial data. We downloaded a base map for the region encompassing the breeding records 167 included using the package rnaturalearth⁵⁰ in R version 4.0.2.⁵¹ For some analyses (see 168 below) we wanted to summarize breeding records and historical temperature by grouping 169 records and weather stations that were recorded close together. To facilitate this grouping, 170 we created a grid of hexagons each with an area of 40,000 km² across the land area of North 171 America with the st make grid function from package sf.⁵² We then clipped this grid to 172 include only hexagons that contained breeding records from the dataset described above. 173 Temperature records. We accessed historical temperature data from meteorological 174 stations in the United States using the package rnoaa⁵³ and in Canada using the package 175 weathercan.⁵⁴ In both cases, we filtered stations to include those that were reported to have 176 at least 50 years of data between 1920 and 2020 (not all stations actually yielded 50 years of 177 data because some were active but missing the data we required). Using these criteria, we 178 identified 2,608 stations from the National Oceanic and Atmospheric Administration 179 (NOAA) and 1,125 stations from Environment and Climate Change Canada (ECCC) that 180 covered the spatial extent of our breeding records (Figure S2). From these stations we 181 downloaded all available daily high temperature records between 1920 and 2020. When 182 summarizing temperature data for the hexagonal grid, we averaged all stations that were 183 within the bounds of each hexagonal grid cell. 184

Temperature anomalies. We downloaded a monthly temperature anomaly raster from

the Berkeley Earth project (www.berkeleyearth.org/data/). This data product compares the monthly average temperature within each 1° square grid to the average temperature over the 187 period 1951-1980. Monthly anomalies in each year are expressed relative to that 30 year 188 baseline period. Because we were interested in conditions during the breeding season, we 189 extracted the anomaly for April, May, and June for each year from 1920-2020. We next 190 clipped the square anomaly grid to the boundaries of each equal area hexagon and calculated 191 a weighted average representing the anomaly within each hexagon and then merged these 192 readings with the breeding records described above. We hereafter refer to this averaged value 193 as the spring temperature anomaly. 194 **Definitions of extreme climate events.** In order to usefully summarize and analyze 195 long-term climatic data, we had to make decisions about what counts as an extreme event. 196 These decisions included choices about how to handle the severity, duration, and timing of 197 cold-snaps and heat-waves. We followed two connected, but distinct strategies for the 198 historical analysis of the timing of extreme events and for the analysis of the biological 190 effects of extreme events on each species. For historical analyses, we needed to select 200 absolute temperature values and we defined these based on the distribution of temperatures 201 actually experienced by the 24 species we studied (see details below). 202 For the analyses focused on fitness effects for individual species, we were interested in the 203 evidence for ECEs resulting from extreme temperatures. The definition of an ECE is 204 inconsistent, 45,46 but typically these events refer to a combination of both extreme climatic 205 conditions defined based on a distribution (e.g., events occurring <5% of the time) and a 206 negative biological response, sometimes requiring a long recovery period. 45 No single 207

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definition is universally applicable across studies⁴⁶, but for the purposes of our study, we

considered our results to indicate evidence for an ECE if 3-day cold or heat events two

standard deviations from the species mean were associated with reduced reproductive

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success for each species.

The exact choice of a 3-day time period for considering cold-snaps and heat-waves was also somewhat arbitrary, but previous work suggests that multi-day poor weather events tend to have a greater impact on offspring survival than single day events. This pattern has been observed in raptors, ^{55–57} in swallows, ^{38,58} and in other passerines. ³⁴ We acknowledge that longer cold-snaps and heat-waves likely have even more severe impacts, but they will also be correspondingly rarer to observe when matched with breeding records and 3 days represents a compromise between a multi-day challenge and enough observation of extreme temperatures to allow us to model fitness effects.

Categorizing patterns of historical cold-snap and heat-wave occurrence. Using 220 the averaged daily high temperature for each grid cell, we determined the date of the latest 221 3-day cold-snap and the date of the earliest 3-day heat-wave for each year and hexagon cell 222 between 1920 and 2020. For this analysis, it was necessary to use absolute temperature 223 values to define a cold-snap and heat-wave so that we could ask how the annual timing of the 224 same temperature conditions may have changed over the past 100 years. We selected the 225 threshold temperatures used by examining the distribution of temperatures recorded during 226 nest attempts for the 24 species included in our study. 227

For each species, we determined the 5th, 10th, and 20th percentile of 3-day temperatures (for cold-snaps) along with the 80th, 90th, and 95th percentile of 3-day temperatures (for heat-waves) experienced during all the nesting attempts in our cleaned dataset (see below for details on nest level temperature). This resulted in 24 temperatures for each percentile value.

We then averaged these species-specific estimates to arrive at a single threshold value for each percentile (Figure S3).

This summary resulted in cold-snap thresholds of 13.9°, 15.5°, and 17.4° C along with
heat-wave thresholds of 32.1°, 33.6°, and 34.8° C. In each year, we only considered
cold-snaps and heat-waves that occurred after the 60th and before the 240th day of the year
(approximately March 1st to August 28th), because we were interested in the timing of these

events in relation to breeding activities. We present all three thresholds in most analyses and in all cases the patterns were qualitatively similar regardless of threshold, but for some summary plots at the continental scale we used the milder thresholds (17.4° and 32.1°) 240 because these allowed for the inclusion of a wider geographic area since the most extreme 241 temperatures for cold and heat were rarely recorded at low and high latitudes, respectively. 242 Historical trends in cold-snap and heat-wave anomalies. We derived a measure of yearly cold-snap and heat-wave timing anomaly for each grid cell to compare with the average temperature anomaly data described above. To do this we averaged the last 245 cold-snap and earliest heat-wave date for the years 1951-1980 in each grid and then for each 246 year and grid combination calculated the deviation from that average date. Negative values 247 for the anomaly indicate years in which the last cold-snap or earliest heat-wave date 248 occurred earlier than the historical average and positive values indicate years in which the 249 last cold-snap or earliest heat-wave occurred later than the historical average. These values 250 were calculated to be comparable to the temperature anomaly data described above. 251 Temperatures experienced by individuals during breeding. Using the temperature 252 data described above, we matched each breeding record to temperature from the most 253 similar station using a two step process. We first matched records to the closest station. We 254 next compared the difference in elevation between the breeding record and the station. If the 255 closest station differed in elevation by >300 m, we searched for the station at the most 256 similar elevation within 50 km. Using this approach, we paired records to weather stations 257 that were 18.6 ± 10.5 km (SD) from the nest and within 47.2 ± 67.1 m elevation. While this 258 approach resulted in the best matched station, we also note that standardized temperature 250 records are usually recorded ~1.5m above the ground, whereas the species included in our 260 dataset could have nested at varying heights and in open cups or nest boxes. Thus, the exact 261 thermal environment for nests likely differed from the temperature records we used, but we could not account for this detailed level of variation. 263

Using the identified station, we determined the coldest and hottest 3-day period experienced during each nesting attempt separately for the incubation and nestling provisioning period. 265 To accomplish this, we found the sequence of 3 days with the lowest combined daytime high temperature and considered this value as the lowest short term temperature exposure. For 267 the highest temperature, we first determined the high temperature for each day in a string of 268 3-days; for each group of 3 days, we considered the lowest daytime high temperature and we 269 then selected the string of 3-days that maximized this value (i.e., the hottest 3-day period 270 experienced). It is important to note that these temperature determinations were continuous 271 and, unlike the historical analysis above, they did not rely on any choice of threshold values. 272 We used species specific timing for incubation and nestling stages to ensure that the 273 temperatures we recorded would actually have been experienced during the reproductive 274 attempt. 275

2.6 2.3 Data analysis

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100 years in spring temperature anomalies and in the timing of cold-snaps and heat-waves 278 across the spatial range of the breeding records included in our dataset. We were interested 279 in determining how these temperature variables have changed over time, how much regional 280 variation there is in those patterns, and the extent to which average temperature anomalies were correlated with cold-snap and heat-wave timing. We initially plotted the average temperature, cold-snap, and heat-wave anomalies over the 283 past 25 years (the time period covering our breeding records) in each of the hexagon grids 284 described above for illustration purposes. We also fit a generalized additive model (GAM) 285 for each anomaly measure using the entire time series with the anomaly in each grid-year combination as the response variable along with a basis smoothed predictor variable for year and a spatial smooth for latitude and longitude to account for spatial autocorrelation. We

Spring temperature and cold-snap timing. We first modeled the change over the last

used these models to describe the overall change in spring temperatures and cold-snap or heat-wave timings and the degree of spatial variation in those changes.

Next, we fit a GAM with the average timing of last cold-snap or earliest heat-wave anomaly for each grid over the past 25 years as the response variable and with the corresponding average spring temperature anomaly, and a spatial smooth for latitude and longitude as predictor variables. These models were used to infer whether spring temperature anomaly and the timing of the latest cold-snap or earliest heat-wave covaried, while accounting for spatial autocorrelation in the dataset.

Impact of cold-snaps and heat-waves on reproductive success. To model the
impact of cold-snaps and heat-waves on reproductive success, we used a two-step approach
to fit GAMs for each species separately for the incubation and nestling provisioning phase.
First, as weather measurements were indicative of fine-scale regional conditions rather than
those measured at specific nest locations, we calculated the response to weather conditions
by averaging the number of chicks fledged grouped by hatch date and nearest weather
station. We then created a model using these averaged records and 3-day temperature
extremes experienced during the incubation period with mean number of chicks fledged as
the response variable.

Next, we fit the same model for each species, but restricted records to those that hatched at least a single chick successfully and included only temperature records during the nestling 307 provisioning stage. This allowed us to minimize the direct effects of temperature extremes 308 during incubation on our models investigating the nestling phase. We separated between 309 these two life history stages because we suspected that temperature extremes experienced 310 during the nestling provisioning phase could have a stronger direct effect on fledging success, 311 as the ability of the parent to buffer against extreme temperatures varies.³¹ We implemented 312 a Gaussian model for both the incubation and nestling provisioning models as they have 313 lower false positive Type I error rates when compared to a Poisson due to overdispersion,

and thus tend to be more conservative.⁵⁹

In both models, the structure of the GAM was identical. Predictors included a basis smooth 316 for the coldest and hottest 3-day period during either reproductive stage (incubation stage or provisioning) as well as a spatial smooth for latitude and longitude to account for spatial 318 autocorrelation. The models also included smoothed predictors for date and a random effect for year. The basis dimension value of k for the spatial smooth was chosen iteratively by comparing the effective degrees of freedom (edf) with the k-index as per Wood 2017⁶⁰. 321 Predictors were checked for whether their smooths contributed unique information to the 322 model (concurvity). We fit each model separately for each of the 24 species included in our 323 analyses. Not all species had records in every year, so the number of years included in each 324 model varies by species, and the basis dimension value was set to this in each model. To 325 facilitate comparison between species, we standardized the coldest and hottest 3-day period 326 within species so that the mean was zero and standard deviation was one for each species. 327 We summarized sensitivity to cold-snaps and heat-waves from these models by calculating 328 pointwise estimates for number of offspring for each species during a 2 standard deviation 329 cold-snap or heat-wave that occurred during either incubation or the nestling provisioning 330 phase while controlling for the other model parameters. We then converted the estimates to 331 model predicted relative fitness by taking the predicted number of offspring fledged divided 332 by the average number of offspring fledged for each species. We considered point estimates 333 with confidence intervals that did not overlap one (the average value of relative fitness) to 334 indicate significant sensitivity to temperature extremes. 335 Species that experienced these reductions in fitness would be considered to have experienced 336 an ECE based on the combined climatic extremity and biological response definition outlined 337 by Smith, 2011. 45 However, Bailey & van de Pol (2016) argue that using an arbitrary 338 distribution cutoff might miss important biological responses at different values or when 339 responses differ between species⁴⁶ (in our case using 2 SD results in point estimates at

approximately the 2.5th and 97.5th percentiles). In an online appendix, we include the full relative fitness surface for each species across the range of coldest and hottest 3-day periods.

These comparisons do not rely on any arbitrary choice of a single point estimate from the climatic distribution.

Latitudinal variation in cold-snap and heat-wave susceptibility.

After fitting the global models described above, we investigated whether the susceptibility to 346 cold-snaps and heat-waves varied from the northern to southern limits of the breeding range in three species: purple martins (*Progne subis*), eastern bluebirds (*Sialia sialis*), and tree swallows (Tachycineta bicolor). We chose these three species because they had the largest sample sizes and because all three also have a wide latitudinal distribution and the global analysis above indicated susceptibility to both cold-snaps and heat-waves. For each species, 351 we split the records into 5 latitude bands with an equal number of nest records per band and 352 then repeated the global analyses exactly as described above within each band. The number 353 of records was sufficient for these species that even after splitting into five datasets each 354 latitude band included >9,000 nesting records for each species. For tree swallows, we limited 355 this analysis to records east of the Rocky Mountains because mountain and western 356 populations have very different breeding timing from eastern populations at similar latitudes. 357

$_{358}$ 2.4 Data and code availability

All data used in the analyses here are available from publicly accessible databases. Breeding records for individual nesting attempts can be accessed from each organization by filing a request and completing a data sharing agreement. Historical temperature data was accessed from the Berkeley Earth project (www.berkeleyearth.org/data/) or from meteorological station records maintained by NOAA in the United States and by ECCC in Canada.

All data processing, analyses, and figures were created in R version 4.0.2.⁵¹ A complete set of

annotated code to reproduce the full analysis, manuscript, and supplemental materials is
permanently archived on Zenodo (DOI: 10.5281/zenodo.8208669). The archived repository
also includes an appendix with complete model summary tables for every GAM described in
the results. Figure S4 provides a conceptual overview of the complete analysis pipeline with
reference to each script in the code repository.

3 RESULTS

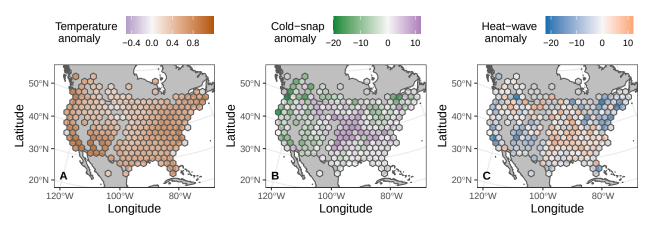


Figure 1: Spring temperature anomaly (A), cold-snap anomaly (B), and heat-wave anomaly (C) for the North American spatial extent studied over the years 1995-2020. Spring temperature anomaly is defined as the average of April, May, and June using data from the Berkeley Earth Project measured in degrees C compared to the baseline period of 1950-1980. Cold-snap anomaly is the difference in average timing of the latest three day period in which temperatures did not rise above 17.4 C from 1995-2020 compared to the average date of the latest cold-snap from 1950-1980, measured in days. Heat-wave anomaly is the difference in timing of the earliest three day period in which temperatures always rose above 32.1 C from 1995-2020 compared to the average date of the earliest heat-wave from 1950-1980, measured in days. Gray cells indicate regions with missing data.

$_{\scriptscriptstyle{71}}$ 3.1 Spring temperature and cold-snap or heat-wave timing

Across the spatial range that we studied, the average spring temperature anomaly from 1995 to 2020 was universally positive, although there was spatial variation in the magnitude of this increase in average spring temperature (Figure 1A). In contrast, change in the date of the latest 3-day cold-snap or the earliest 3-day heat-wave was inconsistent in both sign and magnitude (Figure 1B-C).

In a spatial GAM averaging across the entire range studied, spring temperature anomaly was
consistently positive from 1995-2020 and the last cold-snap tended to occur 3 to 5 days
earlier than the reference period regardless of the cold-snap threshold used (Figure 2A-B).
However, the first heat-wave was not consistently different from the reference period
regardless of the temperature threshold used (Figure 2C).

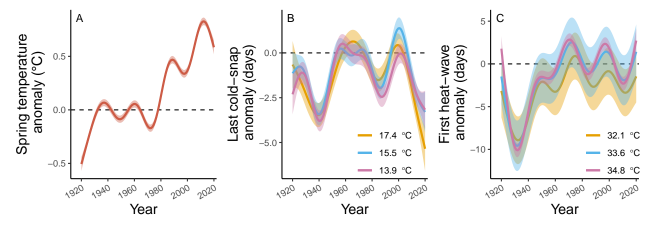


Figure 2: The historical change across the North American spatial extent studied in average spring temperature anomaly (A), timing of the last cold-snap (B), or timing of the first heat-wave (C) over the last 100 years. Lines and 95% confidence intervals are from spatial GAMs that account for latitude and longitude of each grid cell. For the cold-snap and heat-wave panels, three different threshold values are shown to illustrate increasingly more severe cold-snaps or heat-waves.

When considering average anomalies over the last 25 years, there was no evidence that a larger overall spring temperature anomaly was associated with any consistent difference in the change in cold-snap timing (Figure 3A) or heat-wave timing (Figure 3B). When comparing the temperature anomaly in each individual year at the level of hexagonal grids to the cold-snap or heat-wave anomaly in each individual grid-year, there was an association such that grid-years with higher spring temperature anomalies tended to have both earlier last cold-snaps and earlier heat-waves (Figure S5), but there was wide variation for individual grid-years.

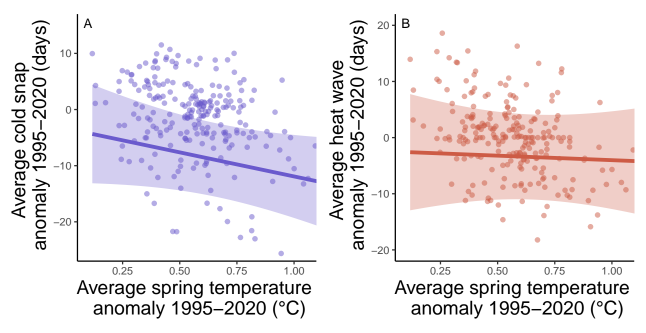


Figure 3: Relationship between average spring temperature anomaly for the North American spatial extent studied for each grid cell from 1995-2020 and (A) average cold-snap anomaly or (B) average heat-wave anomaly over the same period. Points are the average values for each hexagon grid cell. Lines and 95% confidence intervals are derived from GAMs that include a spatial smooth for latitude and longitude to account for spatial autocorrelation.

$_{\scriptscriptstyle{590}}$ 3.2 Impact of cold-snaps and heat-waves on reproductive success

Because of differences in range and breeding timing, the species in our dataset varied widely 391 in their exposure to temperature during breeding (Figure 4A). During incubation, point 392 estimates from GAMs that controlled for date, year, and location, indicated that a two 393 standard deviation cold-snap reduced relative fitness in 8 out of 24 species (Figure 4B). The 394 species that were sensitive to cold-snaps in this period included purple martins, tree 395 swallows, eastern bluebirds, western bluebirds (Sialia mexicana), mountain bluebirds (Sialia currocoides), prothonotary warblers (Protonotaria citrea), Carolina chickadee (Poecile 397 carolinensis), and mountain chickadees (Poecile gambeli). Relative fitness estimates ranged 398 from 0.64 to 0.93 for these species; all values and confidence intervals for point estimates are 399 included in Table S2. Only two of these same species also showed evidence of reduced fitness 400 during a two SD incubation heat-wave (purple martin and eastern bluebird; Figure 4B, Table

S2). No species showed evidence of significantly increased fitness from either cold-snap or heat-wave exposure during incubation. 403 During the nestling provisioning period, 75% of the species (6 of the 8) that were sensitive to 404 cold-snaps in incubation also showed reduced relative fitness from two SD cold-snaps 405 occurring after hatching (prothonotary warbler and mountain chickadee were no longer susceptible; Figure 4C). In addition to these six species, eastern phoebes (Sayornis phoebe), violet-green swallows (Tachycineta thalassina), and six other species had reduced fitness 408 when a two SD cold-snap occurred during the provisioning period (Figure 4C). Overall, the 400 effect sizes for reduction in relative fitness during provisioning were comparable to those seen 410 during incubation (range of estimates for susceptible species was 0.74 to 0.94; Table S2). 411 Nine of the 14 species susceptible to cold-snaps were also susceptible to heat-waves during 412 provisioning (Figure 4C). Two additional species (brown-headed nuthatch: Sitta pusilla and 413 chestnut-backed chickadee: Poecile rufescens) were susceptible to heat-waves but not 414 cold-snaps during provisioning (Figure 4C). Similar to the incubation phase, no species 415 showed evidence of significantly increased fitness from either cold-snap or heat-wave 416

exposure during the nestling phase.

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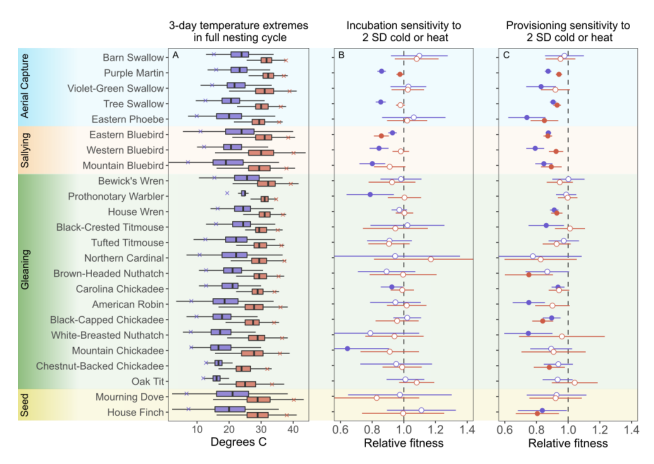


Figure 4: Exposure and sensitivity to cold-snaps and heat-waves across 24 species. (A) Distribution of the coldest and hottest 3-day high temperatures experienced during the full nesting cycle for all species sorted by the coldest average temperature within each foraging style. Boxplots indicate median, IQR, and 1.5 times IQR for the coldest (blue) and hottest (red) 3-day period. Blue and red x marks indicate points that are 2 SD below (cold) or above (hot) the mean exposure. (B and C) Predicted relative fitness for breeding attempts that experienced a 2 SD cold-snap (blue) or 2 SD heat-wave (red) during incubation (B) or provisioning (C). Point estimates are derived from GAMs that control for spatial patterns, year, and date. Circles show point estimate, lines show 95% CI; open circles indicate no difference from mean reproductive success while closed circles indicate a significant effect of temperature on relative reproductive success. Color bands show the primary foraging strategy that each species uses to provision nestlings.

3.3 Latitudinal variation in temperature exposure and

$_{\scriptscriptstyle{119}}$ susceptibility

- 420 We examined latitudinal trends in susceptibility to cold-snaps and heat-waves for eastern
- bluebirds, purple martins, and tree swallows. In all three species, breeding date was later
- farther north, but breeding attempts from northern areas still experienced lower 3-day

- coldest and 3-day hottest temperatures on average (Figure 5A-B, Figure S6A-B, and Figure S7A-B). While the overall patterns of susceptibility to cold-snaps and heat-waves were 424 largely similar across these species (Figure 4B-C), the latitudinal patterns differed somewhat 425 for each species. During both incubation and provisioning, eastern bluebirds had reduced 426 fitness from cold-snaps only near the northern edge of the range (Figure 5C-D; full details on 427 point estimates in Table S3). In contrast, relative fitness was reduced from heat-waves over a 428 wider, but somewhat inconsistent latitudinal extent (Figure 5C-D). 429 Purple martins susceptibility to heat-waves was only apparent near the southern edge of the 430 range and was more pronounced in provisioning than during the incubation period (Figure 431
- S6C-D). Unlike eastern bluebirds, purple martins had reduced relative fitness from cold-snaps during incubation and provisioning at nearly every latitude band. Only the southernmost two bands during provisioning showed no impact on fitness associated with cold-snaps (Figure S6C-D; Table S4).
- Like purple martins, tree swallows had consistently reduced relative fitness when cold-snaps occurred during incubation (Figure S7C). Despite an aggregate effect of heat-waves during incubation (Figure 4C), there was no signal for heat-wave effects during incubation or provisioning in any individual latitude band (Figure S7C-D). During provisioning, tree swallows only showed a sensitivity to cold-snaps at the northern range limit and no clear susceptibility to heat-waves (Figure S7D; Table S5).

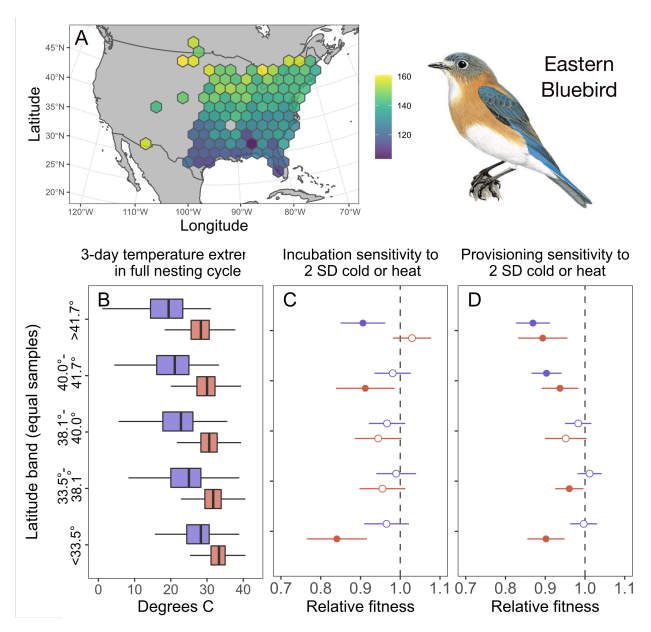


Figure 5: Breeding timing, temperature exposure, and extreme temperature sensitivity in eastern bluebirds. (A) Average date of first egg laying across the records included. (B) Distribution of the coldest and hottest 3-day high temperatures in the full nesting cycles in five latitude bands with an equal number of samples per band. The top of the panel is the northernmost band and bottom is the southernmost. Boxplots indicate median, IQR, and 1.5 times IQR for the coldest (blue) and hottest (red) 3-day period. (C and D) Predicted relative fitness for breeding attempts that experienced a 2 SD cold-snap (blue) or 2 SD heat-wave (red) during incubation (C) or provisioning (D) for each latitude band. Point estimates are derived from GAMs that control for spatial patterns, year, and date. Circles show point estimate, lines show 95 percent CI; open circles indicate no difference from mean reproductive success while closed circles indicate a significant effect of temperature on relative reproductive success. Illustration from Handbook of the Birds of the World by Ian Willis, copyright Lynx Edicions.

4 DISCUSSION

Using community scientist collected breeding records, we show that exposure to three-day 443 cold-snaps during a breeding attempt is associated with reduced relative fitness in 16 of 24 444 common species and that exposure to three-day heat-waves is associated with reduced fitness 445 in 11 of 24 species. In three widespread species, sensitivity to cold-snaps and heat-waves was 446 spatially heterogeneous, indicating a mechanism by which climate change might generate 447 population trends that differ across a species range. Historical temperature records for the 448 same area showed that the date of the latest cold-snap is only slightly earlier and the date of 440 the earliest heat-wave has not changed consistently over the past 70 years despite warmer 450 springs overall. Taken together, our results demonstrate that the likelihood of encountering 451 cold-snaps or heat-waves during breeding might differ as a consequence of climate change 452 and that exposure to these events often results in reduced reproductive success for many 453 common and widespread species. Moreover, in many cases it is impossible to track both average temperature and temperature variability through phenological shifts in the timing of 455 breeding. As a consequence, species that are most sensitive to extreme temperature events may be less able to adapt to changes in average temperature. Within species, variation in 457 sensitivity at northern and southern range limits might contribute to range shifts and local 458 population declines. Understanding the mechanisms that result in different sensitivity to 459 temperature variability within and between species should help accurately predict which 460 populations are most vulnerable to climate change. 461 Across the range of breeding records that we studied, average spring temperatures have 462 increased over the last 70 years, but the average timing of latest cold-snaps and earliest 463 heat-waves during the breeding season has not changed consistently in sign or magnitude. As 464 a consequence, the historical association between average temperature and temperature 465 variability differs from the association observed over the last 25 years. Thus, cues that wild 466 birds use to time seasonal events may not contain the same information about average 467

temperature and temperature variability that they did historically. Changes in breeding phenology have been identified as one of the 'universal responses' to climate change. ^{61,62} In 469 some cases, these shifts can minimize the effects of phenological mismatch with food 470 resources, ^{6,63-65} but in many cases the degree of change in breeding timing seems inadequate 471 to keep pace with average temperatures. ^{23,26,27} These inadequate shifts may be attributable 472 to a lack of ability for rapid evolutionary change or incomplete phenotypic plasticity. ^{66,67} 473 However, average breeding temperature and food availability are not the only gradients that 474 could be disrupted by climate change.²⁹ For species that are especially susceptible to 475 cold-snaps or heat-waves during breeding, changes in the timing of breeding events to track 476 average temperature might result in increased exposure to temperature variability. 31,34 Our 477 results highlight the consequences of temperature variation and suggest that in some cases 478 incomplete plasticity and a degree of phenological mismatch may represent an adaptive 470 response that balances competing selection pressures.⁶⁸ 480 We predicted that the consequences of cold-snaps might be especially severe during the 481 nestling provisioning period and for species that rely on insect activity for foraging. A recent 482 meta-analysis demonstrated that environmental insect food abundance is a strong predictor 483 of nestling body condition and fledging success, especially in species in which insects make 484 up a large component of the diet.⁶⁹ Daily total flying insect biomass and emergence rates are 485 strongly influenced by temperature, 38,70 so cold-snaps can have immediate and direct effects 486 on food availability. These impacts on food availability could compound the increased 487 thermoregulatory challenges and associated increase in energy use during cold-snaps. We did 488 find that more species were sensitive to cold-snaps during nestling provisioning, although several species were sensitive in both stages (6 species in both stages, 2 species in incubation only, 8 species in provisioning only). The 14 species that were sensitive to cold during provisioning included four of the five obligate aerial insectivores (purple martin, tree swallow, 492 violet-green-swallow, and eastern phoebe) and all three species that rely extensively on insect 493 movement for sally-capture foraging (eastern, western, and mountain bluebird). However, 494

white-breasted nuthatches (Sitta carolinensis) and American robins (Turdus migratorius) also showed declines despite a less obvious link to insect availability and the other aerial 496 insectivore did not exhibit sensitivity (barn swallow, *Hirundo rustica*). Thus our results suggest the possibility of foraging mode as a mechanism generating susceptibility to 498 cold-snaps during provisioning, but more data for a broader distribution of phylogenetically 499 independent species with different foraging styles would be needed to evaluate this prediction 500 convincingly. 501 In contrast to cold-snaps, we predicted that the primary challenge from heat-waves would be their impact on thermoregulation, rather than indirect effects of food availability. For 503 example, several studies demonstrate that heat dissipation rate may constrain reproductive 504 investment^{71,72} and hot conditions increase cooling costs.⁷³ Therefore, we expected to find 505 more widespread evidence for heat-wave impacts during both incubation and provisioning. 506 While we did find fairly widespread effects of heat-waves, fewer species were sensitive to 507 heat-waves than to cold-snaps (11 versus 18 species across both stages). For nine of these 508 species the sensitivity was only observed in one breeding stage. Moreover, the effect sizes for 500 reduction in relative fitness from heat-waves were generally smaller than those observed for 510 cold-snaps. While natural heat-waves during breeding can clearly drive reduced fitness in 511 some cases, ^{57,74} over the range and set of species we studied, cold-snaps seemed to generate 512 more consistent and stronger reductions in breeding success even though spring breeding 513 conditions are warmer overall than they were historically. One reason for these smaller 514 effects might be that the species in our study were able to mitigate the risks of hyperthermia 515 by changing behaviors or by choosing nesting sites that were cooler than the overall temperature recorded from nearby weather stations. It is also possible that a different characterization of heat-waves, such as one using average rather than daily high temperature 518 or a longer time period than three days, might have detected more vulnerability. 519 In one sense, it is surprising that we did not find universal evidence for cold-snap and

heat-wave consequences, because ultimately it is clear that a sufficiently severe temperature extreme would reduce fitness. Several non-mutually exclusive factors likely explain the lack of evidence for sensitivity that we observed in some species. First, species ranges are shaped, 523 in part, by direct and indirect effects of temperature. The Because we took the approach of 524 generating point estimates that were 2 SD from the mean of the observed breeding records 525 for each species, the absolute value of temperature challenges that we modeled differed 526 across species. While our approach explores the realized fitness declines from actual 527 temperature exposure, manipulations or extensions beyond these temperature extremes 528 would eventually uncover sensitivities in all species. Many experimental studies have 520 demonstrated physiological or fitness costs from cooling^{76,77} or heating^{78,79} nest boxes during 530 breeding, but these often employ more sustained or extreme temperatures than the natural 531 variation we modeled. On a related note, some species with relatively restricted ranges 532 limited by temperature exposure might be highly sensitive to temperature extremes, but 533 rarely experience those conditions during breeding, preventing us from detecting effects of 534 temperature challenges. Pigot et al. 2010 argued that widespread species range limits are 535 more likely to be defined by temperature, so it may be more common to detect sensitivities 536 to temperature extremes in those same widespread species⁷⁵. Finally, the community science datasets that we used yielded highly variable sample sizes and do not necessarily include representative observations across the range of most species. For widespread and easily 539 observed species it is likely easier to detect effects both because of larger sample sizes and 540 because of more dispersed sampling across the extent of each species range. Samples near 541 the range limits might be especially important for detecting sensitivity to temperature 542 extremes and are not equally available for all species in our dataset (e.g., bluebird trails and 543 nest boxes make monitoring especially easy and widespread for these species). 544 The results from our more detailed analysis of records from eastern bluebirds, purple martins, and tree swallows do indeed suggest that within-species spatial dynamics in 546 sensitivity to temperature variability may be highly relevant. Eastern bluebirds were

vulnerable to heat-waves over most of their range especially during provisioning, but were only sensitive to cold-snaps near the northern range limit. Purple martins were vulnerable to 549 cold-snaps across their range during incubation and in the northern half of the range during 550 provisioning, but were only sensitive to heat-waves near their southern range limit. Tree 551 swallows had no clear sensitivity to heat-waves within narrow latitude bands; they were 552 sensitive to cold-snaps in incubation across most of their range, but only sensitive to 553 cold-snaps during provisioning near their northern range limit. We did not quantify overall 554 abundance changes in our study, but it is interesting to note that since 2007 purple martins 555 have declined near their southern range limit and tree swallows have declined most 556 precipitously near their northern range limit while expanding their range southward.⁸⁰ These 557 patterns qualitatively match with the range limits near which we found differences in 558 sensitivity for each species. In contrast, eastern bluebirds have declined west of the 550 Appalachian Mountains and increased on the Atlantic seaboard over the same period, so it is 560 less clear how the patterns might relate to the sensitivity differences we found.⁸⁰ 561 Nevertheless, the fact that all three species show at least some evidence of increased 562 sensitivity to temperature near a range limit suggests that exposure to increased temperature 563 variability might be an important contributor to range shifts with climate change. However, despite the fact that all three species have similar body sizes, breeding behavior, and distributions, with two being fairly close relatives that are very similar ecologically, there was no common latitudinal pattern. Presumably these idiosyncratic responses are driven by 567 subtle differences, such as exposure to multiple additive stressors, life history details, or 568 habitat requirements. Thus, predicting differential sensitivity to temperature variability will 560 likely require a detailed understanding of the ecology for each species. 570 One important limitation on the conclusions of our study is that our data only allowed us to model overall fledging success, but it is likely that temperature variability also generates 572 sub-lethal effects that could have long term consequences on population demographics. Early 573 developmental conditions, including temperature, are well known to have a wide variety of

long lasting effects on wild birds, such as changes in body size and morphology, physiology, immunology, and survival to recruitment as a breeding adult.⁸¹ Indeed, long term declines in 576 body size is another widespread consequence of global warming.⁵ While changes in body size 577 may sometimes result from rapid evolutionary responses, many of the observed changes in 578 body size could also be explained by changes in developmental temperature.⁸² For example, 579 in tree swallows fledgling body size is positively correlated with average developmental 580 temperature and predicts both adult body size and likelihood of recruiting.⁸² Thus, 581 cold-snaps or heat-waves that are not severe enough to reduce fledging success could still 582 have important consequences on the demographics of bird populations through their long 583 term effects on nestlings. Our results should therefore be considered as a minimum measure 584 of the fitness costs from experiencing extreme temperature conditions. 585 A great deal of research has focused on the effects of climate change on wild bird populations 586 over the past several decades. While the large-scale patterns of climate change on ranges, 587 phenology, and morphology have been described for many species, studies that focus on 588 these patterns are often unable to characterize the organismal and ecological processes that 580 are operating. At the same time, targeted studies that focus on individually marked birds or 590 experimental manipulations often isolate mechanisms but cannot directly link their results to 591 population and range wide consequences. Our results add to a growing number of studies 592 that seek to make connections from individual-to-population level effects²⁵ in an effort to 593 understand how the sensitivity of populations to specific conditions ultimately creates larger 594 patterns that may differ between species. Moreover, we highlight the fact that average 595 temperature and temperature variability are both important and that they-along with other gradients-must be considered simultaneously in order to understand the impacts of climate change. Because many climate change models predict increasingly extreme weather in addition to overall warming, one of the challenges in predicting vulnerability for bird species 599 moving forward is to understand the extent to which average conditions versus extreme 600 events drive demographic consequences for populations as conditions change. 601

AUTHOR CONTRIBUTIONS

CCT & JRS contributed equally to conception and analysis of the manuscript. CCT drafted 603 the paper with input from JRS and both authors contributed to revisions. 604

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REFERENCES

- 1.
- NOAA. NOAA national centers for environmental information, state of the climate: 614 Global climate report for october 2021, published online november 2021, retrieved on november 24, 2021 from https://www.ncdc.noaa.gov/sotc/global/202110. (2021).
- 2. 616

615

- Parmesan, C. & Yohe, G. A globally coherent fingerprint of climate change impacts 617 across natural systems. *Nature* **421**, 37–42 (2003). 618
- 3. 619
- Ozgul, A. et al. The dynamics of phenotypic change and the shrinking sheep of st. 620
- kilda. Science **325**, 464–467 (2009).

- 622 4.
- Weeks, B. C. et al. Shared morphological consequences of global warming in north american migratory birds. Ecology Letters 23, 316–325 (2020).
- 625 5.
- Gardner, J. L., Peters, A., Kearney, M. R., Joseph, L. & Heinsohn, R. Declining body size: A third universal response to warming? Trends in Ecology & Evolution 26, 285–291 (2011).
- 628 6.
- Charmantier, A. et al. Adaptive phenotypic plasticity in response to climate change in a wild bird population. Science 320, 800–803 (2008).
- 631 7.
- Youngflesh, C. et al. Migratory strategy drives species-level variation in bird sensitivity to vegetation green-up. Nature Ecology & Evolution 5, 1–8 (2021).
- 634 8.
- Horton, K. G. et al. Phenology of nocturnal avian migration has shifted at the continental scale. Nature Climate Change 10, 63–68 (2020).
- 637 9.
- Blackford, C., Germain, R. M. & Gilbert, B. Species differences in phenology shape coexistence. *The American Naturalist* **195**, E168–E180 (2020).
- 640 10.
- Samplonius, J. M. *et al.* Strengthening the evidence base for temperature-mediated phenological asynchrony and its impacts. *Nature Ecology & Evolution* **5**, 155–164 (2021).
- 643 11.
- Cohen, J. M., Lajeunesse, M. J. & Rohr, J. R. A global synthesis of animal phenological
 responses to climate change. Nature Climate Change 8, 224–228 (2018).

- 646 12.
- Both, C., Van Asch, M., Bijlsma, R. G., Van Den Burg, A. B. & Visser, M. E. Climate change and unequal phenological changes across four trophic levels: Constraints or adaptations? *Journal of Animal Ecology* **78**, 73–83 (2009).
- 649 13.
- Shipley, J. R. et al. Climate change shifts the timing of nutritional flux from aquatic insects. Current Biology 32, 1342–1349 (2022).
- 652 14.
- Vasseur, D. A. et al. Increased temperature variation poses a greater risk to species than climate warming. Proceedings of the Royal Society B: Biological Sciences 281, 20132612 (2014).
- 655 15.
- Thackeray, S. J. et al. Phenological sensitivity to climate across taxa and trophic levels. Nature **535**, 241–245 (2016).
- 658 16.
- Rosenberg, K. V. et al. Decline of the north american avifauna. Science **366**, 120–124 (2019).
- 661 17.
- Loss, S. R., Will, T. & Marra, P. P. The impact of free-ranging domestic cats on wildlife of the united states. *Nature communications* 4, 1–8 (2013).
- 664 18.
- 665 Senzaki, M. et al. Sensory pollutants alter bird phenology and fitness across a continent.
- Nature **587**, 605–609 (2020).
- 667 19.
- Julliard, R., Jiguet, F. & Couvet, D. Common birds facing global changes: What makes a species at risk? *Global Change Biology* **10**, 148–154 (2004).

- 670 20.
- Robinson, R. A. et al. Emerging infectious disease leads to rapid population declines of common british birds. PLoS One 5, e12215 (2010).
- 673 21.
- Møller, A. P., Rubolini, D. & Lehikoinen, E. Populations of migratory bird species that did not show a phenological response to climate change are declining. *Proceedings* of the National Academy of Sciences 105, 16195–16200 (2008).
- 676 22.
- Franks, S. E. *et al.* The sensitivity of breeding songbirds to changes in seasonal timing is linked to population change but cannot be directly attributed to the effects of trophic asynchrony on productivity. *Global Change Biology* **24**, 957–971 (2018).
- 679 23.
- Devictor, V., Julliard, R., Couvet, D. & Jiguet, F. Birds are tracking climate warming,
 but not fast enough. Proceedings of the Royal Society B: Biological Sciences 275,
 2743–2748 (2008).
- 682 24.
- Both, C., Bouwhuis, S., Lessells, C. & Visser, M. E. Climate change and population declines in a long-distance migratory bird. *Nature* **441**, 81–83 (2006).
- 685 25.
- Socolar, J. B., Epanchin, P. N., Beissinger, S. R. & Tingley, M. W. Phenological shifts conserve thermal niches in north american birds and reshape expectations for climate-driven range shifts. *Proceedings of the National Academy of Sciences* 114, 12976–12981 (2017).
- 688 26.

- Simmonds, E. G., Cole, E. F., Sheldon, B. C. & Coulson, T. Phenological asynchrony:
 A ticking time-bomb for seemingly stable populations? *Ecology Letters* **23**, 1766–1775

 (2020).
- 691 27.
- Radchuk, V. et al. Adaptive responses of animals to climate change are most likely insufficient. Nature Communications 10, 1–14 (2019).
- 694 28.
- Gienapp, P. et al. Predicting demographically sustainable rates of adaptation: Can great tit breeding time keep pace with climate change? Philosophical Transactions of the Royal Society B: Biological Sciences 368, 20120289 (2013).
- 697 29.
- Spence, A. R. & Tingley, M. W. The challenge of novel abiotic conditions for species undergoing climate-induced range shifts. *Ecography* **43**, 1571–1590 (2020).
- 700 30.
- DeGregorio, B. A., Westervelt, J. D., Weatherhead, P. J. & Sperry, J. H. Indirect effect of climate change: Shifts in ratsnake behavior alter intensity and timing of avian
 nest predation. Ecological Modelling 312, 239–246 (2015).
- 703 31.
- Shipley, J. R. et al. Birds advancing lay dates with warming springs face greater risk of chick mortality. Proceedings of the National Academy of Sciences 117, 25590–25594 (2020).
- 706 32.
- Marrot, P., Garant, D. & Charmantier, A. Multiple extreme climatic events strengthen selection for earlier breeding in a wild passerine. *Philosophical Transactions of the Royal Society B: Biological Sciences* **372**, 20160372 (2017).
- 709 33.

- Townsend, A. K. *et al.* The interacting effects of food, spring temperature, and global climate cycles on population dynamics of a migratory songbird. *Global Change Biology*22, 544–555 (2016).
- 712 34.
- Regan, C. E. & Sheldon, B. C. Phenotypic plasticity increases exposure to extreme climatic events that reduce individual fitness. *Global Change Biology* **29**, 2968–2980 (2023).
- 715 35.
- Kearney, M. R., Jusup, M., McGeoch, M. A., Kooijman, S. A. & Chown, S. L. Where do functional traits come from? The role of theory and models. *Functional Ecology* 35, 1385–1396 (2021).
- 718 36.
- Belitz, M. et al. Climate drivers of adult insect activity are conditioned by life history traits. Ecology Letters 24, 2687–2699 (2021).
- 721 37.
- Riddell, E. et al. Exposure to climate change drives stability or collapse of desert mammal and bird communities. Science **371**, 633–636 (2021).
- 724 38.
- Winkler, D. W., Luo, M. K. & Rakhimberdiev, E. Temperature effects on food supply and chick mortality in tree swallows (tachycineta bicolor). *Oecologia* **173**, 129–138 (2013).
- 727 39.
- Albright, T. P. *et al.* Combined effects of heat waves and droughts on avian communities across the conterminous united states. *Ecosphere* 1, 1–22 (2010).
- ₇₃₀ 40.

- McKechnie, A. E. & Wolf, B. O. Climate change increases the likelihood of catastrophic
- avian mortality events during extreme heat waves. Biology Letters 6, 253–256 (2010).
- 733 41.
- Visser, M. E., Lindner, M., Gienapp, P., Long, M. C. & Jenouvrier, S. Recent natural variability in global warming weakened phenological mismatch and selection on seasonal timing in great tits (parus major). *Proceedings of the Royal Society B* 288, 20211337 (2021).
- ₇₃₆ 42.
- Jirinec, V. et al. Morphological consequences of climate change for resident birds in intact amazonian rainforest. Science Advances 7, eabk1743 (2021).
- ₇₃₉ 43.
- Freeman, B. G., Scholer, M. N., Ruiz-Gutierrez, V. & Fitzpatrick, J. W. Climate change causes upslope shifts and mountaintop extirpations in a tropical bird community.
- Proceedings of the National Academy of Sciences 115, 11982–11987 (2018).
- ₇₄₂ 44.
- Thomas, C. D. & Lennon, J. J. Birds extend their ranges northwards. *Nature* **399**, 213–213 (1999).
- ₇₄₅ 45.
- Smith, M. D. An ecological perspective on extreme climatic events: A synthetic definition and framework to guide future research. *Journal of Ecology* 99, 656–663 (2011).
- 748 46.
- Bailey, L. D. & Pol, M. van de. Tackling extremes: Challenges for ecological and evolutionary research on extreme climatic events. *Journal of Animal Ecology* **85**, 85–96 (2016).
- ₇₅₁ 47.

- Williams, H. M., Siegrist, J. & Wilson, A. M. Support for a relationship between demography and modeled habitat suitability is scale dependent for the purple martin progne subis. *Journal of Animal Ecology* **90**, 356–366 (2021).
- ₇₅₄ 48.
- Billerman, S. M., Keeney, B. K., Rodewald, P. G. & Schulenberg, T. S. Birds of the world. (https://birdsoftheworld-org.proxy.library.cornell.edu/bow/home; Cornell
 Laboratory of Ornithology, Ithaca, NY USA, 2020).
- ₇₅₇ 49.
- Hollister, J., Shah, T., Robitaille, A. L., Beck, M. W. & Johnson, M. *Elevatr: Access elevation data from various APIs.* (2020).
- ₇₆₀ 50.
- South, A. Rnaturalearth: World map data from natural earth. (2017).
- ₇₆₃ 51.

762

- R Core Team. R: A language and environment for statistical computing. (R Foundation for Statistical Computing, 2020).
- ₇₆₆ 52.
- Pebesma, E. Simple Features for R: Standardized Support for Spatial Vector Data.
- The R Journal 10, 439–446 (2018).
- ₇₆₉ 53.
- Chamberlain, S. Rnoaa: 'NOAA' weather data from r. (2020).
- ₇₇₂ 54.

771

LaZerte, S. E. & Albers, S. weathercan: Download and format weather data from environment and climate change canada. The Journal of Open Source Software 3, 571 (2018).

- 775 55.
- Fisher, R. J. et al. Extreme precipitation reduces reproductive output of an endangered
- raptor. Journal of Applied Ecology **52**, 1500–1508 (2015).
- 778 56.
- G. MCDonald, P., Olsen, P. D. & Cockburn, A. Weather dictates reproductive success and survival in the australian brown falcon falco berigora. *Journal of Animal Ecology* 73, 683–692 (2004).
- ₇₈₁ 57.
- Corregidor-Castro, A. et al. Experimental nest cooling reveals dramatic effects of heatwaves on reproduction in a mediterranean bird of prey. Global Change Biology (2023).
- ₇₈₄ 58.
- Brown, C. R. & Brown, M. B. Intense natural selection on body size and wing and tail
- asymmetry in cliff swallows during severe weather. Evolution **52**, 1461–1475 (1998).
- ₇₈₇ 59.
- Knief, U. & Forstmeier, W. Violating the normality assumption may be the lesser of two evils. Behavior Research Methods **53**, 2576–2590 (2021).
- 790 60.
- Wood, S. N. Generalized additive models: An introduction with r. (CRC press, 2017).
- ₇₉₃ 61.

792

- Kharouba, H. M. et al. Global shifts in the phenological synchrony of species interactions over recent decades. Proceedings of the National Academy of Sciences 115, 5211–5216 (2018).
- 796 62.

- Horton, K. G., Morris, S. R., Van Doren, B. M. & Covino, K. M. Six decades of north american bird banding records reveal plasticity in migration phenology. *Journal of Animal Ecology* **92**, 738–750 (2023).
- 799 63.
- Iler, A. M. *et al.* Maintenance of temporal synchrony between syrphid flies and floral resources despite differential phenological responses to climate. *Global Change Biology*19, 2348–2359 (2013).
- 802 64.
- Vatka, E., Orell, M. & Rytkönen, S. Warming climate advances breeding and improves synchrony of food demand and food availability in a boreal passerine. *Global Change Biology* 17, 3002–3009 (2011).
- 805 65.
- Twining, C. W., Shipley, J. R. & Matthews, B. Climate change creates nutritional phenological mismatches. *Trends in Ecology & Evolution* **37**, 736–739 (2022).
- 808 66.
- Both, C. & Visser, M. E. Adjustment to climate change is constrained by arrival date in a long-distance migrant bird. *Nature* **411**, 296–298 (2001).
- 811 67.
- Visser, M. E., Both, C. & Lambrechts, M. M. Global climate change leads to mistimed avian reproduction. *Advances in Ecological Research* **35**, 89–110 (2004).
- 814 68.
- Visser, M. E., Marvelde, L. te & Lof, M. E. Adaptive phenological mismatches of birds and their food in a warming world. *Journal of Ornithology* **153**, 75–84 (2012).
- 817 69.

- Grames, E. M., Montgomery, G. A., Youngflesh, C., Tingley, M. W. & Elphick, C. S. The effect of insect food availability on songbird reproductive success and chick body condition: Evidence from a systematic review and meta-analysis. *Ecology Letters* **26**, 658–673 (2023).
- 820 70.
- Dunn, P. O. *et al.* Extensive regional variation in the phenology of insects and their response to temperature across n orth a merica. *Ecology* **104**, e4036 (2023).
- 823 71.
- Nord, A. & Nilsson, J.-Å. Heat dissipation rate constrains reproductive investment in a wild bird. Functional Ecology 33, 250–259 (2019).
- 826 72.
- Tapper, S., Nocera, J. J. & Burness, G. Heat dissipation capacity influences reproductive performance in an aerial insectivore. *Journal of Experimental Biology* **223**, jeb222232 (2020).
- 829 73.
- Riddell, E., Iknayan, K., Wolf, B., Sinervo, B. & Beissinger, S. Cooling requirements fueled the collapse of a desert bird community from climate change. *Proceedings of the National Academy of Sciences* 116, 21609–21615 (2019).
- 832 74.
- Piatt, J. F. et al. Extreme mortality and reproductive failure of common murres resulting from the northeast pacific marine heatwave of 2014-2016. PloS One 15, e0226087 (2020).
- 835 75.
- Pigot, A. L., Owens, I. P. & Orme, C. D. L. The environmental limits to geographic range expansion in birds. *Ecology Letters* **13**, 705–715 (2010).
- 838 76.

- Ardia, D. R., Pérez, J. H. & Clotfelter, E. D. Experimental cooling during incubation leads to reduced innate immunity and body condition in nestling tree swallows.
- Proceedings of the Royal Society B: Biological Sciences 277, 1881–1888 (2010).
- 841 77.
- Nilsson, J. F., Stjernman, M. & Nilsson, J.-Ä. Experimental reduction of incubation temperature affects both nestling and adult blue tits cyanistes caeruleus. *Journal of Avian Biology* **39**, 553–559 (2008).
- 844 78.
- Pérez, J. H., Ardia, D. R., Chad, E. K. & Clotfelter, E. D. Experimental heating reveals nest temperature affects nestling condition in tree swallows (Tachycineta bicolor).
- Biology letters 4, 468–471 (2008).
- 847 79.
- Andreasson, F., Nord, A. & Nilsson, J.-Ä. Experimentally increased nest temperature affects body temperature, growth and apparent survival in blue tit nestlings. *Journal of Avian Biology* **49**, jav–01620 (2018).
- 850 80.
- Fink, D. et al. eBird status and trends, data version: 2021; released: 2022. Cornell
- Lab of Ornithology, Ithaca, New York (2022) doi:https://doi.org/10.2173/ebirdst.2021.
- 853 81.
- Lindström, J. Early development and fitness in birds and mammals. *Trends in ecology*854 & evolution 14, 343–348 (1999).
- 856 82.
- Shipley, J. R., Twining, C. W., Taff, C. C., Vitousek, M. N. & Winkler, D. W. Selection counteracts developmental plasticity in body-size responses to climate change. *Nature Climate Change* **12**, 863–868 (2022).

SUPPLEMENTAL TABLES AND FIGURES

Table S1: Total number of breeding records for each included species.

Common name	Scientific name	Records
American Robin	Turdus migratorius	2101
Barn Swallow	Hirundo rustica	1003
Bewick's Wren	Thryomanes bewickii	1421
Black-Capped Chickadee	Poecile atricapillus	2993
Black-Crested Titmouse	Baeolophus atricristatus	527
Brown-Headed Nuthatch	Sitta pusilla	375
Carolina Chickadee	Poecile carolinensis	5287
Chestnut-Backed Chickadee	Poecile rufescens	951
Eastern Bluebird	Sialia sialis	80447
Eastern Phoebe	Sayornis phoebe	998
House Finch	Haemorphous mexicanus	627
House Wren	$Troglodytes \ aedon$	16946
Mountain Bluebird	$Sialia\ currocoides$	9051
Mountain Chickadee	Poecile gambeli	526
Mourning Dove	Zenaida macroura	524
Northern Cardinal	Cardinalis cardinalis	512
Oak Tit	Baeolophus inornatus	1066
Prothonotary Warbler	Protonotaria citrea	2121
Purple Martin	Progne subis	85368
Tree Swallow	Tachycineta bicolor	64222
Tufted Titmouse	Baeolophus bicolor	1224
Violet-Green Swallow	Tachycineta thalassina	2410
Western Bluebird	Sialia mexicana	20335
White-Breasted Nuthatch	Sitta carolinensis	479

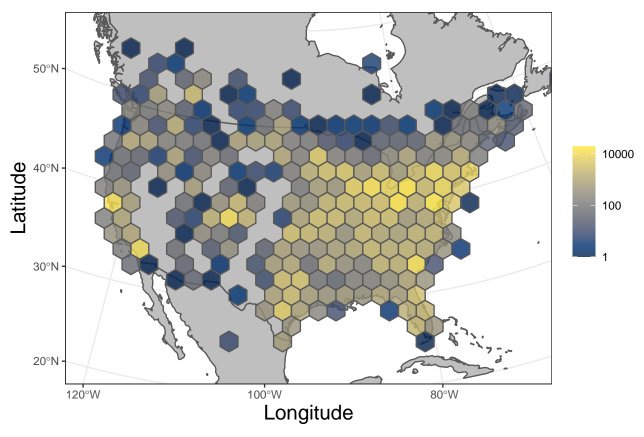


Figure S1: Number of breeding records included within each grid cell for all species and years from the combined datasets after filtering. A total of 301,514 records from 24 species are included from 1995 to 2020.

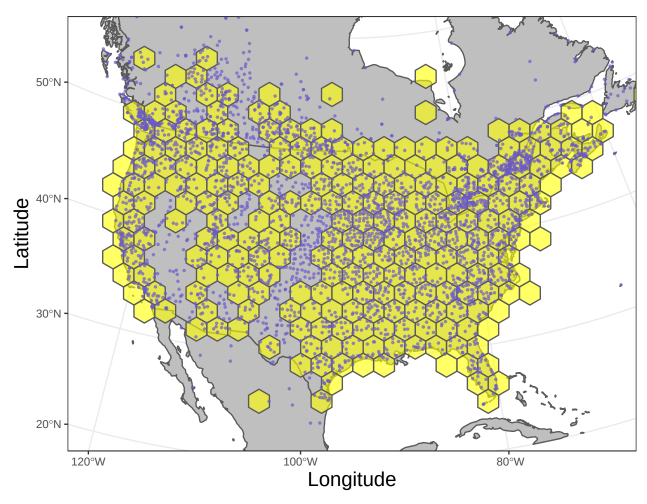


Figure S2: Weather stations used to access historical temperature data. Individual points show station locations with data downloaded from NOAA (n=2608) or Environment and Climate Change Canada (n=1125). Stations that had at least 50 years of available data were included. Yellow hexagons show the spatial extent of the avian breeding records used in the analysis to illustrate how coverage of temperature data corresponds to the main dataset.



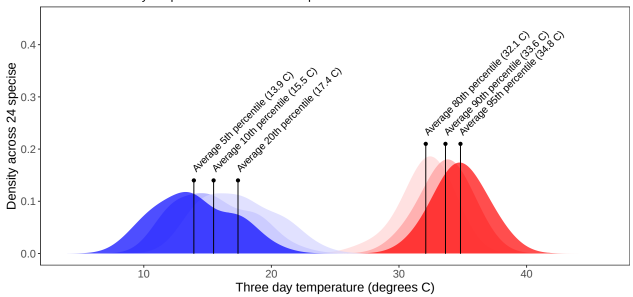


Figure S3: Distribution of extreme climatic events for the 24 species included in this study used to determine threshold temperatures for analysis of historical cold-snap and heat-wave timing. Six distributions are shown, illustrating the 5th, 10th, and 20th percentile of the coldest three day period during breeding attempts and the 80th, 90th, and 95th percentile of hottest three day period during breeding attempts. Black lines indicate the raw averages across 24 species for each distribution and these values were used in the historical analysis described in the text.

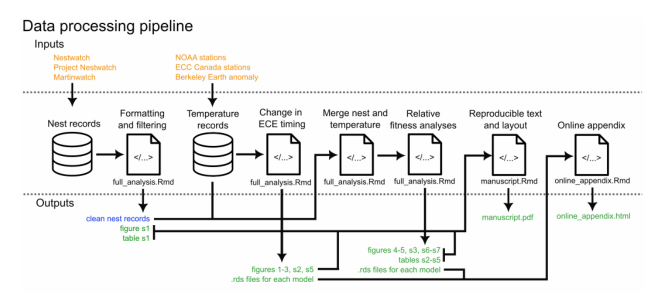


Figure S4: Schematic illustration of the data analysis workflow from raw data inputs to data outputs, manuscript file, and online supplement. To reproduce the analyses, raw data shown in orange will need to be accessed from each appropriate repository in order to produce the output shown in blue. Downstream output files shown in green are available in the archived data and code repository along with all of the scripts referenced in the figure.

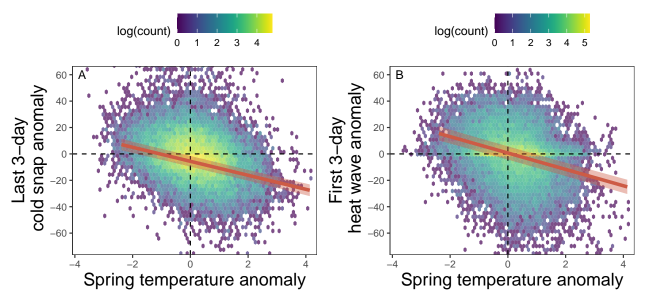


Figure S5: Spring average temperature anomaly in relation to the date of the last three day cold-snap (A) or earliest heat-wave (B). Grid shows the count of total cell level observations at each combination with all years counted. Red line is a fit from a GAM that includes a basis spline for year and tensor smooth for latitude and longitude to account for spatial and temporal autocorrelation. Shaded red region shows the 95% confidence interval of the fit GAM.

Table S2: Point estimates for the expected relative fitness of each species during a two standard deviation cold-snap or heat-wave based on GAMs described in text. Numbers in brackets show 95-percent confidence intervals.

	Incubation	Incubation	Provisioning	Provisioning
	cold-snap	heat-wave	cold-snap	heat-wave
American robin	0.96 [0.80, 1.12]	1.02 [0.89, 1.14]	$0.75 \ [0.65, \ 0.85]$	0.90 [0.79, 1.01]
Barn swallow	1.10 [0.92, 1.28]	1.08 [0.94, 1.22]	0.98 [0.85, 1.10]	0.95 [0.86, 1.05]
Bewick's wren	0.98 [0.85, 1.11]	0.93 [0.78, 1.07]	1.00 [0.90, 1.10]	0.95 [0.86, 1.03]
Black-capped	1.02 [0.93, 1.11]	0.96 [0.82, 1.09]	0.90 [0.84, 0.95]	$0.84 \ [0.77, \ 0.91]$
chickadee Black-crested titmouse	1.02 [0.79, 1.26]	0.95 [0.74, 1.15]	0.86 [0.75, 0.97]	1.01 [0.92, 1.11]
Brown-headed	$0.89 \ [0.71, \ 1.07]$	$1.00 \ [0.78, \ 1.21]$	$0.87 \ [0.73, \ 1.01]$	$0.75 \ [0.60, \ 0.90]$
nuthatch Carolina	0.92 [0.85, 0.99]	0.99 [0.92, 1.06]	0.94 [0.90, 0.98]	0.94 [0.88, 1.01]
chickadee Chestnut- backed	0.95 [0.73, 1.18]	0.99 [0.87, 1.11]	0.94 [0.85, 1.03]	0.88 [0.78, 0.98]
chickadee Eastern bluebird	0.93 [0.90, 0.95]	0.86 [0.81, 0.91]	0.87 [0.86, 0.89]	0.87 [0.85, 0.90]
Eastern phoebe	1.06 [0.86, 1.26]	$1.02 \ [0.89, \ 1.15]$	$0.74 \ [0.62, \ 0.86]$	$0.85 \ [0.76, \ 0.94]$
House finch	1.11 [0.89, 1.33]	1.01 [0.77, 1.25]	0.85 [0.69, 1.00]	$0.80 \ [0.66, \ 0.93]$
House wren	0.97 [0.92, 1.02]	1.00 [0.95, 1.06]	0.91 [0.89, 0.94]	0.93 [0.89, 0.96]
Mountain	0.80 [0.72, 0.88]	0.91 [0.81, 1.01]	0.85 [0.79, 0.90]	0.89 [0.83, 0.96]
bluebird Mountain	0.64 [0.38, 0.91]	0.91 [0.73, 1.10]	0.89 [0.76, 1.03]	0.91 [0.71, 1.11]
chickadee Mourning dove	0.96 [0.63, 1.29]	0.83 [0.56, 1.10]	0.92 [0.73, 1.11]	0.92 [0.76, 1.09]
Northern	$0.96 \ [0.55, \ 1.37]$	1.15 [0.79, 1.51]	$0.80 \ [0.49, \ 1.10]$	0.81 [0.58, 1.04]
cardinal Oak titmouse Prothonotary	1.01 [0.89, 1.13] 0.79 [0.64, 0.94]	1.08 [0.97, 1.19] 1.00 [0.90, 1.11]	0.93 [0.84, 1.03] 0.99 [0.92, 1.05]	1.04 [0.90, 1.19] 1.00 [0.93, 1.06]
warbler Purple martin Tree swallow	0.86 [0.83, 0.89] 0.85 [0.82, 0.89]	0.98 [0.95, 1.00] 0.98 [0.95, 1.01]	0.87 [0.85, 0.89] 0.91 [0.89, 0.93]	0.94 [0.92, 0.96] 0.93 [0.91, 0.95]
Tufted titmouse	0.91 [0.77, 1.05]	0.91 [0.78, 1.04]	0.97 [0.88, 1.07]	0.93 [0.84, 1.02]
Violet-green	1.03 [0.92, 1.14]	1.03 [0.92, 1.13]	0.83 [0.74, 0.92]	0.92 [0.83, 1.01]
swallow Western	0.84 [0.78, 0.90]	0.98 [0.93, 1.03]	0.79 [0.74, 0.85]	0.92 [0.88, 0.97]
bluebird White-breasted	0.79 [0.48, 1.10]	0.94 [0.76, 1.13]	0.75 [0.60, 0.90]	0.96 [0.69, 1.23]
nuthatch				

Table S3: Point estimates for the expected relative fitness of eastern bluebirds during a two standard deviation cold-snap or heat-wave based on GAMs described in text. Breeding records are split into five latitude bands with an equal number of records per band; the southernmost band includes nests below 33.5 degrees latitude and is followed by 33.5 to 38.1 degrees, 38.1 to 40.0 degrees, 40.0 to 41.7 degrees, and finally nests above 41.7 degrees in the northernmost band. Numbers in brackets show 95-percent confidence intervals.

	Incubation	Incubation	Provisioning	Provisioning
	cold-snap	heat-wave	cold-snap	heat-wave
Northernmost	0.98 [0.92, 1.04]	1.02 [0.96, 1.08]	0.74 [0.69, 0.79]	0.82 [0.77, 0.88]
North	1.00 [0.92, 1.09]	0.98 [0.88, 1.08]	0.85 [0.77, 0.93]	0.85 [0.75, 0.94]
Middle	0.99 [0.92, 1.06]	0.86 [0.76, 0.97]	0.87 [0.81, 0.93]	0.83 [0.77, 0.90]
South	1.04 [0.96, 1.11]	0.89 [0.80, 0.98]	0.92 [0.86, 0.97]	0.80 [0.73, 0.86]
Southernmost	0.98 [0.89, 1.07]	0.88 [0.77, 0.99]	0.86 [0.79, 0.93]	0.74 [0.66, 0.83]

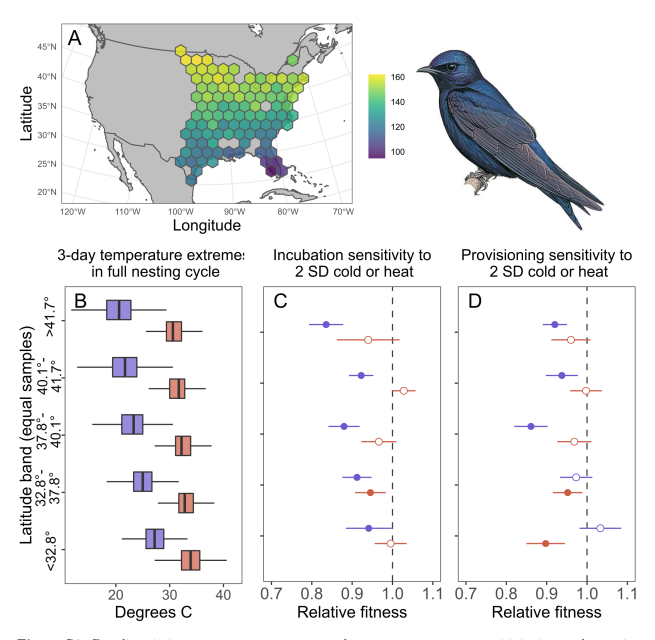


Figure S6: Breeding timing, temperature exposure, and extreme temperature sensitivity in purple martins. (A) Average date of first egg laying across the records included. (B) Distribution of the coldest and hottest 3-day high temperatures in the full nesting cycles in five latitude bands with an equal number of samples per band. The top of the panel is the northernmost band and bottom is the southernmost. Boxplots indicate median, IQR, and 1.5 times IQR for the coldest (blue) and hottest (red) 3-day period. (C and D) Predicted relative fitness for breeding attempts that experienced a 2 SD cold-snap (blue) or 2 SD heat-wave (red) during incubation (C) or provisioning (D) for each latitude band. Point estimates are derived from GAMs that control for spatial patterns, year, and date. Circles show point estimate, lines show 95 percent CI; open circles indicate no difference from mean reproductive success while closed circles indicate a significant effect of temperature on relative reproductive success. Illustration from Handbook of the Birds of the World by Hilary Bum, copyright Lynx Edicions.

Table S4: Point estimates for the expected relative fitness of purple martins during a two standard deviation cold-snap or heat-wave based on GAMs described in text. Breeding records are split into five latitude bands with an equal number of records per band; the southernmost band includes nests below 32.8 degrees latitude and is followed by 32.8 to 37.8 degrees, 37.8 to 40.1 degrees, 40.1 to 41.7 degrees, and finally nests above 41.7 degrees in the northernmost band. Numbers in brackets show 95-percent confidence intervals.

	Incubation	Incubation	Provisioning	Provisioning
	cold-snap	heat-wave	cold-snap	heat-wave
Northernmost	0.84 [0.79, 0.88]	0.94 [0.86, 1.02]	0.92 [0.89, 0.95]	0.96 [0.91, 1.01]
North	0.92 [0.89, 0.95]	1.03 [1.00, 1.06]	0.94 [0.90, 0.98]	1.00 [0.96, 1.04]
Middle	0.88 [0.84, 0.92]	0.97 [0.92, 1.01]	0.86 [0.82, 0.90]	0.97 [0.93, 1.01]
South	0.91 [0.88, 0.95]	0.95 [0.91, 0.98]	0.97 [0.93, 1.01]	0.95 [0.92, 0.99]
Southernmost	0.94 [0.88, 1.00]	1.00 [0.96, 1.04]	1.03 [0.98, 1.08]	0.90 [0.85, 0.95]

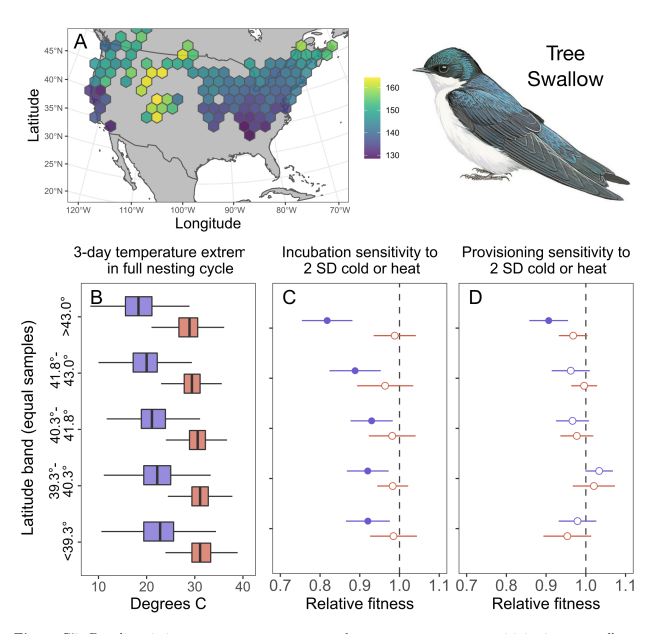


Figure S7: Breeding timing, temperature exposure, and extreme temperature sensitivity in tree swallows. (A) Average date of first egg laying across the records included. (B) Distribution of the coldest and hottest 3-day high temperatures in the full nesting cycles in five latitude bands with an equal number of samples per band. The top of the panel is the northernmost band and bottom is the southernmost. Boxplots indicate median, IQR, and 1.5 times IQR for the coldest (blue) and hottest (red) 3-day period. (C and D) Predicted relative fitness for breeding attempts that experienced a 2 SD cold-snap (blue) or 2 SD heat-wave (red) during incubation (C) or provisioning (D) for each latitude band. Point estimates are derived from GAMs that control for spatial patterns, year, and date. Circles show point estimate, lines show 95 percent CI; open circles indicate no difference from mean reproductive success while closed circles indicate a significant effect of temperature on relative reproductive success. Analysis by latitude band included only species in the eastern range, see methods for details. Illustration from Handbook of the Birds of the World by Hilary Bum, copyright Lynx Edicions.

Table S5: Point estimates for the expected relative fitness of tree swallows during a two standard deviation cold-snap or heat-wave based on GAMs described in text. Breeding records are split into five latitude bands with an equal number of records per band; the southernmost band includes nests below 39.3 degrees latitude and is followed by 39.3 to 40.3 degrees, 40.3 to 41.8 degrees, 41.8 to 43 degrees, and finally nests above 43 degrees in the northernmost band. Numbers in brackets show 95-percent confidence intervals.

	Incubation	Incubation	Provisioning	Provisioning
	cold-snap	heat-wave	cold-snap	heat-wave
Northernmost	0.82 [0.75, 0.88]	0.99 [0.93, 1.04]	0.91 [0.86, 0.96]	0.97 [0.93, 1.00]
North	0.89 [0.82, 0.95]	0.96 [0.89, 1.03]	0.96 [0.91, 1.01]	1.00 [0.96, 1.03]
Middle	0.93 [0.88, 0.98]	0.98 [0.92, 1.04]	0.97 [0.93, 1.01]	0.98 [0.94, 1.02]
South	0.92 [0.87, 0.97]	0.98 [0.94, 1.02]	1.03 [1.00, 1.07]	1.02 [0.97, 1.07]
Southernmost	0.92 [0.87, 0.98]	0.98 [0.93, 1.04]	0.98 [0.93, 1.03]	0.95 [0.89, 1.01]