Corticosterone and the Stress Response in Young Western Screech-Owls: Effects of Captivity, Gender, and Activity Period

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Accepted by G.K.S. 8/12/96

ABSTRACT

We used a standard handling protocol to examine the stress response of captive young western screech-owls during their active (nighttime) and inactive (daytime) periods and to compare the stress responses of captive and free-living owls. Circulating corticosterone levels were significantly higher during the inactive period than in the active period in this nocturnal species. This suggests that the daily pattern of corticosterone secretion is reversed in nocturnal birds and is correlated with activity period rather than with the light/dark cycle. Young (ca. 4–5 mo old) screech-owls of both sexes showed increases in plasma corticosterone up to 30 min after capture, followed by significant decreases at 60 min. This pattern is similar to those of other species of birds examined previously, except that decreases in corticosterone at 60 min rarely have been observed. Such decreases may be the result of physiological differences between adult and young birds, habituation to handling in captive birds, or the effects of body condition. Corticosterone levels and the response to capture and handling were comparable in captive and free-living owls, which suggests that the captive owls were not subjected to chronically high levels of stress.

Introduction

Adrenocortical hormones are released in response to stressful stimuli in all vertebrates, including birds (e.g., Harvey et al. 1984; Smith et al. 1994). The release of corticosterone, the primary avian adrenocorticoid, into the bloodstream facilitates many of the physiological adjustments that enable birds to restore homeostasis in the face of disruptive events (see, e.g., Harvey et al. 1984; Cherel et al. 1988; Gray et al. 1990; Buttemer et al. 1991). Adrenal activity also produces behavioral changes in birds that may contribute to the amelioration of stressful situations (Wingfield 1994). For example, corticosterone stimulates increased locomotor and foraging activity (Gray et al. 1990; Asheimer et al. 1992) and reduced reproductive activity (Wingfield 1984, 1994). These behavioral responses direct birds’ efforts toward energy acquisition and survival and away from energetically costly behaviors, such as breeding.

The development of a standardized capture and handling stress protocol (Wingfield et al. 1992) has allowed comparison of stress responses among unrelated avian species experiencing different ecological and seasonal conditions (see, e.g., Wingfield et al. [1995] and references therein). Consequently, the interplay among the physiological and ecological factors that produce the stress response in breeding and nonbreeding adult birds is being examined closely (Wingfield 1994; Wingfield et al. 1995).

One area that has not been explored systematically is the effect of stress on the adrenal response of young birds. The hypothalamic-pituitary-adrenocortical axis may not be fully mature at hatching. For example, in domestic fowl (Gallus domesticus), the adrenal glands increase their responsiveness to adrenocorticotropic hormone during the first several months of life, as corticosterone secretion develops into the adult-like pattern (Webb and Mashaly 1985). Thus, it is important to determine if newly independent juvenile birds of either or both sexes respond to stress in a manner similar to adults.

Furthermore, previous investigations of avian stress responses have been limited to the birds’ active periods, and all avian species heretofore examined are active during the daytime. Because of this, it is not clear if the rate of onset or magnitude of the stress response differs between the active and inactive periods in birds. In addition, there is often a daily rhythm in plasma corticosterone secretion (see, e.g., Boissin and Assenmacher 1971; Joseph and Meier 1973; Beuving and Vonder 1977; but see Marra et al. 1995), and maximal corticosterone levels occur during the inactive (nocturnal) period. As a result, it is not known if elevated corticosterone secretion is a function of nighttime or of inactivity. Examination of corticosterone secretion in species of nocturnal birds would differentiate between the two.

During their active period, birds with already elevated corticosterone levels do not show additional responsiveness when subjected to the capture and handling stress protocol (Smith
et al. 1994). Previous studies indicate that captivity may be
eone factor that results in elevated corticosterone levels. For
instance, Marra et al. (1995) found that secretory profiles of
corticosterone differed significantly between captive and free-
living individuals in two species of sparrows in the genus Zono-
trichia. Captive individuals exhibited levels between two and
three times higher than those of free-living birds despite the
fact that they had the opportunity to acclimate to captivity for
more than 1 mo prior to sampling. This begs the question of
whether captive birds respond to capture and handling stress
with an adrenocortical response.

Our objectives in the present study were to (1) examine the
capture and handling stress response in young birds of both
sexes, (2) determine if and how the stress response differs
during the active and inactive periods, and (3) examine differ-
ences in corticosterone titers and the stress response in captive
and free-living birds.

Material and Methods

Study Species

In this study we examined the stress response in captive and
free-living young western screech-owls (Otus kennicottii). These
small (ca. 250 g) nocturnal owls are common in low-
elevation forests, riparian woodlands, and suburban areas
throughout the western United States (Johnsgard 1988). They
are nonmigratory, territorial, and secondary cavity nesters, and
they breed in both natural cavities and artificial nest boxes
(J. R. Belthoff, personal observations). In southwestern Idaho,
where our studies were conducted, young owls hatch in mid-
April and fledge approximately 28 d later. Following a 6–10-wk
dependency period, young owls disperse to winter territories
beginning in late June and continuing through mid-August,
although most individuals disperse in mid-July (Belthoff and
Dufty 1995; E. A. Ellsworth and J. R. Belthoff, unpublished data). Once young have settled, they defend the area from
conspecifics and commence breeding the following spring if
resources (e.g., mates, nest cavities, cover, food) are adequate
on the territory.

Laboratory Samples

During the period of May 12–25, 1995, we collected 12 nestling
western screech-owls and transported them to the animal care
facility on the campus of Boise State University. The nests from
which owls were collected (n = 5) were in artificial nest boxes
mounted along the Snake River in Elmore and Owyhee Coun-
ties and along the Boise River in Ada County, Idaho. On the
basis of development at the time of collection, all owls appeared
to be about 23–24 d old, or about 1 wk from leaving the
nest box.

In captivity, owls were kept in wire-mesh cages measuring
approximately 0.5 m × 0.5 m × 1.0 m, which were placed in
individual sound-attenuating isolation chambers. In the front
of each chamber was a clear plastic window (ca. 29 cm × 45
cm). Once in the chambers, owls could not interact visually or
vocally. We provided owls with laboratory mice, day-old
chickens, and water ad lib. The owls remained undisturbed
except for feeding and cage maintenance, which required less
than 1 h each day. Once each day we also removed each owl
from its cage to read its backpack-mounted pedometer (Mi-
cronta Mini-Jog Mate, catalog no. 63-667, Radio Shack), which
recorded locomotor activity as part of another study (see
Ritchison et al. [1992] and Belthoff and Dufty [1995] for other
details on pedometers and their use). Thus, each bird also was
handled by one of the investigators for approximately 1 min
each day during its captivity. Light entered the animals’ room
through a large (12 m × 1.5 m), south-facing set of windows,
which allowed the owls to experience the natural photoperiod.
All owls remained in good health while in captivity.

Between August 25 and September 12, 1995, we performed
the capture and handling stress protocol on 4–5-mo-old cap-
tive owls during each of two time periods, termed active and
inactive. For western screech-owls, the active period corre-
sponds to nighttime and the inactive period to daytime, which
is opposite to diurnal species. For the active period, samples
were collected 1 h after dark, at approximately 2230–2300
hours. For the inactive period, samples were collected between
1300 and 1500 hours. Owls were randomly assigned to one of
two treatment sequences such that six owls initially were sam-
ples during the active period and the other six initially were
sampled during the inactive period. An average of 6 d was
allowed to pass before taking blood during the second period.
The owls weighed an average of 213 g (range: 174–281 g)
when this study began. Prior to this study, the 12 owls had
been bled on a weekly basis (beginning 1 wk after capture; 600
µL of whole blood removed per sample) as part of another
study to assess circulating hormones in relation to locomotor
activity patterns and dispersal (see Belthoff and Dufty 1995).

To assess the effects of handling stress on circulating cortico-
sterone levels during active and inactive periods, five blood
samples were obtained from each owl during each activity
period. These samples were obtained 1, 5, 10, 30, and 60 min
following removal of the owl from its home cage (Wingfield
et al. 1992). Blood samples were collected by puncture of the
brachial vein with a 26-gauge needle. Whole blood was col-
llected into between one and three heparinized microhematocrit
tubes. These were kept on ice until centrifuged (within 1 h)
at 8,000 rpm for 3 min. Plasma was harvested and frozen
at −20°C until the hormone assay was performed. Between
sampling periods, owls were held in cotton-mesh sacks and
left undisturbed.

The sex of young screech-owls generally cannot be deter-
dined accurately on the basis of morphological or plumage
characteristics alone (J. R. Belthoff, personal observations).
Therefore, after we removed the plasma for radioimmunoassay, we provided the remaining hematocrit samples to a commercial laboratory (Zoogen, Davis, Calif.) for analysis. The laboratory used a heterologous sex-specific probe on DNA isolated from the owl blood to determine the gender of the captive owls.

The study sample consisted of eight males and four females. Following exercise in flight cages, during which time captive owls learned to hunt live mice, the owls were released in accordance with agency permit specifications.

Field Samples

We also obtained blood samples (ca. 600 μL whole blood) from free-living western screech-owls and compared the pattern of corticosterone secretion and the range of plasma corticosterone values with those of captive owls. Fledgling screech-owls (*n* = 11) were captured at night (range: 2150–2335 hours) from June 6 to July 10, 1994, and on June 22, 1995, with mist nets placed within their home ranges. These owls were 2–3 mo younger than the captive birds and had not yet dispersed; some were still dependent on adults for provisioning. Some birds were lured to the nets by broadcasting the calls of adult screech-owls. Ten of the birds previously had been captured as nestlings, bled as described above, given numbered aluminum leg bands, and affixed with backpack radio transmitters (Model SOPB-2190, Wildlife Materials, Carbondale, Ill.; see Belthoff and Ritchison [1989, 1990] for description of transmitters and attachment methods). These birds were sexed as described earlier, and the sample consisted of six males and four females. The remaining fledgling, of unknown sex, had not been captured previously.

Unlike in the stress protocol used for captive birds, when free-flying young were captured, we collected only one blood sample per bird. We recorded the length of time from capture to completion of the sampling procedure for all birds. Six blood samples were collected in 5 min or less, four in 5–10 min, and one in 30 min. Blood was placed in 1.5-mL microcentrifuge tubes, which were kept on ice for 2–3 h, transported to the laboratory, and centrifuged. The plasma and hematocrit were separated and stored as described previously.

Corticosterone Radioimmunoassay

Plasma corticosterone levels in captive owls were measured with the protocol described by Wingfield et al. (1992). Plasma volumes were 20–50 μL. Approximately 2,000 cpm of tritiated corticosterone was added to each sample, along with 300 μL of distilled water, and samples were refrigerated (4°C) for 3 h. Five milliliters of freshly distilled dichloromethane was added to each sample to extract corticosterone, and samples were refrigerated overnight. The lower (organic) phase was removed and dried in a water bath (40°C) under nitrogen gas, and samples were reconstituted with 550 μL of buffer. Duplicate assay tubes received 200-μL aliquots for each sample. An additional 100-μL aliquot was added to a scintillation vial (along with 4.5 mL of scintillant), which enabled us to determine recovery values for the extraction process. Intra- and interassay coefficients of variation were 1%–13%.

Experimental Design and Statistical Analyses

In captive owls, we examined the effects of sex, activity period, and sampling time on circulating corticosterone using a three-factor mixed factorial design (Zolman 1993). In this design, sex was a between-group factor, and both time and activity period represented within-subjects factors or repeated measures. Using ANOVA, we examined the main effects and their interactions following log transformation of the data. If significant effects were detected, we calculated least squares means and compared them using pairwise *t*-tests. Methodological differences (i.e., collection of single vs. multiple blood samples) preclude any direct comparison of free-living and captive screech-ows. However, for field samples we used the nonparametric Mann-Whitney *U*-test to compare corticosterone levels in samples taken after 5 min or less of handling to those with relatively longer handling times (5–10 min). Finally, all data are presented as mean ± standard error, and rejection levels were set at *P* = 0.05.

Results

Captive Screech-Owls

The pattern of the adrenocortical response of captive young screech-ows to handling stress is shown in Figure 1. One bird (male 064) did not exhibit an adrenocortical response to handling stress in either time period. Results for this bird are shown separately in Figure 1 and are not included in the analyses. We detected no effect of sex on the pattern of corticosterone secretion (overall for males: 27.3 ± 1.56 ng/mL, *n* = 70 samples from seven males; overall for females: 27.5 ± 2.06 ng/mL, *n* = 40 samples from four females; Fig. 1A), and sex did not interact with any of the other factors in the model. There was an effect of activity period on the adrenocortical response (Table 1): samples obtained during the inactive (daytime) period (overall for day: 29.5 ± 1.82 ng/mL, *n* = 55 samples from 11 birds) exhibited significantly higher corticosterone levels than those obtained during the active (nighttime) period (overall for night: 25.3 ± 1.82 ng/mL, *n* = 55 samples from 11 birds; Fig. 1B). The lack of a significant interaction between time and activity period (Table 1) indicates that the pattern of corticosterone secretion was similar during day and night, even though absolute corticosterone levels were higher during the day.

For the remaining 11 birds included in the analysis, basal corticosterone levels (i.e., those collected after only 1 min of
Corticosterone levels were significantly greater in blood samples collected after 5–10 min than in samples collected within 5 min (Mann-Whitney U-test, $U = 24, P = 0.011$). The single sample collected after 30 min exhibited the highest corticosterone level.

**Discussion**

**Captive Young Western Screech-Owls**

Captive young western screech-owls of both sexes exhibited similar patterns of corticosterone secretion in response to the handling stress protocol. This is consistent with the results of earlier studies on adult birds that revealed no gender differences in the adrenocortical response in most, but not all, situations (Wingfield et al. 1992; Astheimer et al. 1994; Smith et al. 1994; Wingfield et al. 1995). Those gender differences in corticosterone secretion that do occur may be associated with parental behavior that evolved in response to unpredictable weather during the breeding season. For example, there is often an inverse relationship between the degree of parental care and maximum corticosterone levels (Wingfield et al. 1995). Parental behavior is unlikely to be developed in sexually immature birds, and this could account for the indistinguishable stress responses in young male and female screech-owls.

Although the overall pattern of secretion was similar in both periods, corticosterone levels were significantly higher during the inactive period. Thus, young screech-owls may have responded differently to handling stress in their active and inactive periods. One explanation for this phenomenon is that handling stress may be more severe when it occurs during a bird's inactive period. That is, birds that are not alert or are asleep may be more startled and alarmed, and produce a stronger adrenal response, than birds that are active and more aware of events happening around them.

*Figure 1.* Circulating corticosterone levels (mean ± SE) of captive juvenile western screech-owls across sampling time periods by A, sex (four females, seven males); B, activity period; and C, all sexes and activity periods. Panel C also presents the corticosterone values of one male (064) that did not show a stress response and was excluded from subsequent analyses. None of the pairwise comparisons in A or B are significant at the $P < 0.05$ level. In C, bars labeled with the same letter do not differ significantly.

*Figure 2.* Circulating corticosterone levels (mean ± SE) in free-living juvenile western screech-owls sampled after three different handling durations. Sample sizes are indicated.

**Free-Living Screech-Owls**

There also was an effect of time on the pattern of corticosterone secretion in free-living fledglings (Fig. 2). Plasma corticoste-
Alternatively, the difference may have resulted from a daily rhythm in corticosterone secretion. Such rhythms have been documented in diurnal species, with increased baseline corticosterone levels occurring during the inactive period (e.g., Boissin and Assenmacher 1971; Joseph and Meier 1973; Beuving and Vonder 1977). Unlike species investigated in these previous studies, screech-owls are nocturnal; thus, their inactive period is during the daytime. If the daily rhythm of corticosterone is likewise reversed, then the stronger stress response we observed during the day may simply reflect higher baseline corticosterone values at that time. Although none of the activity period × time pairwise comparisons achieved significance at the 0.05 level, the baseline (i.e., 1-min) sample nearly did so ($P = 0.0811$), and mean daytime corticosterone values were similarly elevated above mean nighttime values at all sampling times. This supports the idea that while the overall pattern of adrenocortical secretion is similar in both activity periods, the baseline corticosterone levels differ.

Plasma corticosterone levels in young western screech-owls rose significantly in response to the stress protocol between the 1-, 5-, and 10-min samples and plateaued at 30 min, which is similar to results from previous studies (Wingfield et al. 1992; Asheimer et al. 1994; Smith et al. 1994; Wingfield et al. 1995). However, there was a significant decline in circulating corticosterone between 30 and 60 min, which was rarely reported in previous investigations. Most of these earlier studies focused on adults, and circulating corticosterone typically peaked in the 10- or 30-min samples and remained elevated through the 60-min sample.

Interestingly, a reduction in corticosterone titers between the 30- and 60-min samples of the stress protocol also occurs in young captive American kestrels (*Falco sparverius*; J. A. Heath and A. M. Dufty, Jr., unpublished data). In addition, Freeman and Flack (1980) found that plasma corticosterone levels in 3-wk-old domestic chicks typically rise between samples obtained 0–5 min and 5–10 min after handling but then fall in samples taken at 15–20 min and 30–35 min. This pattern may simply reflect incomplete development of the hypothalamo-pituitary-adrenal axis in young birds. Alternatively, any consistent differences in adrenal activity between young and adults may result from different age-related selective pressures. In adults, the adrenocortical response varies in different ecological situations in ways that enhance survival of individuals or their young (Wingfield 1994; Wingfield et al. 1995). A muted adrenocortical response in young birds also may have survival benefits. For example, it could facilitate maintenance of protein stores or reduce movements away from protective parents or out of newly acquired territories. We have not examined the stress response in adult western screech-owls, so it is not known whether the pattern of corticosterone secretion seen in young screech-owls is typical of the species or is age related.

Another potential explanation for the significant drop in corticosterone levels after 60 min is that the owls (and kestrels) were maintained in captivity for many weeks, handled on a daily basis, and bled at least once per week. Thus, the birds could have habituated to being handled, learned that soon they would be returned to their cages, and subsequently moderated their adrenal response to handling. It is unclear whether the domestic chicks (Freeman and Flack 1980) were handled on a regular basis, yet they exhibited a similar pattern of corticosterone secretion.

Finally, there is some evidence that body condition affects the stress response in a variety of vertebrates. For example, poorly nourished young American kestrels show no significant decrease in corticosterone between 30 and 60 min of the stress protocol, while well-nourished birds do (J. A. Heath and A. M. Dufty, Jr., unpublished data). Similarly, green sea turtles (*Chelonia mydas*) with and without fibropapillomas exhibit different responses to capture stress (Aguirre et al.

### Table 1: Results of mixed factorial ANOVA on log-transformed data

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</table>

*Note. Sex was a between-groups factor, and period and time were within-subjects factors.*
Thus, the fact that screech-owls in our study were fed ad lib. and were likely in excellent body condition also could have contributed to the significant decrease in corticosterone between 30 and 60 min. Clearly, stress responses in adult and naive young birds, as well as those in well- and poorly nourished individuals, need to be examined before the reduction in corticosterone after 60 min in young western screech-owls can be interpreted more precisely.

One juvenile male, bird 064, did not produce any stress response during the 1 h of handling in either the active or inactive periods (Fig. 1C). This bird showed no evidence of disease: its body weight, plumage, and behavior all appeared normal. Asheimer et al. (1994) determined that male sparrows undergoing postbreeding molt exhibit minimal increases in plasma corticosterone. However, bird 064 was engaged only in the first prebasic molt typical of the entire cohort of captive screech-owls. Hence, we cannot determine whether this bird had an immature or an abnormal hypothalamo-pituitary-adrenal axis.

Free-Living Young Western Screech-Owls

The initial increase in plasma corticosterone levels displayed by captive juvenile screech-owls was mirrored in free-living juvenile owls captured in mist nets. The two data sets were not compared statistically because of fundamental procedural differences in the timing of sample collection and the number of samples obtained. Nonetheless, corticosterone values for free-living birds sampled in 5 min or less and 5–10 min after capture are similar to, albeit slightly higher than, those found in captive owls over the first 10 min of the handling stress protocol. The highest plasma corticosterone value from free-living young owls was measured in the single sample obtained 30 min after capture. Although this value was more than twice the corresponding mean value for captive owls, it coincided with the period of peak corticosterone levels in the captive birds.

Finally, the similarity between basal corticosterone levels measured in captive owls (the initial stress protocol sample) and free-living owls (the samples obtained in ≤ 5 min) contrasted with the results of Marra et al. (1995), who found that captive sparrows manifest significantly higher basal plasma corticosterone levels throughout the day than do free-living sparrows. It is possible that the owls maintained in captivity for more than 3 mo had adequate time to adjust to captive conditions, whereas the sparrows (35-d captivity) did not.

Conclusion

In summary, our results demonstrate that young western screech-owls exhibit an adrenocortical response to handling stress that is similar in both sexes. Furthermore, by using a nocturnal species, we have uncoupled the active/inactive periods from the light/dark cycle. Circulating corticosterone levels were higher during the owls’ inactive (daytime) period, which suggests that peaks in the diurnal rhythm of corticosterone are associated with the species’ daily activity pattern rather than with the photoperiod. In addition, unlike the results from many previous studies, mean corticosterone levels in young owls declined 30–60 min after capture. We suggest that this may reflect age-related differences in the stress response, acclimation to captivity and handling, or an effect of body condition. Finally, the results obtained from captive owls appear to be similar to those from free-living juvenile screech-owls.

Acknowledgments

We thank L. Belthoff, J. Doremus, B. and C. Dufty, E. and M. Ellsworth, J. Emerson, J. Heath, D. Lowe, and S. Mitchell for assistance in the laboratory and field. We gratefully acknowledge the financial support of the Idaho State Board of Education (specific research grant 595-042), the National Science Foundation (award IBN-9509079), and Boise State University (faculty research fund and faculty research associateship). We also thank the Boise District of the U.S. Bureau of Land Management, the Snake River Birds of Prey National Conservation Area, and particularly J. Doremus for facilitating our fieldwork on owls. We collected young owls under a U.S. Fish and Wildlife Service permit (PRT 785053) and a State of Idaho Department of Fish and Game scientific collecting permit (SCP 930810).

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