Climate effects on nesting phenology in Nebraska turtles

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Abstract
A frequent response of organisms to climate change is altering the timing of reproduction, and advancement of reproductive timing has been a common reaction to warming temperatures in temperate regions. We tested whether this pattern applied to two common North American turtle species over the past three decades in Nebraska, USA. The timing of nesting (either first date or average date) of the Common Snapping Turtle (Chelydra serpentina) was negatively correlated with mean December maximum temperatures of the preceding year and mean May minimum and maximum temperatures in the nesting year and positively correlated with precipitation in July of the previous year. Increased temperatures during the late winter and spring likely permit earlier emergence from hibernation, increased metabolic rates and feeding opportunities, and accelerated vitellogenesis, ovulation, and egg shelling, all of which could drive earlier nesting. However, for the Painted Turtle (Chrysemys picta), the timing of nesting was positively correlated with mean minimum temperatures in September, October, December of the previous year, February of the nesting year, and April precipitation. These results suggest warmer fall, and winter temperature may impose an increased metabolic cost to painted turtles that impedes fall vitellogenesis, and April rains may slow the completion of vitellogenesis through decreased basking opportunities. For both species, nest deposition was highly correlated with body size, and larger females nested earlier in the season. Although average annual ambient temperatures have increased over the last four decades of our overall fieldwork at our study site, spring temperatures have not yet increased, and hence, nesting phenology has not advanced at our site for Chelydra. While Chrysemys exhibited a weak trend toward later nesting, this response was likely due to increased recruitment of smaller females into the population due to nest protection and predator control (Procyon lotor) in the early 2000s. Should climate change result in an increase in spring temperatures, nesting phenology would presumably respond accordingly, conditional on body size variation within these populations.

KEYWORDS
Chelydra, Chrysemys, climate change, nesting phenology, precipitation, weather
1 | INTRODUCTION

There is near consensus in the scientific community that climate change is a reality and is accelerating, and its biological effects are already being felt (Pachauri et al., 2014). Considerable evidence already exists demonstrating the impacts of climate change (especially warming) on the reproductive phenology of plants (Ahas, 1999; Price & Waser, 1998; Sherry et al., 2007), invertebrates (Roy & Sparks, 2000), amphibians (Benard, 2015; Gibbs & Breisch, 2001; While & Uller, 2014), fish (Parmesan & Yohe, 2003; Root et al., 2003), birds (Bowers et al., 2016; Bradey et al., 1999; Charmantier & Gienapp, 2014; Crick et al., 1997; Fritts et al., 2018), mammals (Réale et al., 2003), lizards (Bull & Burzacott, 2002; Telemecko et al., 2009), snakes (Moreno-Rueda et al., 2009), and turtles (citations in Table 1). The emerging pattern in temperate regions is that as temperatures have warmed, reproductive phenologies (e.g., courtship, nesting, and birth) have advanced in time (i.e., occur earlier in the spring or summer).

Earlier reproduction has the potential to severely disrupt an organism’s life cycle. On the positive side, for temperate animals it might permit the production of additional clutches or broods, and neonates may have more time to feed and grow in the fall prior to their first winter (Carroll & Ultsch, 2007; Rhen & Lang, 1999; Schwanz & Janzen, 2008; Tucker et al., 2008). However, on the negative side, early reproduction might expose eggs or neonates to atypical or mismatched conditions, increasing mortality rates (Benard, 2015; Jara et al., 2019; Muir et al., 2012; Pike et al., 2006; Saino et al., 2011). The potential impact of earlier nesting is especially complicated for species that exhibit temperature-dependent sex determination during development in the nest, as most turtles do (Janzen & Paukstis, 1991). Hence, understanding the impacts of climate on the reproductive phenology of turtles (and other organisms) is critical to conservation and management in the face of climate change, but also because turtles are among the most endangered organisms on the planet (Stanford et al., 2020).

We have been studying the reproduction and demography of turtle populations in western Nebraska since 1981 (Iverson, 1991; Iverson et al., 1997; Iverson & Smith, 1993). For this study, we sought to examine the effects of climate variables on the nesting phenology in two species: Common Snapping Turtle (*Chelydra serpentina*) and Painted Turtle (*Chrysemys picta*). Detailed descriptions of the reproductive biology of these two species at this site have been previously reported (Iverson et al., 1997; Iverson & Smith, 1993). Both species nest annually in late May to late June or early July, but the timing of nesting varies among years by as much as two weeks in *Chelydra* and over a month in *Chrysemys* (see below). That variation is likely related to variability in weather, but the specific climatic variables that drive nest timing, and how those variables might be changing over time, have not been studied at this site.

Preliminary data from our site suggested that cooler springs delayed the onset of nesting in turtles (date of first nest only; Janzen et al., 2018). For this expanded study, we predicted that spring temperatures would be inversely correlated with the Julian date of the first nest produced each year and the mean date of that first clutch. We also explored the potential effects of monthly precipitation and mean monthly maximum and minimum temperatures during the previous summer, autumn, and winter, when females are undergoing vitellogenesis of the clutch produced the following year (Rollinson et al., 2012). We hypothesized that warmer autumn conditions might contribute to more complete follicle development before winter, and hence, the production of earlier clutches the next spring. In addition, warmer conditions in winter (e.g., Mitchell et al., 2017) or spring (Edge et al., 2017; Janzen et al., 2018) were expected to advance nesting phenology during the following season. We also investigated whether body size, clutch size, or egg size affected nest timing, speculating that larger turtles or those with relatively large eggs or clutches might nest earlier in the season.

Finally, given the deep continental location of our study site, and the finding that climate change is generally progressing more rapidly in continental versus coastal North America (Loarie et al., 2009), we expected turtle nesting at our site to have advanced in time over the course of our long-term study. Furthermore, that advancement should be more evident than for populations farther east.

2 | METHODS

2.1 | Data collection

We monitored nesting turtles that emerged from Gimlet Lake (41°45.24′N, 102°26.12′W), a shallow, sandhill lake on the Crescent Lake National Wildlife Refuge, Garden County, Nebraska, USA (see Iverson & Smith, 1993 for study site description) during 18 (*Chelydra*; Table 2) or 23 years (*Chrysemys*; Table 3). The primary nesting areas were monitored daily during the nesting season (May–July) from at least 06:00 to 22:00 hr by two to five observers. Turtles were weighed, measured (maximum carapace length and maximum plastron length), and marked after nesting. It was not possible to monitor both species for the entire nesting season for every year between 1986 and 2017, but data were available for most years (Tables 2 and 3). In some years, we also sampled *Chelydra* that nested at nearby Island Lake (41°43.95′N, 102°24.16′W).

For each species, we recorded the date in May or June each year that the first gravid female of each species emerged from the lake with the intention of nesting (i.e., gravid and attempted or completed a nest). Additionally, we calculated the mean nest date each year for all emergence dates (even if a nest was not completed that day). For females that failed to complete a nest when first sighted in a given year (e.g., if she was disturbed by Refuge personnel activities) and then nested in a subsequent night during the following several days (i.e., before she could produce a second clutch), her nest was scored as having been deposited on the night she was first observed constructing a nest.

*Chelydra* produced a maximum of one clutch per year at this site (Iverson et al., 1997), but some female *Chrysemys* produced at least three clutches per season (Iverson & Smith, 1993). Hence, for
| Species               | Nest Trait                          | Phenology inversely correlated with temperatures prior to nesting | Earlier nesting phenology through time | Reference                                      |
|----------------------|-------------------------------------|------------------------------------------------------------------|----------------------------------------|-----------------------------------------------|
| Caretta caretta (FL) | Median Nest Date                    | Yes                                                              | Yes                                    | Weishampel et al. (2004)                      |
| Caretta caretta (FL) | Median Nest Date                    | Yes                                                              | Yes                                    | Pike et al. (2006)                           |
| Caretta caretta (FL) | First Day Nesting                   | Yes                                                              | No                                     | Pike et al. (2006)                           |
| Caretta caretta (NC) | First Day Nesting                   | Yes                                                              | No                                     | Hawkes et al. (2007)                         |
| Caretta caretta (Greece) | First Day Nesting | Yes                                                              | Yes                                    | Mazaris et al. (2008), Patel et al. (2016)  |
| Caretta caretta (FL) | First Day Nesting                   | Yes                                                              | NA                                     | Pike (2008)                                  |
| Caretta caretta (Greece) | First Day Nesting | Yes                                                              | NA                                     | Mazaris et al. (2009)                        |
| Caretta caretta (FL) | Median Nest Date                    | Yes                                                              | No                                     | Weishampel et al. (2010)                     |
| Caretta caretta (multiple) | First Day Nesting | Yes                                                              | NA                                     | Mazaris et al. (2012)                        |
| Caretta caretta (FL) | First Day Nesting                   | Yes                                                              | NA                                     | Lamont and Fujisaki (2014)                   |
| Caretta caretta (FL) | Median Nest Date                    | No                                                               | NA                                     | Lamont and Fujisaki (2014)                   |
| Chelonia mydas (FL)  | First Day Nesting/ Median Nest Day  | No                                                               | NA                                     | Pike (2009)                                  |
| Chelonia mydas (FL)  | Median Nest Date                    | Yes                                                              | No                                     | Weishampel et al. (2010)                     |
| Chelonia mydas (East Africa) | Peak Nest Date | Opposite                                                        | No                                     | Dalleau et al. (2012)                        |
| Chelydra serpentina (ON) | First Day Nesting | Yes                                                              | No                                     | Obbard and Brooks (1987), Janzen et al. (2018), Edge et al. (2017) |
| Chelydra serpentina (MI) | First Day Nesting | Yes                                                              | NA                                     | Congdon et al. (1987), Edge et al. (2017)    |
| Chelydra serpentina (NE) | First Day Nesting/ Mean Nesting Day | Yes                                                              | No                                     | Janzen et al. (2018), this paper             |
| Chelydra serpentina (IL) | First Day Nesting | Yes                                                              | Yes                                    | Janzen et al. (2018)                         |
| Chelydra serpentina (SC) | First Day Nesting | No                                                               | No                                     | Janzen et al. (2018)                         |
| Chrysemys picta (ON)  | First Day Nesting                   | Yes                                                              | No                                     | Christens and Bider (1987), Janzen et al. (2018), Edge et al. (2017) |
| Chrysemys picta (MI)  | First Day Nesting/ Median Nest Date | Yes                                                              | NA                                     | Rowe et al. (2003), Edge et al. (2017)       |
| Chrysemys picta (NE)  | First Day Nesting                   | Yes                                                              | No                                     | Iverson and Smith (1993), this paper         |
| Chrysemys picta (IL)  | Median Nest Date                    | Yes                                                              | Yes                                    | Schwanz and Janzen (2008), Janzen et al. (2018) |
| Chrysemys picta (IL; 2 Rivers) | First Day Nesting | Yes                                                              | No                                     | Janzen et al. (2018)                         |
the latter species, mean nest date refers only to the first clutch of the season. The end of production of first clutches for *Chrysemys* was estimated by assuming that at least ten days are required to produce a second clutch by a given female (though usually 12 or more days; Iverson & Smith, 1993) and noting the dates for females known to be depositing their second clutches. The daily frequency of nesting females in the interval between 11 days after the first nest date and the date of the first known second clutch was examined for a gap (or at least a greatly reduced nesting frequency) that was presumed to indicate the transition between first and second clutches. The last day for a first clutch was estimated to be the last day before that gap. We realize that this method is imprecise and likely excludes some females that produced their first clutches, while most of the population was producing second clutches, but our sample sizes should be large enough to minimize this potential bias.

Climate data from July of the previous year through May of the year of nesting were obtained from the NOAA weather station located immediately adjacent to the turtle nesting area (<100 m). We initially compiled a series of climatic variables for the years 1970 through 2017, including mean monthly maximum and minimum daily temperatures, and monthly precipitation. We excluded climate data for the month of June since it overlapped with the nesting season of our study species.

![Species Nest Trait](image)

| Species                      | Nest Trait              | Phenology inversely correlated with temperatures prior to nesting | Earlier nesting phenology through time | Reference                      |
|------------------------------|-------------------------|-----------------------------------------------------------------|----------------------------------------|---------------------------------|
| *Dermochelys coriacea*       | First 10th Percentile  | Opposite                                                        | NA                                     | Neeman et al. (2015)            |
|                               | Nesting                 |                                                                 |                                        |                                 |
| *Emydoidea blandingii*        | First Day Nesting       | Yes                                                             | NA                                     | Congdon et al. (1983)           |
| (MI)                         |                         |                                                                 |                                        |                                 |
| *Emydoidea blandingii*        | 90% Nested              | Yes                                                             | NA                                     | Buckardt et al. (2020)          |
| (IL/WI)                      |                         |                                                                 |                                        |                                 |
| *Gopherus agassizii*          | Mean Gravid Date        | Yes                                                             | NA                                     | Lovich et al. (2012)            |
| (IL)                         |                         |                                                                 |                                        |                                 |
| *Gopherus polyphemus*         | First Day Nesting       | No                                                              | NA                                     | Levengood et al. (2015)         |
| (GA)                         |                         |                                                                 |                                        |                                 |
| *Graptemys geographica*       | First Day Nesting       | Yes                                                             | NA                                     | Nagle and Congdon (2016)        |
| (PA)                         |                         |                                                                 |                                        |                                 |
| *Graptemys ouachitensis*      | First Day Nesting       | Yes                                                             | NA                                     | Geller (2012)                   |
| (WI)                         |                         |                                                                 |                                        |                                 |
| *Malaclemys terrapin*         | First Day Nesting       | Yes                                                             | Yes                                    | Wood et al. (2013)              |
| (NJ)                         |                         |                                                                 |                                        |                                 |
| *Malaclemys terrapin*         | First Day Nesting       | Yes                                                             | No                                     | Janzen et al. (2018)            |
| (MD)                         |                         |                                                                 |                                        |                                 |
| *Trachemys scripta*           | First Day Nesting       | NE                                                              | Yes                                    | Tucker et al. (2008), Janzen et al. (2018) |
| (IL)                         |                         |                                                                 |                                        |                                 |
| *Trachemys scripta*           | First Day Nesting       | No                                                              | No                                     | Janzen et al. (2018)            |
| (SC)                         |                         |                                                                 |                                        |                                 |
| *Kinosternon flavescens*      | First Day Nesting       | No                                                              | No                                     | Janzen et al. (2018)            |
| (NE)                         |                         |                                                                 |                                        |                                 |
| *Kinosternon subrubrum*       | First Day Nesting       | No                                                              | No                                     | Janzen et al. (2018)            |
| (SC)                         |                         |                                                                 |                                        |                                 |
| *Sternotherus odoratus*       | First Day Nesting       | No                                                              | No                                     | Janzen et al. (2018)            |
| (IL)                         |                         |                                                                 |                                        |                                 |

Note: Abbreviations below species names indicate study site. Abbreviation: NA, not assessed.
TABLE 2  First, last, and mean (±1 SD) nesting dates for *Chelydra serpentina* at Crescent Lake National Wildlife Refuge, Garden County, Nebraska by year

| Year | N   | Mean ± SD | First | Last |
|------|-----|-----------|-------|------|
| 1993 | 36  | 170.8 ± 4.5 | 163   | 179  |
| 1994 | 32  | 156.9 ± 3.0 | 152   | 164  |
| 1998 | 14  | 161.4 ± 4.4 | 157   | 174  |
| 1999 | 29  | 165.4 ± 5.0 | 155   | 173  |
| 2000 | 21  | 165.1 ± 8.3 | 151   | 163  |
| 2004 | 15  | 158.5 ± 2.8 | 150   | 162  |
| 2005 | 36  | 164.5 ± 3.4 | 158   | 170  |
| 2006 | 29  | 155.0 ± 2.7 | 152   | 162  |
| 2007 | 91  | 160.0 ± 7.1 | 148   | 165  |
| 2008 | 54  | 171.6 ± 2.3 | 167   | 177  |
| 2009 | 65  | 168.9 ± 2.6 | 162   | 175  |
| 2010 | 52  | 170.0 ± 4.0 | 161   | 178  |
| 2011 | 21  | 176.6 ± 3.5 | 165   | 176  |
| 2012 | 57  | 156.4 ± 2.3 | 153   | 161  |
| 2013 | 49  | 163.2 ± 2.9 | 158   | 168  |
| 2014 | 29  | 163.4 ± 5.1 | 154   | 174  |
| 2015 | 45  | 162.2 ± 3.0 | 158   | 176  |
| 2017 | 25  | 160.4 ± 3.9 | 156   | 171  |

Note: Dates are Julian days (152 = 1 June, except 153 in leap years). Overall means (and number of sample years) at bottom.

### 2.2 Statistical approach

We investigated whether nest deposition (first nests per season, mean nests, and nest dates by individuals) for *Chelydra* and *Chrysemys* was influenced by climatic predictor variables and whether they changed over time during our study period. Additionally, we assessed whether population-level measurements of body size and reproductive variables (carapace length, plastron length, female mass postnesting, mean egg mass, and clutch size) have changed during our study, as these variables can influence timing of nest deposition of an individual.

We analyzed our data two ways. First, we conducted least squares regression analyses to assess long-term trends in climate variables versus year and relationships between temperature and precipitation variables and time (Julian day of nest deposition and years) for the first nest of a season and mean nest dates. Significant p-values for regressions were conservatively adjusted for multiple comparisons by using a sequential Bonferroni correction to (Holm, 1979). We conducted these analyses using Statview software (Abacus Concepts). Second, we assessed relationships between climatic variables and the above-mentioned life history traits on nesting phenology (Julian day of nest deposition) of all nests. For these analyses, we fit linear mixed-effect models via maximum likelihood in R using the package “lme4” (Bates et al., 2015; R Core Team, 2017). We evaluated candidate models using the Akaike information criterion (AIC) where the level of importance was assessed by model weights (w) and overall ranking in the candidate set. For both species, we included carapace length as a fixed effect to account for an individual’s growth and size over time. For *Chrysemys*, we used the random effects of Female ID (identification) to account for individual variation and year to account for differences in sample sizes. For *Chelydra*, we used year as the random effect, but not female ID because 76% of nests could not be associated with a female ID (see below).

To improve model convergence and determine relationships with nest foray dates, we z-standardized the continuous covariates for our mixed-effect model analyses. We examined relationships between our covariates and dropped one of two variables if their Pearson’s correlation coefficient was >|0.70|, with one exception; mean minimum and maximum air temperatures are correlated, but serve important, separate roles in regulating water temperatures and metabolic rates of freshwater turtles (however, minimum and maximum means for a given month were not included in the same models because of collinearity). In total, we had 33 climatic variables for our analyses.

Each weather variable can affect different aspects of reproductive ecology, including timing of emergence from hibernation, food resources, water temperatures, metabolism, basking conditions, and

### TABLE 3  First, last, and mean (±1 SD) nesting dates for *Chrysemys picta* at Crescent Lake National Wildlife Refuge, Garden County, Nebraska by year

| Year | N   | Mean ± SD | First | Last |
|------|-----|-----------|-------|------|
| 1986 | 26  | 155.0 ± 5.2 | 141   | 164  |
| 1988 | 29  | 151.5 ± 4.2 | 144   | 160  |
| 1990 | 40  | 156.5 ± 3.6 | 150   | 163  |
| 1993 | 20  | 154.0 ± 4.9 | 142   | 161  |
| 1994* | 4+  | NA        | 137   | NA   |
| 1998* | 4+  | NA        | 140   | NA   |
| 1999* | 5+  | NA        | 148   | NA   |
| 2000 | 31  | 153.5 ± 5.4 | 143   | 161  |
| 2001 | 39  | 152.5 ± 4.3 | 145   | 159  |
| 2002 | 31  | 155.7 ± 4.2 | 148   | 163  |
| 2003 | 24  | 159.2 ± 5.6 | 149   | 165  |
| 2004 | 31  | 164.2 ± 3.2 | 157   | 167  |
| 2005 | 48  | 177.1 ± 3.3 | 171   | 185  |
| 2006 | 36  | 159.3 ± 3.1 | 151   | 166  |
| 2007 | 56  | 157.8 ± 5.1 | 147   | 169  |
| 2008 | 67  | 172.2 ± 3.8 | 161   | 178  |
| 2009 | 59  | 168.4 ± 4.0 | 157   | 175  |
| 2010 | 52  | 158.9 ± 4.1 | 152   | 167  |
| 2012 | 71  | 146.1 ± 8.4 | 128   | 156  |
| 2013 | 98  | 159.8 ± 3.7 | 153   | 167  |
| 2014 | 82  | 158.5 ± 4.7 | 147   | 166  |
| 2015 | 98  | 167.0 ± 4.6 | 153   | 174  |
| 2017 | 43  | 157.7 ± 3.7 | 151   | 165  |
| Means | 159.2 (20) | 148.5 (23) | 166.6 (20) |

Note: Dates are Julian days (152 = 1 June, except 153 in leap years). Asterisks indicate years when census ended before all first clutches laid. Overall means (and number of sample years) at bottom.
3 | RESULTS

3.1 | Nesting summary

We tallied 705 nesting forays for *Chelydra* and individually identified 230 (23.6%) females, although many of the others were marked but eluded us after nesting by returning to the water before capture. Of the known females, 45 were only recorded on a nesting foray once during the study, but the average number of years that known females emerged to nest was 2.3 (range 1–8). We found no differences in nesting dates, changes in nesting dates over time (years), or body size metrics between Gimlet and Island Lakes, except for clutch size, our top model, which was significantly smaller at Island Lake (Tables S1–S4). However, sample sizes were disproportionate, with 204 nests from Gimlet Lake and 58 from Island Lake, and likely influenced this result. Therefore, we merged datasets of both lakes for climatic variable analyses.

We also recorded 981 total forays for first nests for *Chrysemys* and associated 503 of those to known females (51.3%). Of the known females, 160 were identified on a first clutch nesting foray only once during the study, but the average number of years that identified females emerged on forays for first nests was 1.9 (range 1–10). As the study progressed, clutch size for *Chrysemys* remained unchanged, but there was a significant increase in carapace length of nesting females (Tables S3 and S4).

Julian day of nest deposition was highly variable among years (e.g., Figure 1). The earliest nesting date for *Chelydra* occurred on Julian day 148 (28 May 2007) and the latest first nest was deposited on Julian day 179 (28 June 1993) (Table 2). The earliest nesting date for *Chrysemys* occurred on Julian day 128 (7 May 2012) and the latest first nest was deposited on Julian day 185 (4 July 2005). The average date of the first nest for *Chelydra* across 18 years was Julian day 157 (6 June), and the average date of nest deposition across those 18 years was Julian day 164 (13 June; Table 2). For *Chrysemys*, the average date of the first nest was Julian day 149 (29 May), and the average nest date was Julian day 159 (8 June; Table 3). Date of the first nest and mean date of nesting within years were highly correlated (p < .0001) for both *Chelydra* (R = .82) and *Chrysemys* (R = .93), although the first nesting dates in a given year between these two species were not correlated (N = 17; R = 0.38; p = 0.14) nor were the mean dates (N = 14; R = 0.34; p = 0.24).

3.2 | Climate summary

Annual precipitation at this site averaged 43.3 cm between 1970 and 2017, and wet season (May–June) rainfall averaged 14.9 cm. However, no measure of precipitation (monthly, seasonal, or annual) changed significantly with time over those 48 years (p > 0.17 for all regressions). In contrast, mean annual temperature at our study site has warmed at a rate of about 0.5°C per decade (Figure 2). Mean daily minimum temperatures for every month of the year except February and December increased significantly from 1970 to 2017 (Table 4). However, mean daily maximum temperatures increased significantly only for January (p = 0.037), but only if no adjustment in that p-value was made for multiple comparisons (Table 4). Mean April–May temperature also did not change over that period (Figure 2), although mean September–October temperature increased significantly, by about 0.5°C per decade (Figure 2).

3.3 | Climate effects

Based on our mixed model analysis, variation in the nesting date by year for *Chelydra* was best explained by mean May minimum temperatures (Table 5; Tables S5 and S6), where each degree C increase in mean May minimum temperature advanced the first nesting date by ca. four days, and the mean nest date by ca. two days (Figure 3). In addition, each degree increase in mean maximum December temperature advanced first nesting by 1.5 days and mean nesting date by 1.3 days. Together, these two variables explained 62% of the variation in first nesting date (Annual First Nesting Date = −0.810*Decmax – 2.942*Maymin + 188.585; p = .0007) and 48% of the variation in mean nesting date (Annual Mean Nesting Date = −0.676*Decmax – 2.56*Maymin + 191.019; p = .0079).

For all *Chelydra* nests (mixed model analyses), eight of our climatic variables significantly influenced nest deposition (Tables S5 and S6). Notably, an increase in mean May minimum (6.1 to 9.6°C) and mean May maximum temperatures (16.9 to 24.0°C) were the most influential variables, each advancing nesting from approximately Julian days 169 to 155 and day 170 to 158, respectively. Similarly, both an increase in mean December maximum (−1.7 to 7.4°C) and mean April maximum (11.1 to 18.8°C) temperatures advanced nesting from Julian day 171 to 160 and day 170 to 158, respectively. Conversely, an increase in precipitation during July of the preceding year delayed nesting from Julian day 159 to 170 when precipitation increased from 0.52 to 5.77 cm.

Based on our mixed model analysis, for *Chrysemys*, nesting date by year was best explained by mean September minimum temperatures (Table 5), where each degree increase in mean temperature delayed first nesting date by 1.5 days and mean nesting date by 1.3 days. Furthermore, each degree increase in mean May maximum
temperature advanced first nesting by 1.9 days, but did not significantly affect mean nesting date (Figure 4). Together, these two variables explained 51% of the annual variation in first nesting (Annual First Nesting Date = 0.167*Septmin - 1.482*Maymax + 168.712; $p = .05$) and 55% of the variation in mean nesting (Annual mean Nesting Date = 1.601*Septmin - 0.99*Maymax + 165.136; $p = .04$).

**FIGURE 1** Box-and-whisker plots of Julian day of nest deposition of *Chelydra serpentina* (top; mean of annual means is Julian day 164; $N = 18$) and *Chrysemys picta* (bottom; mean of annual means is Julian day 159; $N = 20$) by year at Crescent Lake National Wildlife Refuge, Garden County, Nebraska.
For all *Chrysemys* nests (mixed model analyses), five of our climatic variables significantly influenced nest deposition (Tables S7 and S8). An increase in either mean February minimum temperature from −15.2 to −3.9°C or mean minimum temperatures in the previous September (from 3.9 to 12.6°C) delayed nesting up to 15 days from Julian day 150 to 165. Similarly, we found that an increase in either October minimum temperatures (−1.9 to 5.6°C) or mean December minimum temperatures in the preceding year (−17.1 to −4.8°C) each delayed nesting from Julian 153 until day 165. Lastly, April precipitation delayed nesting from Julian day 155 to 165 when rainfall increased from 0.44 to 4.3 cm.

3.4 | Life history effects

Body size and reproductive variables (carapace length, plastron length, clutch size, and spent mass of a postnesting female) were all significantly inversely correlated with nesting dates (Tables S9 and S10). We dropped spent body mass and plastron length as predictor variables for both species because of high correlation coefficients (>|.87|) with carapace length. For *Chelydra*, females with a carapace length of 225 mm were predicted to nest on Julian day 167, whereas a larger female with a carapace length of 395 mm was predicted to nest on day 162. Similarly, females of *Chrysemys* with a carapace length of 150 mm were predicted to nest on Julian day 162, whereas a larger female with a carapace length of 206 mm was predicted to nest on Julian day 157.

3.5 | Temporal effects

Least squares analyses of first and mean nesting dates for both *Chelydra* and *Chrysemys* revealed no change over time during our study period (Figure 5). Similarly, from our mixed model analyses, we

| TABLE 4 Correlations between year (1970–2017; N = 48) and mean monthly minimum and maximum temperatures at Crescent Lake National Wildlife Refuge, Garden County, Nebraska |
| --- |
| **Month** | **Minimum** | **Maximum** |
|   |   | **R** | **p** | **R** | **p** |
| Jan | .626 | <.0001 | .305 | .037 |
| Feb | .322 | .025 | .220 | .132 |
| Mar | .574 | <.0001 | .179 | .225 |
| Apr | .501 | <.0001 | .179 | .224 |
| May | .522 | <.0001 | .210 | .152 |
| Jun | .601 | <.0001 | .080 | .588 |
| Jul | .727 | <.0001 | .153 | .300 |
| Aug | .675 | <.0001 | .036 | .806 |
| Sep | .720 | <.0001 | .049 | .740 |
| Oct | .648 | <.0001 | .231 | .115 |
| Nov | .594 | <.0001 | .177 | .230 |
| Dec | .393 | .006 | .095 | .519 |

Note: Regression coefficients (R) are followed by p-values. Only p-values < .0001 are significant after sequential Bonferroni adjustment for multiple comparisons.
found no changes in nesting dates over time for *Chelydra* ($\beta = 0.04$, SE = 0.18, 95% CI = −0.31, 0.40) or *Chrysemys* ($\beta = 0.26$, SE = 0.17, 95% CI = −0.08, 0.61). Mean annual carapace length did not change over time for *Chelydra* ($R = −.46; p = .07; n = 18$), a pattern supported in our mixed model analyses (Tables S3 and S4). However, for *Chrysemys*, mean annual carapace length decreased over time (Figure 6). In contrast, our mixed model analysis suggested that carapace length in *Chrysemys* increased over time (Tables S3 and S4), although the latter results are complicated by the uneven annual sample sizes (Table 3) and the clear trend of an increase in body size over the last third of the study (Figure 6) during the years with large sample sizes (Table 3). Furthermore, the relationship between body size and nest date over the full study period in *Chrysemys* was confounded by population-level demographic changes due to variation in female mortality and nest survivorship (Figure 6; see discussion for details).

4 | DISCUSSION

Spring temperatures are generally inversely correlated with nest phenology in turtles (Table 1), with 27 of 38 previous studies (representing 15 species) demonstrating this pattern. Only two studies (both for marine turtles) exhibited the opposite trend (see Table 1), but both of those studies used different measures of nest timing (peak nesting date; first 10th percentile) than all other studies, including ours.

### TABLE 5 Spring climate variables correlated with nest date for two turtle species at Crescent Lake National Wildlife Refuge, Garden County, Nebraska

| Climate Variable | Chelydra | Chrysemys |
|------------------|----------|-----------|
| May min | .0006* | .05 |
| May max | .004 | .03 |
| October min | .005 | .05 |
| December max | .01 |
| April max | .05 |
| March max | .05 |
| January max | .05 |
| April min | .03 |
| December min | .02 |
| December max | .06 |
| July rain | .04 |

**Note:** *p*-Values appear below variable name, and all listed climate variables are means. Positive correlations are bolded; all others are inverse relationships. *p*-Values with asterisks are significant after sequential Bonferroni adjustment for multiple comparisons (separately for temperature and precipitation). Sample sizes and sample years in Table 1.

At our site, warmer springs also advanced nesting in *Chelydra*, as did increased mean December maximum temperatures. The mechanisms driving this pattern likely operate through thermoregulation or local food chain productivity (Schwanz & Janzen, 2008). Increased local environmental temperatures during the winter and spring presumably permit earlier emergence from hibernation, increased metabolic rates (e.g., via basking), and accelerated vitellogenesis, ovulation, and egg shelling, all of which would drive earlier nesting (Mitchell et al., 2017; Obbard & Brooks, 1987). Similarly, an increase in local food chain productivity due to increased temperatures could also provide more resources necessary to speed up reproductive demands, although this mechanism is probably secondary to thermoregulatory affects. However, nest timing in *Chrysemys* was not strongly influenced by spring temperatures, as predicted, but rather, delayed by warm temperatures in the fall and winter.

Rollinson et al. (2012) demonstrated that snapping turtles complete vitellogenesis primarily by the end of the previous fall, whereas for *Chrysemys* the process occurs both in the fall and spring (see also Callard et al., 1978). Our data suggest that vitellogenesis in *Chelydra* may in fact continue into December, even though air temperatures are quite cold that month (long-term mean daily maximum = 3.8°C; minimum, −10.1°C). However, for *Chrysemys*, warm temperatures in early fall might be expected to accelerate vitellogenesis and hence result in early nesting during the following summer (the opposite of our findings). Perhaps the metabolic costs of a warm fall exceed the benefits to vitellogenesis in *Chrysemys*. Part of the difficulty in explaining these novel patterns is our lack of water temperature data,
July temperatures were not related to nest timing in Chelydra serpentina nesting the following year by slowing vitellogenesis, perhaps can therefore only speculate that increased precipitation in July de-

\[ y = -3.969x + 187.225, r^2 = .531 \]

\[ y = -1.899 + 250.642, R^2 = .412, \text{and } p = .004. \]

\[ y = 1.503x + 134.65, r^2 = 0.171 \]

\[ y = -1.94x + 189.014, r^2 = 0.165 \]

\[ y = -1.482x + 173.611, r^2 = 0.411 \]

\[ y = -1.260x + 177.987, R^2 = .311, \text{and } p = .0162 \]

\[ y = -1.740x + 143.330, R^2 = .25, \text{and } p = .026, \text{and} \]

\[ y = -1.315x + 186.523, R^2 = .10, \text{and } p = .18, \text{respectively} \]

FIGURE 3 First Julian nesting date of Chelydra serpentina from 1993 to 2017 at Crescent Lake National Wildlife Refuge, Garden County, Nebraska, in response to mean May minimum daily temperature (TOP) and mean December maximum daily temperature (BOTTOM). For May temperatures, regression is statistically significant before and after sequential Bonferroni adjustment \((p = .0006; \text{see Table 4); for mean Julian nesting date } y = -1.899x + 250.642, R^2 = .412, \text{and } p = .004. \)

For December temperatures, regression is statistically significant before but not after sequential Bonferroni adjustment \((p = .0041; \text{see Table 4); for mean Julian nesting date } y = -1.260x + 177.987, R^2 = .311, \text{and } p = .0162 \)

which presumably lag air temperatures. Interpreting differences between the two species is further complicated by the much higher propensity of Chrysemys to bask aerially compared to Chelydra. Indeed, basking in Chrysemys may explain why nest timing in that species was so much less predictable by air temperatures than in Chelydra.

We were surprised to find that increased precipitation in July was correlated with a delay in nesting in Chelydra, over ten months later. Although previous studies have examined the impact of precipitation on nest timing during the nesting season (see review in Czaja et al., 2018), no study has examined the effects on nest timing of precipitation outside the nesting season. Increased precipitation in July at our site was correlated with colder mean daily July maximum temperatures \((R = .38; p = .007; N = 48 \text{ years}), \text{but average July temperatures were not related to nest timing in Chelydra. We can therefore only speculate that increased precipitation in July delays nesting the following year by slowing vitellogenesis, perhaps via its effect on lowering water temperatures. The importance of April precipitation in delaying nesting in Chrysemys was surprising. We suspect that high precipitation in April reduces basking opportunities and decreases water temperatures, both of which would be expected to delay nesting in Chrysemys, which must complete vitellogenesis in the spring (Rollinson et al., 2012).

Our analyses revealed that larger female Chelydra and Chrysemys tended to nest earlier in the season than smaller females. Because clutch size and egg size are correlated with body size in both species (Iverson et al., 1997; Iverson & Smith, 1993), early nests included more and bigger eggs. Earlier nesting by larger female turtles has previously been reported for Graptemys geographica in Pennsylvania (Nagle & Congdon, 2016). These results suggest that the size class distribution of a population can impact its nesting phenology, complicating phenology comparisons across years, populations, and species.

Although mean body size of nesting female Chelydra did not vary over time at our study site \((R = -.46; p = .07), \text{mean annual carapace length of nesting female Chrysemys did decrease significantly} \)
of nest protection that began in 1993 and a major depredation period in 2003 and 2004 (evident in Figure 6; Tables S3 and S4). This decrease was most likely the result of nesting female Chrysemys picta (open dots; N = 23) by year at Crescent Lake National Wildlife Refuge, Garden County, Nebraska. None of the relationships was significant (TOP: R = -.13, p = .60; R = .32, p = .14, respectively; BOTTOM: R = .02, p = .93; R = .33, p = .16, respectively)

Because both species in our study demonstrated size-related variation in nesting dates, future studies of turtle nesting phenology should consider the effects of body size as a complicating factor when making comparisons over time and space, as changes in nesting phenology may actually be related to or compounded by changes in population structure. For example, a population of turtles experiencing a decline in body size over time (e.g., via increased poaching or predation) would be expected to exhibit a pattern of delayed nesting even if the climate had not changed. In the future, failure to account for changes in size class distributions over time in turtle nesting phenology studies may mask or artificially suggest temporal patterns.

Climate change over the last five decades has produced warmer temperatures overall at our site, with the greatest impact being a noticeable increase in nighttime minimum temperatures (Table 4). It was also our subjective impression that nighttime skies grew increasingly hazy over the study period, and although the cause(s) are not yet clear, the increasingly cloudy skies and warmer nights were likely related. Despite the significant overall warming at our site over the past several decades and a clear inverse relationship between spring temperatures and nesting timing, nesting phenology in at least Chelydra has not changed between 1993 and 2017 at our site. This is likely at least in part a reflection of the fact that spring temperatures at our site have not changed over that period (Table 5; Figure 3), even though annual temperatures have (Figure 2). The meteorological reasons for this spring difference are not yet evident. In any case, should spring day-time temperatures eventually warm at our site, the nest phenology of at least these two turtle species in western Nebraska will likely be affected.

Of the 38 studies summarized in Table 1, 23 (nine species) evaluated nesting phenology through time, and surprisingly, only seven of them (five species) documented the expected temporal shift to
earlier nesting. However, other studies of each of the latter five species demonstrated no significant shift in phenology. Because climate change has generally been more rapid in continental versus coastal areas of North America (Loarie et al., 2009), we expected continental turtle populations to show more advanced nest dates than those closer to the coast. For example, Trachemys scripta in South Carolina exhibited no shift over time, whereas in Illinois it did (Table 1). However, among the four studied populations of Chelydra, three (Ontario, Canada, and Nebraska and South Carolina, USA) exhibited no shift in nest timing, while that in Illinois did (Table 1). Similarly, among four populations of Chrysemys, (Ontario, Canada, Nebraska and southern Illinois, USA) exhibited no shift, but that in northern Illinois did. Clearly, the observed geographic variation in nest timing in Chelydra and Chrysemys does not match the predicted continental scenario. Interestingly, the only two populations of these two taxa exhibiting a temporal change are riparian, whereas those with no change are all lentic systems. Whether this is an important factor explaining these patterns will require additional fieldwork, including the collection of water temperature data.

Regrettfully, water temperatures were not recorded during our study, since they might be expected to be better predictors of nesting dates (e.g., as sea surface temperatures have been for marine turtles; Table 1). However, even those data would be complicated by differences in habitat use by our study species. In our experience, Chelydra seems to occur in shallow (warmer?) water and does limited aerial basking, whereas Chrysemys seem to inhabit deeper water and exhibits extensive aerial basking (see also Ernst & Lovich, 2009). Similarly, more detailed analyses of temperatures (water and air) beyond simply monthly means (especially during the spring and fall temperature windows that are most highly correlated with nest timing) might clarify the mechanism for the relationship between temperature and nest timing more precisely (e.g., see Edge et al., 2017; Schwanz & Janzen, 2008).

It is also possible that the inability to detect a change in the nesting phenology at our site, as well as many other sites in Table 1, could be due to the stalling in increasing global mean surface temperatures from the late 1900s through the 2000s known as the "climate change hiatus" (e.g., Kosaka & Xie, 2013). Steady rather than increasing temperatures during that period could explain the lack of statistically significant change in nesting phenology in studies including data collected during that period, although this hiatus is not evident in our climate histories (Figures 2–4).

Although our study focused on the proximate effects of climate on turtle nesting phenology, its impacts on other areas of life history remain poorly studied (Butler, 2019). For example, climate change is likely to affect the length of the nesting season (e.g., Hedrick et al., 2018; Lamont & Fujisaki, 2014; Pike et al., 2006); internest intervals (Hays et al., 2002); clutch frequency (Mazaris et al., 2012; Tucker et al., 2008); egg and clutch size (Hedrick et al., 2018; Lamont & Fujisaki, 2014; Mazaris et al., 2012); survival of early nests and early nesting females (Mazaris et al., 2009; Schofield et al., 2009); hatching success (Hawkes et al., 2007); posthatching survival (especially in species like Chrysemys picta with hatchlings that overwinter terrestrially; Costanzo et al., 1995; Muir et al., 2012); juvenile growth rates (Avery et al., 1993; Gibbons, 1970); and population sex ratios via temperature-dependent sex determination (Schwanz & Janzen, 2008; Tucker et al., 2008). None of these potential effects have been examined at our site, nor have the fitness costs of earlier nesting.

As previously noted, only 7 of 23 studies that evaluated turtle nesting phenology over time have documented that nest dates have advanced over recent decades. Clearly, the collection of more data is necessary before we can generalize that climate change has altered nesting phenology in nonmarine turtles. Part of the problem is that such studies depend on demanding long-term studies. For example, of the 38 studies reviewed in Table 1, only three field sites (Ontario, Canada, and Illinois and Nebraska, USA) have recorded nesting histories exceeding 20 years (see also Janzen et al., 2018). Thus, our ability to detect long-term changes in nest timing in many turtle populations may be constrained by sample size, speaks to the value of long-term studies, and argues for the continuation of those currently in place.

Furthermore, among sea turtles, most of the nesting phenology research done to date has focused on only two species (with complicated migratory cycles), while most of the work done on nonmarine turtles has focused on Chelydra and Chrysemys (Table 1). Studies of other taxa, especially those in regions other than temperate North America, are sorely needed.

While our study provides insight into how morphological and environmental variability influences nesting dates of Chelydra and Chrysemys in Nebraska, these effects are complex and at times likely synergistic or antagonistic. Regardless, this and other studies demonstrate that turtles exhibit extensive plasticity in their reproductive biology. For example, from this study, the range of dates of first nesting varied across years by 19 days for Chelydra and 43 days for Chrysemys. However, it remains to be seen whether this plasticity is sufficient to overcome the threat of additional significant climate change, particularly warming (McGaugh et al., 2010; Refsnider & Janzen, 2016; Schwanz & Janzen, 2008; Urban et al., 2014).

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CONFLICT OF INTEREST
None declared.

AUTHOR CONTRIBUTION
Ashley R. Hedrick: Conceptualization (equal); Investigation (equal); Writing-original draft (equal); Writing-review & editing (equal).
Daniel U. Greene: Data curation (equal); Software (equal); Writing-original draft (equal); Writing-review & editing (equal).
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John B. Iverson: Conceptualization (equal); Investigation (equal); Software (equal); Writing-original draft (equal); Writing-review & editing (equal).

DATA AVAILABILITY STATEMENT
The original phenology data files for Chelydra and Chrysemys are available from the DRYAD Digital Repository: https://doi.org/10.5061/dryad.2v6wpzkn.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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