Breeding on the leading edge of a northward range expansion: differences in morphology and the stress response in the arctic Gambel’s white-crowned sparrow

Jesse S. Krause1 · Helen E. Chmura1 · Jonathan H. Pérez1 · Lisa N. Quach1 · Ashley Asmus2 · Karen R. Word1 · Michaela A. McGuigan1 · Shannan K. Sweet4 · Simone L. Meddle3 · Laura Gough2 · Natalie Boelman4 · John C. Wingfield1

Received: 14 January 2015 / Accepted: 2 September 2015 / Published online: 1 October 2015
© The Author(s) 2015. This article is published with open access at Springerlink.com

Abstract Individuals at the forefront of a range shift are likely to exhibit phenotypic traits that distinguish them from the population breeding within the historic range. Recent studies have examined morphological, physiological and behavioral phenotypes of individuals at the edge of their range. Several studies have found differences in the hypothalamic–pituitary–adrenal (HPA) axis activity in response to acute restraint stress in individuals at the range limits. HPA axis activation leads to elevations in glucocorticoids that regulate physiology and behavior. Here we compare the hormonal profiles and morphometrics from Gambel’s white-crowned sparrows (Zonotrichia leucophrys gambelii) breeding at the northern limit of the population’s range to those birds breeding within the historic population range. Birds breeding at the northern limit experienced a harsher environment with colder temperatures; however, we found no differences in arthropod prey biomass between the northern limit and more southern (historic) sites. Males at the northern limit had higher body condition scores (mass corrected for body size) compared to individuals within the historic range, but no differences were found in beak and tarsus lengths, wing chord, muscle profile or fat stores. In males during the pre-parental stage, before breeding commenced, HPA axis activity was elevated in birds at the northern limit, but no differences were found during the parental or molt stages. Females showed no differences in HPA axis activity during the parental stage. This study suggests that “pioneering” individuals at the limits of their breeding range exhibit physiology and morphology that are distinct from individuals within the historic range.

Keywords Adrenocortical · Corticosterone · Hypothalamic–pituitary–adrenal axis · Glucocorticoids · Morphometrics · Climate change · Allostatic load

Introduction

The geographical range of a species is determined by the ecological niche where evolution has selected the traits that maximize fitness (MacArthur 1972). Species ranges are often highly plastic, shifting or expanding to match spatial shifts of their niche (Sexton et al. 2009). As an individual moves to the range margins, the specialized traits may no longer be optimal for the environment that is encountered because habitat quality has declined. (Sexton et al. 2009). Changes in species’ geographic range are one of the most commonly reported responses to climate change. A meta-analysis of 1700 plant and animal species indicates that ranges are shifting at a rate of 6.1 km per decade towards the poles (Parmesan and Yohe 2003). Northward range
expansion may expose individuals to more spatially disparate habitats, shifts in phenology and abundance of prey items, novel parasites and predators, and to thermal challenges that have not previously been experienced.

Individuals at the leading edge of a range expansion may exhibit distinct behavioral, morphological, and physiological traits that enhance fitness in a challenging environment. There have been many behavioral traits associated with a vertebrate’s ability to colonize a new area that include aggression, sociability, boldness, and exploration (Cote et al. 2010). Morphological characteristics, such as body size (Bowler and Benton 2005), are also commonly reported as observed changes in range expanding individuals, and evidence exists for both larger and smaller individuals moving into new areas (Bowler and Benton 2005). In addition, circulating levels of glucocorticoids are commonly measured because these hormones regulate physiology and behavior and are thought to help these individuals in their novel environment (Addis et al. 2011; Dunlap and Wingfield 1995; Liebl and Martin 2012).

Modulation of glucocorticoid secretion by the hypothalamic–pituitary–adrenal (HPA) axis may be particularly important in vertebrate populations undergoing a range expansion because it is a highly plastic trait that is profoundly influenced by the environment. Stressors in the environment, such as storms, predation attempts, food shortages, and territorial disputes induce increases in glucocorticoids that regulate changes in physiology and behavior to enhance fitness (Sapolsky et al. 2000; Wingfield and Sapolsky 2003). For birds, the main glucocorticoid, corticosterone, is important for energetics (Astheimer et al. 2000; Landys et al. 2004), foraging (Angelier et al. 2008; Astheimer et al. 1992), dispersal (Silverin 1997) and escape behavior (Breuner and Hahn 2003). Basal levels of corticosterone are important for regulating metabolic processes (Sapolsky et al. 2000).

Acute activation of the HPA axis, in response to stress, is beneficial to the organism, but chronic activation can have deleterious effects (Sapolsky et al. 2000; Wingfield and Sapolsky 2003). Since glucocorticoids are an important mediator of life history trade-offs, either under- or over-production of these steroids could have significant fitness consequences (i.e., the decision to abandon a nest) (Angelier and Wingfield 2013).

Existing evidence indicates that birds breeding at the edge of a population’s range show increased HPA activity, as well as higher variance in activity in response to acute restraint stress, compared to individuals within the historic range (Addis et al. 2011; Liebl and Martin 2012; Walker et al. 2015; Wingfield et al. 1995a). Range expansion into a new territory with novel characteristics such as harsher climate, interactions with novel species, altered habitat characteristics such as plant species richness, abundance, and density; and altered food availability and quality; may make elevated HPA axis activity beneficial for coping with these environmental stressors (Wingfield et al. 2015).

The Arctic is changing rapidly in response to climate change and provides a unique system to investigate characteristics of individuals at the leading edge of a range expansion. The expansion of deciduous woody shrubs, mainly willows (Salix spp.), alders (Alnus spp.), and birch (Betula spp.) (Sturm et al. 2001; Tape et al. 2006), has transformed the landscape enabling colonization by higher trophic level species previously limited by habitat quality. One colonist of this new habitat is a migratory bird, the Gambel’s white-crowned sparrow (Zonotrichia leucophrys gambelii), which winters in the North American Southwest and migrates to its breeding grounds that span a large geographical area extending from Southern to Northern Canada and Alaska (Chilton et al. 1995). Based on observations by our laboratory over the last 27 years, Gambel’s white-crowned sparrows have tracked the northward expansion of woody shrubs, north of the Brooks Mountain Range in Alaska, and thus have expanded their breeding distribution on the North Slope towards the Arctic Ocean (Wingfield, personal observation). This is corroborated by Breeding Bird Survey data from the US Geological Survey that indicate that the number of birds breeding north of the Brooks Range has increased in the past 8 years (Sauer et al. 2014) (Supplemental Fig. 1).

To date, little is known about any adaptive traits that will be beneficial to arctic breeding songbirds colonizing new areas. Here we established whether individuals at the forefront of the range exhibit physiological and morphological characteristics that are distinct from those of individuals in the historic range as a consequence of differing environments. Potential differences in physiology and morphology could be a consequence of either phenotypic plasticity or genetic adaptation. We hypothesized: (1) that birds breeding at the northern limit would experience harsher environments characterized by colder temperatures, higher wind speeds, and lower arthropod biomass (an indicator of food availability). As a consequence of the harsher environment, we hypothesized: (2) that HPA activity in response to acute restraint stress, as well as morphological measures, would be distinct between birds at the edge of the population’s range compared to birds within the historic range, and that these parameters differ among stages of breeding. We predicted that birds at the range limits would show greater basal and stress-induced levels of glucocorticoids and enhanced body condition that include increased mass, fat stores and flight muscle profile when compared to birds within the historic range for each stage of breeding. At both the range limit and historic range, we predicted that HPA axis activity would be highest during the pre-parental stage when conditions are harshest, and that birds would have
reduced fat stores, muscle and mass compared to the parental and molt stages.

Materials and methods

Study sites and species

Our field operation was based at Toolik Lake Field Station at mile point (MP) 284 on the James Dalton highway in the foothills of the Brooks Mountain Range on the North Slope of Alaska, USA (69°38′N, 149°36′W). The James Dalton Highway starts at MP 1 in the south and terminates at 414 in the north at Deadhorse; this creates an ideal latitudinal transect for sample collection. Climate, bird, and arthropod data were collected at the range-limit site, MP 386 (Franklin Bluffs; 69°50′N, 149°45′W), as well as at the four historic range sites, MP 303 (Sagavanirktok Department of Transportation; 68°45′N, 148°53′W), MP 291 (Innavait Creek; 68°37′N, 149°18′W), MP 284 (Toolik Lake Field Station), and MP 265 (Roche Mountonee; 68°22′N, 149°19′W) (Fig. 1). The distance from the Toolik Lake Research Station to the Franklin Bluffs is 102 miles (164 km) by road while the geographical distance is 86 miles (122 km).

Breeding Bird Survey data indicate that the number of Gambel’s white-crowned sparrows encountered along a point-count transect greatly diminishes north of the Brooks Range in Alaska, and although these numbers seem to fluctuate between years, they are currently increasing (Sauer et al. 2014) (Supplemental Fig. 1). The decline in bird numbers from south of the Brooks Range to the northern limit of the range can be attributed to reductions in suitable habitat as shrubs become patchier and more disparate. In 1987, the breeding range extended north to MP 303 on the James Dalton Highway (Fig. 1). In 2005, birds were present at MP 334 (Happy Valley; 69°09′N, 148°49′W) although no evidence of breeding was found. However in 2011, birds were found breeding further north around MP 386 and two non-breeding males were found 16 km further north at MP 396 (69°58′N, 148°42′W). The edge of the population’s range has moved a geographical distance of 75 miles (120 km) north in 24 years with a population estimated to be between 24 and 40 birds, depending on the sampling year, which have successfully reproduced (Wingfield, unpublished data). In the present study, birds inhabiting areas colonized after 2005 (MP 386) are termed “range limit” and birds found in existing range (MP 266–303) are termed “historic range”.

Gambel’s white-crowned sparrow is a socially monogamous, long-distance migrant that raises a single brood during the short arctic summer (Chilton et al. 1995). Birds typically arrive on their breeding grounds in mid-May when snow cover is abundant and they remain in small flocks until enough snow melts to permit territory establishment (Wingfield et al. 2004). Females are responsible for nest
construction which can last from 1 to 3 days (Oakeson 1954). A typical clutch consists of four to five eggs with one egg laid per day (Norment 1992; Oakeson 1954). The brood patch may develop up to 5 days prior to lay but typically becomes fully edematous at the onset of incubation (Bailey 1952). Mean clutch initiation dates within the historic range were 1 June in 2011 and 3 June in 2012 (Wingfield et al., unpublished data). The eggs are incubated for approximately 12 days until hatch and nestlings remain in the nest approximately 9 days until fledge (Norment 1992). Fledglings are fed by the parents until they are fully independent. Molt in adults typically starts after nestlings fledge, at the beginning of July and last until mid- to late August (Morton et al. 1969). Fall departure from the North Slope of Alaska can occur from late August to early September (Ramenofsky and Chmura, unpublished data).

Gambel’s white-crowned sparrows prefer to breed in a habitat that contains shrubs, grass, and a nearby water source (Boelman et al. 2015; Chilton et al. 1995). Nests are typically found either in Betula or Salix species ranging in height from 20 cm to ~1 m (Boelman et al. 2015). Thus a certain threshold of shrub density, complexity, and height appears to be a requisite for suitable nesting habitat (Boelman et al. 2015; Norment 1993; Oakeson 1954). All blood samples were collected from the birds in accordance with University of California Davis Institutional Animal Care and Use Committee approval under protocol 17812.

Climatic variables

Weather data were collected by meteorological stations at one range-limit site (Kane 2013) and four historic range sites (Environmental Data Center 2013). Our analyses focused on minimum temperatures, wind speed, and wind chill as indicators of environmental harshness at three different stages of the breeding season which included pre-parental, parental, and molt. Wind chill was calculated because for a set temperature, increases in wind velocity enhance cooling rates through convective heat loss and as a result lower the apparent temperature. Wind chill was calculated with the equation \( T_{WC} = 13.12 + 0.6215 T_a - 11.37V^{0.16} + 0.3965T_a V^{0.16} \) where \( T_{WC} \) is wind chill, \( T_a \) is ambient temperature and \( V \) is wind velocity (Osczewsjk 1995). Snow cover was less than 20% at the onset of the study and as a result was not included in our analyses.

Arthropod biomass

Canopy-dwelling arthropod biomass was measured weekly during the 2012 field season using a standard sweep net protocol to capture arthropods (Boelman et al. 2015). At each site, an area characterized by dwarf deciduous shrub tundra was designated for sampling and an iron rod was driven into the ground to serve as a central point for a 100-m transect. After generating a random bearing for the transect, by spinning a pen, a total of ten sweep net samples were collected at 10-m intervals along the transect. Individual samples were collected by making 20 horizontal passes along the vegetation. Individual sample contents were placed into a plastic bag containing a 1-cm² piece of Hot Shot No-pest strip to euthanize the arthropods, and then put in a −20 °C freezer upon return to the laboratory. Arthropods were then separated from plant material, placed into Petri dishes, dried for 24 h at 40 °C, and then weighed to the nearest milligram.

Bird morphometrics and HPA axis activity

Birds were sampled at three distinct stages across the breeding season in 2011 and 2012. The pre-parental stage lasted from arrival on territories, as determined by the return of color-banded individuals to known territories, to average clutch initiation date and was classified using mean clutch initiation dates from historic sites in conjunction with brood patch development from both historic and range-limit sites. During the pre-parental stage, developing brood patches were recorded in females at both range-limit and historic sites suggesting that breeding phenology was similar between the two sites. In addition, no edematous brood patches were recorded suggesting that incubation had not commenced (see Bailey 1952 for description). The parental phase included incubation and feeding young until molt was initiated, at which point birds were classified as no longer in breeding condition. Samples for the pre-parental stage, parental, and molt were collected on 20 May to 4 June, 23 June to 11 July, and 7–28 July, respectively.

A standard capture restraint protocol was used to measure HPA axis activity (Wingfield et al. 1992). A total of 57 birds were captured using Japanese mist nets in conjunction with song playback. Following venipuncture of the wing vein with a 26-gauge needle, a baseline blood sample was collected into 75-µL microhematocrit tubes within 3 min of capture. Between blood sampling at 10-, 30- and 60-min post-capture the bird was placed into an opaque cloth bag. Samples from females were also collected but we only had a sufficient number to include birds from the parental stage. After blood collection, each bird was banded with an aluminum US Geological Survey band along with a unique combination of color bands for identification in the field. Prior to release, morphometrics of wing chord, tarsus length, and beak length (nares to tip) were measured using calipers (±0.1 mm); and mass was measured using a Pesola spring scale (±0.1 g). Composite fat stores (furcular and abdominal) on a scale from 0 (lean) to 5 (fat) (Wingfield and Farner 1978) and muscle profile on a scale from 0 to 3 (Bairlein and Simons 1995) were also recorded. The blood was kept...
chilled for 6–12 h until processing. Blood samples were centrifuged at 10,000 r.p.m. for 5 min to separate the plasma. Plasma was aspirated with a fixed-needle Hamilton syringe, placed into a microcentrifuge tube, and stored at −80 °C. At the end of the field season, all samples were transported frozen on dry ice to the University of California, Davis.

**Radioimmunoassay for corticosterone**

A radioimmunoassay was used to quantify plasma concentrations of corticosterone (Wingfield et al. 1992). Tritiated corticosterone was purchased from Perkin Elmer (NET399250UC) and corticosterone antibody from Esoterix (B3-163). Samples were assayed in duplicate and counted for 10 min or to within 2% accuracy on a Beckman LS 6500 counter. Mean recoveries were 84.57%. Inter- and intra-assay variations were 15.4 and 10.7%, respectively.

**Statistical analyses**

Statistical analyses were performed using JMP 11 (1989-2007; SAS Institute, Cary, NC) and in R using package Pracma [Borchers (2015), practical numeric math functions (Pracma); R Package version 1.6.4. http://CRAN.R-project.org/package=pracma]. Climatic variables including minimum temperature, wind chill and mean wind speed were analyzed using a two-way factorial ANOVA to investigate the effects of breeding stage (pre-parental, parental and molt) and range (limit and historic). Post hoc tests were performed using Tukey’s honest significant difference (HSD) test. Sweep net data were analyzed with a repeated-measures ANOVA to investigate the effects of range and sampling week on arthropod biomass.

Tarsus, beak, and wing chord were compared using a t-test to investigate morphometric differences between historic and range-limit birds. In order to correct for body size effects on mass, we conducted a principal component (PC) analysis using beak and tarsus lengths and wing chord to create a structural size index (PC1). The residuals from the regression of mass against the PC1 were used to create a corrected body condition index. Morphometrics including fat stores, body index, and muscle profile were analyzed using a two-way factorial ANOVA including the main effects of range and breeding stage and their interaction. Tukey’s HSD post hoc analyses were used following a significant ANOVA to determine differences among groups.

Hormonal data were analyzed using two statistical approaches following log transformation. All observations within a life history stage were from unique individuals while at the range-limit site four birds were sampled at two different life history stages. Multivariate repeated-measures ANOVA was used to evaluate the effects of range (historic vs. range limit), stress response (effect of acute restraint handling), and their interaction on corticosterone at baseline, 10, 30 and 60 min post-sample at each breeding stage. Data were checked for sphericity, and when not met, the Greenhouse–Geisser correction was used. Differences in integrated corticosterone measurements were tested using a general linear model with the effects of range, breeding stage and their interaction. Post hoc analyses were performed using a Student’s t-test to compare integrated corticosterone within a breeding stage. Integrated corticosterone was calculated by using the trapezoidal rule between time points and subtracting baseline. Data are reported as the mean ± SEM.

**Results**

**Climatic variables**

Temperatures were different across stages of breeding ($F_{2,2} = 190.66, P < 0.01$; Fig. 2a). There was a main effect of range on minimum temperature ($F_{1,1} = 48.15, P < 0.01$) and mean wind speed ($F_{1,1} = 198.34, P < 0.01$). The interaction of range and breeding stage was significant for minimum temperature ($F_{2,2} = 10.11, P < 0.01$) and mean wind speed ($F_{2,2} = 5.60, P = 0.003$). Temperatures were colder at range limit than historic range during the pre-parental ($t = 5.98, P < 0.001$) and parental ($t = 3.72, P < 0.001$) stages but not during molt ($t = 1.90, P < 0.06$). Tukey’s post hoc analyses indicated that mean wind speed was higher at the range limit at each stage (Supplemental Fig. 2). When investigating effects of wind chill, the magnitude of the difference in temperatures between historic range and range limit were significantly increased ($F_{1,1} = 132.38, P < 0.001$) at the pre-parental ($t = 23.01, P < 0.01$), parental ($t = 13.85, P < 0.01$) and molt ($t = 13.96, P < 0.01$; Fig. 2a) stages.

**Arthropod biomass**

Dry arthropod biomass significantly increased as the season progressed for both historic and range-limit sites ($F_{4,45} = 36.11, P < 0.01$; Fig. 2b). However, biomass was not different between the two locations ($F_{4,48} = 0.21, P = 0.64$). The interaction between range and sampling week was significant ($F_{4,45} = 58.69, P = 0.05$). Tukey’s post hoc analyses indicated that biomass was higher at the historic site compared to the range-limit site during the week of 18 June 2012 (day of year 170).

**Bird morphometrics**

**Body size**

Measures of wing chord (male, $t = 1.05, P = 0.26$; female, $t = 0.94, P = 0.35$), tarsus (male, $t = 1.34$,
Female body condition index was higher at the range-limit site ($F_{1,44} = 5.62$, $P = 0.02$; Fig. 3a). Male body condition index did not change across the breeding season ($F_{2,44} = 0.11$, $P = 0.89$) and the interaction of breeding stage and range was not significant ($F_{2,44} = 0.18$, $P = 0.83$). For females the body index during the parental stage was not different between range-limit and historic sites ($t = -1.21$, $P = 0.26$; Fig. 3b).
Fat stores

In males, fat stores were different across breeding stages ($F_{2,44} = 3.74, P = 0.03$; Fig. 3c). Males stored less fat during the pre-parental compared to the parental ($t = 5.84, P < 0.001$) and molt ($t = 4.63, P < 0.001$) stages. In males, the effect of range ($F_{1,44} = 0.01, P = 0.89$) and the interaction of range and breeding stage ($F_{2,44} = 0.39, P = 0.67$) were not significant. Female fat stores were not different between historic range and range limit ($t = -0.45, P = 0.66$; Fig. 3d).

Muscle profile

Muscle profiles in males varied across breeding stages ($F_{2,44} = 8.81, P < 0.01$; Fig. 3e). In males at both the range limit and historic range, muscle profiles were larger during the pre-parental compared to parental ($t = 7.26, P < 0.001$) and molt ($t = 6.76, P < 0.001$) stages. Muscle profiles in males were not different between birds at the historic or range-limit sites ($F_{1,44} = 0.03, P = 0.85$). The interaction between stage of breeding and range in males was not significant ($F_{2,44} = 0.01, P = 0.98$). The muscle profile for females was smaller at the range limit ($t = -2.30, P = 0.047$; Fig. 3f).

HPA axis activity: corticosterone measurement

Male basal corticosterone values showed an interaction between range and breeding stage ($F_{5,40} = 5.18, P = 0.01$). Basal corticosterone values were higher in males at the range limit during the pre-parental stage only ($t = 5.48, P = 0.05$). For males, corticosterone levels increased from basal levels in response to capture and restraint stress during pre-parental ($F_{3,6} = 49.74, P < 0.01$; Fig. 4a), parental ($F_{3,16} = 39.62, P < 0.01$; Fig. 4b) and molt stages ($F_{3,19} = 11.17, P < 0.01$; Fig. 4c). During the pre-parental stage, the stress response was elevated in birds at the range limits compared to the historic range ($F_{1,8} = 13.29, P < 0.01$) but no differences were found during the parental ($F_{1,18} = 0.002, P = 0.96$) or molt stages ($F_{1,15} = 0.88, P = 0.36$). During the pre-parental stage corticosterone levels were higher in the range limit birds at 10 ($t = 2.38, P = 0.03$), 30 ($t = 2.27, P = 0.04$) and 60 ($t = 3.14, P = 0.009$) min. The interactions between stress and range were not significant during pre-parental ($F_{3,6} = 1.85, P = 0.20$), parental ($F_{3,15} = 0.51, P = 0.60$), and molt stages ($F_{3,13} = 0.07, P = 0.97$).

In males, integrated corticosterone levels were highest during the pre-parental stage compared to parental and parental stages.
molt stages \((F_{6,48} = 40.91, P = 0.001)\). The interaction of breeding stage and range was significant \((F_{2,48} = 6.47, P = 0.003)\) so integrated corticosterone levels were compared within each stage. Integrated corticosterone was higher in birds at the range limit than the historic range during the pre-parental stage \((t = 3.97, P = 0.004)\) but no differences were found during the parental \((t = 0.02, P = 0.98)\) or molt \((t = 1.65, P = 0.11)\) stages.

For females, corticosterone levels increased from basal levels in response to capture and restraint stress \((F_{3,6} = 32.62, P < 0.01; \text{Fig. 4d})\). The corticosterone levels in response to acute restraint stress were not different between the historic and range-limit sites \((F_{1,8} = 4.31, P = 0.07)\). Integrated corticosterone was also not different between historic and range-limit sites \((t = -2.07, P = 0.07)\).

**Discussion**

The range of Gambel’s white-crowned sparrows has expanded northwards over the past 30 years and we suggest that this has been facilitated by an expansion of deciduous woody shrubs leading to an increase in available breeding habitat. Relative shrub cover has increased 63 % over the past 50 years along the Sagavanirktok River which is the second largest drainage on the North Slope and encompasses our study sites \((\text{Tape et al. 2006})\). As a consequence of the range expansion, compared to more southern sites, birds breeding at the northern edge of the range experienced a harsher environment that featured colder temperatures, especially when accounting for wind chill. This is not surprising, since it is known that spring and summer temperatures are typically warmer in the foothills of the Brooks Range and decline towards the Arctic Ocean \((\text{Kade et al. 2005}; \text{Nelson et al. 1997})\). Sweep net arthropod biomass was significantly different at only one time point, which may suggest a slightly later emergence of arthropods at the range limit; however, no differences were found later in the season. We also cannot discount other potential differences in food resources such as the abundance of berries. This study investigated the adjustments in morphology and physiology necessary for range expansion and colonization of a harsh environment.

Male sparrows breeding at the limits of their range possessed significantly higher body condition indices. This pattern could be caused either by differences in the birds that initially arrive at the range limit or by differences in birds that successfully gain territories and breed at northern sites; we do not have data that will allow us to conclusively disentangle these competing explanations. However, existing evidence suggests that these are phenotypic traits that are associated with individuals that are responsible for dispersing into new areas. Studies in naked mole rats \((\text{Heterocephalus glaber})\), side-blotched lizards \((\text{Uta stansburiana})\), and Belding’s ground squirrels \((\text{Urocitellus beldingi})\) have shown that dispersing individuals often have a higher body mass than non-dispersers \((\text{Sinervo et al. 2006}; \text{Holekamp 1986}; \text{O’Riain et al. 1996})\). Additionally, increased mass may be important for surviving in a harsher environment by increasing thermal inertia according to Bergmann’s rule \((\text{Kendeigh 1969})\). Tarsus, wing chord and beak length were not different between the two sites suggesting that there are no structural differences in size between the two populations. This evidence, in conjunction with our current findings, suggests that heavier birds for their body size and, therefore, perhaps higher quality Gambel’s white-crowned sparrows, may be responsible for extending their population’s range. Interestingly, the difference in mass could not be attributed to larger muscle profiles or fat stores in the birds at the range limit. This would suggest that other components of the bird’s morphology are generally larger or heavier and were not elucidated by our measurements.

We found basal levels of corticosterone to be elevated in male birds at the range limit during the pre-parental stage. Our findings are similar to results obtained from Puget Sound white-crowned sparrows \((\text{Zonotrichia leucophrys pugetensis})\) breeding in novel high-altitude sites. Populations of \(\text{Z. leucophrys pugetensis}\) have significantly elevated levels of basal corticosterone with increasing altitude, which the authors attributed to harsher, colder and less familiar conditions than previously encountered by this sub-species breeding at lower altitudes \((\text{Addis et al. 2011})\). The \(\text{Z. leucophrys pugetensis}\) study findings are, however, contradicted by those from studies of an introduced population of house sparrows \((\text{Passer domesticus})\) undergoing range expansion in Africa and in female dark-eyed juncos \((\text{Junco hyemalis})\) invading the urban areas from surrounding montane areas in San Diego, California in which the basal corticosterone levels did not differ between birds on the edge of the population’s range compared to birds breeding within the historic range \((\text{Atwell et al. 2012}; \text{Liebl and Martin 2012})\). However, in both of these studies it is unlikely that the birds were faced with thermally challenging temperatures.

One potential explanation for elevated basal corticosterone levels in birds at the northern limit of the range is that food resources were limited during the early season, forcing birds to draw on energy stores to meet increased metabolic demands associated with a harsher environment. Corticosterone has an effect on many catabolic processes such as liberation of lipids from adipose tissues and the breakdown of protein for glucose production \((\text{Sapolsky et al. 2000})\). In birds, corticosterone implants lead to declines in muscle profile and fat stores \((\text{Astheimer et al. 2000})\). However, in males the inability to meet energetic demands is not supported by our fat or muscle data as we did not
detect differences in fat stores or muscle profiles between individuals at the range limit and southern sites during any stage of breeding. This does not disprove the assumption that increased corticosterone levels are contributing to the regulation of metabolism, but instead suggests that any catabolic effects of corticosterone must have been offset by increased food intake.

An alternative explanation for elevated basal corticosterone levels in birds at the northern limit of the range is that food resources were limited during the early season, but that rather than drawing on energy reserves the birds were able to increase foraging effort to meet this demand. Corticosterone implants have been shown to increase foraging (Astheimer et al. 1992; Angelier et al. 2007) and increase activity range (Breuner and Hahn 2003). Our sweep net sampling showed that arthropod biomass was comparable at the two sites, other than the week of 18 June 2012, suggesting that canopy and airborne arthropods were equally abundant across the range. However, during the pre-parental stage, arthropod dry biomass was low and birds may have been reliant upon other food sources such as berries (Vaccinium uliginosum, Empetrum nigrum, Vaccinium vitis-idaea, and Arctous rubra) and catkins (Betula spp. and Salix spp.). Plant community surveys in the area surrounding Franklin Bluffs, which is dominated by wet non-tussock tundra, indicate that berry-producing species are uncommon and thus would be a limited resource (Kade et al. 2005). This is contrasted by the abundance of berry-producing species found in moist acidic tussock tundra within the historic range (Walker et al. 1994). If food resources are different between the sites during the pre-parental period, then the disparity in observed corticosterone might be a requisite of northern birds to increase foraging effort and activity range to maintain body weight. The difference in basal corticosterone was absent during the parental or molt stages in both sexes despite the colder ambient temperatures during the parental stage. However, during these later breeding stages, arthropod dry biomass is increasing and temperatures remain in the vicinity of the thermal neutral zone. Therefore, elevated levels of corticosterone are no longer necessary to increase foraging effort, activity range, or catabolic processes in order to meet energetic demands.

HPA axis activity and release of corticosterone in response to capture restraint stress was significantly higher in males breeding at the range limit compared to individuals at historic sites, but only during the pre-parental stage. This is consistent with studies in other bird species such as house sparrows (Liebl and Martin 2012), Puget Sound white-crowned sparrows (Addis et al. 2011), Lapland longspurs (Calcarius lapponicus), snow buntings (Plectrophenax nivalis) (Walker et al. 2015), bush warblers (Cettia diphone) (Wingfield et al. 1995a) but also in western fence lizards (Sceloporus occidentalis) which showed that individuals at the edge of their range have an elevated HPA axis activity in response to acute restraint stress. In addition, a significant relationship exists between latitude and HPA activity in birds (Jessop et al. 2013). The activity of the HPA axis is highly plastic and is influenced by environmental harshness in order to promote survival (Boonstra 2004; Wingfield 2008). The influence of environmental factors is exemplified in captive and field studies, when in the absence of environmental and social stimuli, HPA axis activity is greatly attenuated (Romero and Wingfield 1999) while it is up-regulated in response to increased stimuli such as storms (Astheimer et al. 1995; Rogers et al. 1993; Wingfield et al. 1983). Corticosterone has been proposed to be a key mediator of trade-offs between the current life history stage and the emergency life history stage (Wingfield et al. 1995b). Entry into the emergency life history stage results in the abandonment of the current life history stage and redirection of resources to self-preservation which can have detrimental consequences on reproductive output.

Elevation in HPA activity at the range limit of a population, and the rapid rise of corticosterone levels in response to stress, may be critical for determining the success of the individual in a novel environment as it promotes rapid changes in behavior and physiology. The fact that not all individuals within our study population demonstrated the same HPA axis activity, suggests that there are benefits associated with elevated corticosterone that enhance self-preservation but may also be associated with long-term fitness costs (i.e., reproductive output, molt, growth, protection against oxidative damage) (Angelier et al. 2013).

HPA axis activity declined at the transition from the pre-parental to parental stages of breeding and is consistent with previous research in Low- and High-Arctic breeding birds (Holberton and Wingfield 2003; Krause et al. 2015; Meddle et al. 2003; Reneerkens et al. 2002; Walker et al. 2015). However, contrary to our original prediction, we did not find a difference in HPA axis activity between birds at the limits of the range compared to those from the historic range during the parental or molt stages. This may suggest that as arctic birds become parental, regardless of location in the range, HPA axis activity must be dampened in order to ensure parental investment (Walker et al. 2015; Wingfield et al. 1995b). Since corticosterone mediates life history trade-offs, maintenance of elevated HPA axis activity through the parental stage may trigger unwarranted entry into the emergency life history stage. Previous research indicated that elevated corticosterone levels and entry into the emergency life history stage during incubation or the nestling phase were correlated with reproductive failure because resources were allocated away from reproduction towards self-maintenance (Astheimer et al. 1995; Spée et al. 2010; Thierry et al. 2013; Wingfield et al. 1983). During molt, the HPA axis activity is typically suppressed, presumably to facilitate the growth of high-quality...
feathers (Cornelius et al. 2011; Romero and Wingfield 1999). There was a trend, although not significant, for corticosterone to be reduced during molt at the range limits.

Conclusion

Some species’ ranges are changing at an unprecedented rate and individuals are moving into areas that may differ in quality, thus exposing them to potentially challenging new conditions. In this study, breeding sites for Gambel’s white-crowned sparrows at the northern edge of their range were significantly colder during the early season compared to southern historic sites. Investigating the basic physiology of these individuals at range limits will enable us to better understand the mechanisms that drive species range shifts and ultimately the colonization of novel habitats. Higher body condition index in birds at the northern limit of their range may suggest that selection at some level has occurred. Enhanced HPA axis activity may be just one of many physiological mechanisms underlying adaptation to the challenges of a sub-optimal habitat at the very edge of a population’s range (Wingfield et al. 2015). Male birds at the range limit compared to the historic range had a higher HPA axis activity during the pre-parental stage but no differences were found during the parental or molt stages. This plasticity in the HPA axis may be a result of individual plasticity or genetic selection, but this is difficult to establish without further research. Future studies should address other physiological mechanisms at work, such as changes in hormone receptor density (Liebl and Martin 2013) and modulation of hormone-binding proteins to investigate whether regulation is occurring at multiple levels (Breuner et al. 2003). Such studies are critical for understanding and predicting the extent of species adaptation in an environment that is changing at an unprecedented rate.

Acknowledgments

This work was supported by the National Science Foundation Office of Polar Programs ARC 0909133, ARC 0908444, and ARC 0908602 to John C. Wingfield, Natalie Boelman, and Laura Gough. Additionally, Jonathan H. Pérez was supported in part by a National Science Foundation Graduate Research Fellowship (NSF GFRP 1148897) and Simone L. Meddle acknowledges Roslin Institute strategic grant funding from the Biotechnology and Biological Sciences Research Council. We would like to thank Molly Timm, Jeb Timm, and Jake Schas for logistical support and sample collections in the field. We thank Douglas L. Kane, Robert Gieck, Ken Irving, and Emily Youcha from the University of Alaska Fairbanks for the weather data collected at Franklin Bluffs. We would also like to thank Tom P. Hahn and Marilyn Ramenofsky for valuable feedback on the manuscript.

Author contribution statement

JSK, SLM and JCW orginally conceived the experiment. JSK, SLM, SKS, HEC, MAM, LNQ, AA, JHP, and JCW conducted the field work. Laboratory work was performed by JSK, HEC, AA, and LNQ. Data were analyzed and the paper was written by JSK, LNQ, and HEC with JSK performing most of the work. Editorial advice was provided by all authors listed.

Open Access

This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

Addis E, Davis J, Miner B, Wingfield J (2011) Variation in circulating corticosterone levels is associated with altitudinal range expansion in a passerine bird. Oecologia. doi:10.1007/s00442-011-2001-5

Angelier F, Wingfield JC (2013) Importance of the glucocorticoid stress response in a changing world: theory, hypotheses and perspectives. Gen Comp Endocrinol 190:118–128. doi:10.1016/j.ygcen.2013.05.022

Angelier F, Shaffer SA, Weimerskirch H, Trouvé C, Chastel O (2007) Corticosterone and foraging behavior in a pelagic seabird. Physiol Biochem Zool 80:283–292. doi:10.1086/512585

Angelier F, Bost C-A, Giraudel M, Bouteloup G, Dano S, Chastel O (2008) Corticosterone and foraging behavior in a diving seabird: the Adélie penguin, Pygoscelis adeliae. Gen Comp Endocrinol 156:134–144. doi:10.1016/j.ygcen.2007.12.001

Angelier F, Wingfield JC, Trouvé C, De Grissac S, Chastel O (2013) Modulation of the prolactin and the corticosterone stress responses: do they tell the same story in a long-lived bird, the Cape petrel? Gen Comp Endocrinol 182:7–15

Astheimer LB, Buttemer WA, Wingfield JC (1992) Interactions of corticosterone with feeding, activity and metabolism in passerine birds. Ornis Scand 23:355–365. doi:10.2307/3676661

Astheimer LB, Buttemer WA, Wingfield JC (1995) Seasonal and acute changes in adrenocortical responsiveness in an Arctic-breeding bird. Horm Behav 29:442–457. doi:10.1006/hbeh.1995.1276

Astheimer LB, Buttemer WA, Wingfield JC (2000) Corticosterone treatment has no effect on reproductive hormones or aggressive behavior in free-living male tree sparrows, Spizella arborea. Horm Behav 37:31–39. doi:10.1006/hbeh.1999.1555

Atwell JW, Cardoso GC, Whittaker DJ, Campbell-Nelson S, Robertson KW, Ketterson ED (2012) Boldness behavior and stress physiology in a novel urban environment suggest rapid correlated evolutionary adaptation. Behav Ecol 23:960–969. doi:10.1093/beheco/ars059

Bailey RE (1952) The incubation patch of passerine birds. Condor 54:121–136. doi:10.2307/1365062

Bairlein F, Simons D (1995) Nutritional adaptations in migrating birds. Isr J Zool 41:357–367

Boelman NT et al (2015) Greater shrub dominance alters breeding habitat and food resources for migratory songbirds in Alaskan arctic tundra. Glob Change Biol 21:1508–1520. doi:10.1111/gcb.12761

Boonstra R (2004) Coping with changing northern environments: the role of the stress axis in birds and mammals. Integr Comp Biol 44:95–108. doi:10.1093/icb/44.2.95

Borchers DE, Bentz TD (2005) Causes and consequences of animal dispersal strategies: relating individual behaviour to spatial dynamics. Biol Rev 80:205–225. doi:10.1017/S1464793904000645

Breuner CW, Hahn TP (2003) Integrating stress physiology, environmental change, and behavior in free-living sparrows. Horm Behav 43:115–123. doi:10.1016/S0018-506X(02)00020-X
Breuner CW et al (2003) Differential mechanisms for regulation of the stress response across latitudinal gradients. Am J Physiol Regul Integr Comp Physiol 285:R594–R600. doi:10.1152/ajpregu.00748.2002

Chilton G, Baker M, Barrentine C, Cunningham M (1995) White-crowned sparrow no. 183. The birds of North America. The American Ornithologist’s Union. Academy of Natural Sciences of Philadelphia

Cornelius JM, Perfito N, Zann R, Breuner CW, Hahn TP (2011) Physiological trade-offs in self-maintenance: plumage molt and stress physiology in birds. J Exp Biol 214:2768–2777. doi:10.1242/jeb.057174

Cote J, Clobert J, Brodin T, Fogarty S, Sih A (2010) Personality-dependent dispersal: characterization, ontogeny and consequences for spatially structured populations. Philos Trans R Soc B Biol Sci 365:4065–4076. doi:10.1098/rstb.2010.0176

Dunlap KD, Wingfield JC (1995) External and internal influences on indices of physiological stress. I. Seasonal and population variation in adrenocortical secretion of free-living lizards, Sceloporus occidentalis. J Exp Zool 271:36–46. doi:10.1002/jez.1402710105

Environmental Data Center (2013) Meteorological monitoring program at Toolik, Alaska. Toolik Field Station, Institute of Arctic Biology, University of Alaska Fairbanks, Fairbanks, AK 99775. http://toolik.alaska.edu/edc/abiotic_monitoring/data_query.php. Accessed 2 Mar 2013

Holberton RL, Wingfield JC (2003) Modulating the corticosterone stress response: a mechanism for balancing individual risk and reproductive success in Arctic-breeding sparrows? Auk 120:1140–1150

Holekamp KE (1986) Proximal causes of natal dispersal in Belding’s ground squirrels (Spermophilus Beldingi). Ecol Monogr 56:365–391. doi:10.2307/1942552

Jessop TS, Woodford R, Symonds MRE (2013) Macrostress: do large-scale ecological patterns exist in the glucocorticoid stress response of vertebrates? Funct Ecol 27:120–130. doi:10.1111/j.1365-2486.2012.02057.x

Kade A, Walker DA, Raynolds MK (2005) Plant communities and soils in cryoturbated tundra along a bioclimate gradient in the Low Arctic, Alaska. Phytocoenologia 35:761–820

Kane DL (2013) Climate data from the North Slope Hydrology Research project. University of Alaska Fairbanks, Water and Environmental Research Center, Fairbanks. http://ine.uaf.edu/word/projects/NorthSlope/. Accessed 13 Mar 2013

Kendeigh SC (1969) Tolerance of cold and Bergmann’s rule. Auk 86:13–25. doi:10.2307/4083537

Krause JS, McGuitian MA, Bishop VR, Wingfield JC, Meddle SL (2015) Decreases in mineralocorticoid but not glucocorticoid receptor mRNA expression during the short Arctic breeding season in free-living Gambel’s white-crowned sparrows (Zonotrichia leucophrys gambelii). J Neuroendocrinol 27:66–75. doi:10.1111/jne.12237

Landys MM, Ramenofsky M, Guglielmo CG, Wingfield JC (2004) The low-affinity glucocorticoid receptor regulates feeding and lipid breakdown in the migratory Gambel’s white-crowned sparrow Zonotrichia leucophrys gambelii. J Exp Biol 207:143–154. doi:10.1242/jeb.007734

Liebl AL, Martin LB (2012) Exploratory behaviour and stressor hyper-responsiveness facilitate range expansion of an introduced songbird. Proc R Soc B Biol Sci 279:4375–4381. doi:10.1098/rspb.2012.1606

Liebl AL, Martin LB (2013) Stress hormone receptors change as range expansion progresses in house sparrows. Biol Lett. doi:10.1098/rsbl.2013.0181

MacArthur RH (1972) Geographical ecology: patterns in the distribution of species. Princeton University Press, Princeton

Meaddle SL, Owen–Ashley NT, Richardson ML, Wingfield JC (2003) Modulation of the hypothalamic–pituitary–adrenal axis of an Arctic-breeding polygynandrous songbird, the Smith’s longspur, Calidris pucta. Proc R Soc Lond Ser B Biol Sci 270:1849–1856. doi:10.1098/rspb.2003.2455

Morton ML, King JR, Farner DS (1969) Postnuptial and postjuvenile molt in white-crowned sparrows in Central Alaska. Condor 71:376–385

Nelson FE, Shiklomanov NI, Mueller GR, Hinkel KM, Walker DA, Bockheim JG (1997) Estimating active-layer thickness over a large region: Kuparuk river basin, Alaska, U.S.A. Arct Alp Res 29:367–378. doi:10.2307/151985

Normet CJ (1992) Comparative breeding biology of Harris’ sparrows and Gambel’s white-crowned sparrows in the Northwest Territories, Canada. Condor 94:955–975

Normet CJ (1993) Nest-site characteristics and nest predation in Harris’ sparrows and White-crowned sparrows in the Northwest Territories, Canada. Auk 110:769–777

Oakeson BB (1954) The Gambel’s sparrow at Mountain Village, Alaska. Auk 71:351–365

O’Riain MJ, Jarvis JUM, Faulkes CG (1996) A dispersive morph in the naked mole-rat. Nature 380:619–621

Osczevski RJ (1995) The basis of wind chill. Arctic 48:372–382

Parresan C, Yohé G (2003) A globally coherent fingerprint of climate change impacts across natural systems. Nature 421:37–42

Reneerkens J, Morrison G, Ramenofsky M, Piersma T, Wingfield JC (2002) Baseline and stress-induced levels of corticosterone during different life cycle stages in a shorebird on the High Arctic breeding grounds. Physiol Biochem Zool 75:200–208. doi:10.1086/340853

Rogers CM, Ramenofsky M, Ketterson ED, Nolan V Jr, Wingfield JC (1993) Plasma corticosterone, adrenal mass, winter weather, and season in nonbreeding populations of dark-eyed juncos (Junco hyemalis hyemalis). Auk 110:279–285. doi:10.2307/4088556

Romero LM, Wingfield JC (1999) Alterations in hypothalamic–pituitary–adrenal function associated with captivity in Gambel’s white-crowned sparrows (Zonotrichia leucophrys gambelii). Comp Biochem Physiol B Biochem Mol Biol 122:13–20. doi:10.1016/S0305-0491(98)10161-X

Sapolsky RM, Romero LM, Munck AU (2000) How do glucocorticoids influence stress responses? Integrating permissive, suppressive, and preparative actions. Endocr Rev 21:55–89. doi:10.1210/er.21.1.55

Sauer JR, Hines JE, Fallon JE, Pardieck KL, Ziolkowski DJ Jr, Link WA (2014) The North American breeding bird survey, results and analysis 1966–2012. US Geological Survey Patuxent Wildlife Research Center, Laurel

Sexton JP, McIntyre PJ, Angert AL, Rice KJ (2009) Evolution and adaptation variation in adrenocortical secretion of free-living lizards, Sceloporus occidentalis. J Exp Zool 271:36–46. doi:10.1002/jez.1402710105

Silverin B (1997) The stress response and autumn dispersal behaviour in willow tits. Anim Behav 53:451–459. doi:10.1006/anbe.1996.0295

Sinervo B, Calsbeek R, Comendant T, Both C, Adamopoulou C, Clobert J (2006) Genetic and maternal determinants of effective dispersal: the effect of sire genotype and size at birth in side-blotched lizards. Am Nat 168:88–99. doi:10.1086/505765

Spé M, Beaulieu M, Dervaux A, Chastel O, Le Maho Y, Raclot T, Spée M, Raclot T, Calsbeek R, Adamopoulou C, Sinervo B, Both C, Comendant T, Adamopoulou C, Clobert J, (2006) The North American breeding bird survey, results and analysis 1966–2012. US Geological Survey Patuxent Wildlife Research Center, Laurel

Sturm M, Racine C, Tape K (2001) Climate change: increasing shrub abundance in the Arctic. Nature 411:546–547

Tape KEN, Sturm M, Racine C (2006) The evidence for shrub expansion in Northern Alaska and the Pan-Arctic. Glob Change Biol 12:686–702. doi:10.1111/j.1365-2486.2006.01128.x

Springer
Thierry A-M, Massemin S, Handrich Y, Raclot T (2013) Elevated corticosterone levels and severe weather conditions decrease parental investment of incubating Adélie penguins. Horm Behav 63:475–483. doi:10.1016/j.yhbeh.2012.12.011

Walker MD, Walker DA, Auerbach NA (1994) Plant communities of a tussock tundra landscape in the Brooks Range Foothills, Alaska. J Veg Sci 5:843–866. doi:10.2307/3236198

Walker BG, Meddle SL, Romero LM, Landys MM, Reneerkens J, Wingfield JC (2015) Breeding on the extreme edge: modulation of the adrenocortical response to acute stress in two High Arctic passerines. J Exp Zool Part A Ecol Genet Physiol 323:266–275. doi:10.1002/jez.1923

Wingfield JC (2008) Comparative endocrinology, environment and global change. Gen Comp Endocrinol 157:207–216. doi:10.1016/j.ygcen.2008.04.017

Wingfield JC, Farner DS (1978) The annual cycle of plasma irLH and steroid hormones in feral populations of the white-crowned sparrow, Zonotrichia leucophrys gambelii. Biol Reprod 19:1046–1056. doi:10.1095/biolreprod19.5.1046

Wingfield JC, Krause JS, Perez JH, Chmura HE, Nemeth Z, Words KR, Calisi RM, Meddle SL (2015) A mechanistic approach to understanding range shifts in a changing world: what makes a pioneer? Gen Comp Endocrinol. doi: 10.1016/j.ygcen.2015.08.022

Wingfield JC, Sapolsky RM (2003) Reproduction and resistance to stress: when and how. J Neuroendocrinol 15:711–724. doi:10.1046/j.1365-2826.2003.01033.x

Wingfield JC, Moore MC, Farner DS (1983) Endocrine responses to inclement weather in naturally breeding populations of white-crowned sparrows (Zonotrichia leucophrys pugetensis). Auk 100:56–62. doi:10.2307/4086277

Wingfield JC, Vleck CM, Moore MC (1992) Seasonal changes of the adrenocortical response to stress in birds of the Sonoran desert. J Exp Zool 264:419–428. doi:10.1002/jez.1402640407

Wingfield JC, Kubokawa K, Ishida K, Ishii S, Wada M (1995a) The adrenocortical response to stress in male bush warblers, Cettia diphone: a comparison of breeding populations in Honshu and Hokkaido, Japan. Zool Sci 12:615–621

Wingfield JC, O’Reilly KM, Astheimer LB (1995b) Modulation of the adrenocortical responses to acute stress in Arctic birds: a possible ecological basis. Am Zool 35:285–294. doi:10.1093/icb/35.3.285

Wingfield JC et al (2004) Arctic spring: the arrival biology of migrant birds. Acta Zool Sin 50:948–960