Movement Patterns and Habitat use for Sympatric Species

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Abstract

**Background:** Movement is an important characteristic of an animal’s ecology, reflecting perception of and response to environmental conditions. To effectively search for food, movement patterns likely depend on habitat characteristics and the sensory systems used to find prey. We examined movements associated with foraging for two sympatric species of lizards inhabiting the Alvord Basin in the Great Basin Desert of southeastern Oregon. The two species have largely overlapping diets but find prey via different sensory cues, which link to their differing foraging strategies — the long-nosed leopard lizard, *Gambelia wislizenii*, is a visually-oriented predator, while the western whiptail, *Aspidoscelis tigris*, relies heavily on chemosensory cues to find prey.

**Methods:** Using detailed focal observations, we characterized the habitat use and movement paths of each species. We placed markers at the location of focal animals every minute for the duration of each 30-min observation. Afterwards, we recorded whether each location was in the open or in vegetation, as well as the movement metrics of step length, path length, net displacement, straightness index, and turn angle, and then made statistical comparisons between the two species.

**Results:** The visual forager spent more time in open areas, moved less frequently over shorter distances, and differed in patterns of plant use compared to the chemosensory forager. Path characteristics of step length and turn angle differed between species.

**Conclusions:** The visual predator moved in a way that was consistent with the notion that they require a clear visual path to stalk prey whereas the movement of the chemosensory predator increased their chances of detecting prey by venturing further into vegetation. Sympatric species can partition limited resources through differences in search behavior and habitat use.

Introduction

For ambulant organisms, movement involves dynamic interactions that balance the animal’s capabilities and ecological preferences with local environmental conditions. Their ability to move can depend not only on their locomotor structures, but also on their ability to detect and respond to current conditions [1]. On large spatiotemporal scales, movement patterns influence population distributions as well as interactions among species and their environment [2–4]. At the individual level, when and where organisms move can directly influence survival and ultimately fitness [5, 6]. Fine-scale locality data coupled with movement path analyses can identify the factors influencing movement as well as how individual animals perceive and respond to their environment [2, 7–9].

Over the past several decades, studies on animal movement have provided insights into the evolutionary biology, ecology, and physiology of many different taxa [10–14]. Examining spatiotemporal movement has led to significant insights into social behavior [15]. Movement indices reveal intraspecific variation [11, 14, 16–18] or behavioral flexibility [19–22] in foraging. Additionally, variation in habitat structure and resource use can lead to changes in movement patterns [23–28]. Using a fine-scale spatial approach to
studying movement patterns, we aim to elucidate how two sympatric lizard species can partition overlapping food resources.

In the Alvord Basin in southeastern Oregon, two sympatric species of lizard, the long-nosed leopard lizard (*Gambelia wislizenii*) and the western whiptail (*Aspidoscelis tigris*), present an interesting contrast. While phylogenetically in distant clades (Iguania vs Scleroglossa) [29], *G. wislizenii* and *A. tigris* provide an opportunity to compare how two cohabitating species behaviorally partition largely overlapping resources. A previous comparison of these species in the same location showed differences in behavior between the lizards, with *G. wislizenii* spending less time moving than *A. tigris* [30]. We build on their comparison of overall movement levels by analyzing movement patterns and resource acquisition from a habitat-use perspective.

In the southwestern United States, the two species largely overlap in desert habitats with finite resources where the harsh environment requires animals to take maximal advantage of any available food or shelter [31, 32]. Their opportunistic nature, as well as their limited food and habitat resources allowed us to make an interspecies comparison of movement ecology. The two species share an overlapping diet consisting of grasshoppers, beetles, spiders, antlions, and caterpillars but differ somewhat in their food choices, as *G. wislizenii*, with their larger body size, can prey on small lizards while *A. tigris* are proficient at digging for termites and buried larvae [33–37].

Despite sharing habitat and consuming similar prey, the species vary in their foraging strategies and rely differently on sensory modalities for prey detection [34, 35, 38, 39]. Leopard lizards primarily use visual cues for finding food, pursuing prey that moves within their visual detection range (≤ 10 m away) [18, 40–42]. The movement rate for leopard lizards can vary with environmental conditions and among individuals [18, 41]. In contrast, whiptail movement is wide-ranging and can vary with habitat structure [43, 44]. Further, whiptails primarily locate prey (even below the surface) through chemical sampling, only sometimes using vision to detect food items [35, 43, 45].

When compared to strictly visual hunters, chemosensory foragers move more frequently and spend a higher proportion of time moving [46, 47]. However, search strategy does not necessarily indicate the size of the area searched or the path traveled. Our overarching hypothesis is that differences in prey detection are associated with the species’ space use and movement patterns. Our goal was to determine if there were consistent differences in habitat use and movement patterns between animals that generally use different prey detection cues: visual long-nosed leopard lizards and chemosensory western whiptails. We predicted that a more visually-oriented predator would move along straighter paths, travel less, and make more use of open spaces than a chemosensory forager. Finding associations between prey detection strategies and differences in movement patterns can help identify the importance of sensory systems in shaping both foraging strategies and interactions between potential competitors.

**Methods**
Our study was conducted in desert scrub habitat of the Alvord Basin, located in the Great Basin Desert, southeastern Oregon (42°18’N, 118°37’W; datum = WGS 84; elevation 1295 m) from 20 June – 14 July 2017. The study site was a 16-ha gridded plot characterized by open desert sand and hardpan interspersed with patches of shrubs, mainly sage (Artemisia tridentata) and greasewood (Sarcobatus vermiculatus). We conducted observations on the long-nosed leopard lizard, G. wislizenii (visual predator; N = 61), and the western whiptail, A. tigris (chemosensory predator; N = 51), during their morning activity period (0800 – 1030 h). Each observation was conducted by a pair of observers at a distance that allowed for clear observation without disturbing foraging behavior (ca. 2–5 m) [35, 44, 48]. Observations lasted 30 min with one observer tracking lizard movements and the other observer placing markers at the lizard’s location at each 1-min interval. To minimize disturbance, markers were placed after the animal left the immediate area where a marker was to be placed. For each marker, we recorded whether the location was in vegetation or in the open. When the location was in vegetation, we recorded the plant species. Unmarked lizards were captured using a lasso attached to an extendable pole, measured (sex, body mass, snout vent length (SVL)), and marked with a unique paint code before being released at the capture site. We recorded movement data only once for each individual animal.

Habitat And Movement Analysis

A full observation generated 31 location-time points for each animal, consisting of x-y coordinates and associated vegetation measurements. From the sequence of locations, we calculated the path characteristics of step length, path length, net displacement, straightness index, and turn angle (Table 1). Several observations did not last the full 30 min, in which case we did not calculate path length and net displacement. In addition, we calculated a visibility index, measured as the proportion of locations from which the next location was visible to the lizard. We deemed a location to be visible to the lizard from the previous location if the line between the two locations was free of vegetation at the ground level (i.e., from the lizard’s line of sight). To measure

| Measurement       | Definition                                                                 |
|-------------------|---------------------------------------------------------------------------|
| Step Length       | The straight-line distance between consecutive 1-min locations.           |
| Path Length       | The sum of all step lengths for an observation period.                    |
| Net Displacement  | The straight-line distance between initial and final locations.           |
| Straightness Index| The ratio of net displacement to path length (value from 0 to 1).         |
| Turn Angle        | The change in direction between consecutive steps (value from 0° to 180°, with 0° = orientation of the focal animal at the previous step). |
visibility, a researcher sighted along the horizon just above the substrate over one location marker in the
direction of the next location marker, allowing us to assess what the animal could see from each vantage
point, and assess whether the direction taken had a clear or obstructed view. We recorded each sequential
step in the lizard’s path as either clear or obstructed, then determined the proportion of locations that were
clear for each observation (i.e., visibility index). We used air temperature and wind speed measures
obtained from a weather station adjacent to our site to estimate local conditions at each 1-min interval of
an observation.

**Data Analyses**

We used Minitab 18 (College Park, PA) for most analyses and R [49] for circular statistics, applying a
significance level of $P \leq 0.05$ to all tests. To examine the distributions of step length and turn angle as
well as habitat use patterns, we pooled the measurements by species. We tested data for normality and
applied non-parametric Mann-Whitney tests to pairwise comparisons of data that were not normally
distributed. We tested for species differences in habitat use and the distribution of step lengths and turn
angles using chi-square analyses. We applied circular statistics to comparisons of mean turn angles, and
to test for uniformity and conformity to von Mises distributions [50]. We examined within species
variation in step length using a mixed-effects model with lizard id as a random factor and sex, body size
(SVL), and weather variables as fixed factors. We did not observe any behavioral change in the lizards
during the duration or our observations in the field. We tested for a possible observer effect by using a
general linear model (GLM) to examine if animal movements varied over the course of the observation
period, with time in the observation period as our independent variable.

**Results**

**Movement patterns**

The two species differed in most characteristics of their movement paths including turn angle,
straightness index, step length, and path length (Table 2). The distribution of turn angles for both species
was significantly different from a uniform circular distribution (Rayleigh’s uniformity test: visual predator
(long-nosed leopard lizards, *G. wislizenii*): $z = 69.1$, df = 568, $P < 0.001$; chemosensory predator (western
whiptails, *A. tigris*): $z = 81.85$, df = 1004, $P < 0.001$). The distribution of turn angles and median turn angle
differed between the species (Chi-square test $X^2 = 10.6$, df = 3, $P = 0.014$; Fig. 1; Table 2), with the visual
predator making a greater proportion
of forward-directed movements and having smaller median turn angles. The strength of directionality (mean vector length) was 0.34 for the visual predator and 0.28 for the chemosensory predator. Leopard lizards moved less frequently than whiptails (visual predator: 44% vs chemosensory predator: 75% of intervals had movement; $z = 16.97$, $P < 0.001$). During intervals of movement, median step length was shorter for the visual predator (Table 2) and the distribution of step lengths differed between the species, with the visual predator being less likely to have longer step lengths (Chi-square test: $X^2 = 188.6$, df = 5, $P < 0.001$; Fig. 2).

Median path length also was shorter for the visual predator than for the chemosensory predator (Table 2), indicating that our chemosensory lizard moved greater overall distances during observations. However, median net displacement was comparable for both species (Table 2), indicating similarity in search area. Path straightness varied by species, with the visual predator traveling straighter paths (Table 2; Fig. 3). Turn angle and step length were not correlated for either species (visual predator: $r = 0.055$, $P = 0.090$; chemosensory predator: $r = -0.031$, $P = 0.474$). Step length for the chemosensory predator was greater for larger animals (SVL: $F_{1,41} = 9.77$, $P = 0.003$) and unrelated to sex, wind speeds or air temperatures. For the visual predator, step length was unrelated to sex, SVL or weather variables.

The movement pattern of an individual, as indicated by step lengths and turn angles, did not vary for either species over the course of an observation, indicating that our presence did not affect lizard behavior (GLM; $G. wislizenii$ step length: $F_{1,1714} = 0.02$, $P = 0.875$ and turn angle: $F_{1,476} = 0.4$, $P = 0.529$; $A. tigris$ step length: $F_{1,1446} = 0.3$, $P = 0.583$ and turn angle: $F_{1,911} = 0.43$, $P = 0.511$).

### Habitat Use

The two species varied in time spent under cover, with the visual predator occurring more frequently in the open than the chemosensory predator ($X^2 = 308.0$, df = 2, $P < 0.001$; Fig. 4). Based on differences in the visibility index, the visual predator was more likely than the chemosensory predator to move to locations visible from the previous location ($X^2 = 392.9$, df = 1, $P < 0.001$). In addition, the two species made use of

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**Table 2**

Summary of movement parameters (median (range)) for *G. wislizenii* (visual forager) and *A. tigris* (chemosensory forager), compared using Mann-Whitney tests.

| Measurement          | *G. wislizenii* | *A. tigris* | U     | P         |
|----------------------|-----------------|-------------|-------|-----------|
| Step Length (m)      | 2.1 (0.02–45.3) | 4.2 (0.01–42.8) | 1120731 | < 0.0001 |
| Path Length (m)      | 28.9 (0.0–203.1) | 92.3 (2.2–251.3) | 2537 | < 0.001 |
| Net Displacement (m) | 16.6 (0.0–116.5) | 21.5 (1.3–126.3) | 2005 | 0.243 |
| Straightness Index   | 0.663 (0.09–1.00) | 0.308 (0.02–0.92) | 1095 | < 0.001 |
| Turn Angle           | 44° (0–180) | 53° (0 – 180) | 820076 | 0.010 |
different species of plants. When under cover, the visual predator primarily used greasewood (*Sarcobatus vermiculatus*), but the chemosensory predator frequented both sage (*Artemisia tridentata*) and greasewood in roughly equal measure ($X^2 = 82.1$, df = 6, $P<0.001$; Fig. 5).

**Discussion**

Organisms occurring in sympatry must share or partition limiting resources to coexist. Frequently, sympatry involves species differing slightly in microhabitat, activity, or food usages. Our study shows a clear distinction between the movements and nuances in habitat use of a visual predator (*G. wislizenii*) that co-occurs with a predator foraging primarily by using chemical cues (*A. tigris*), demonstrating that movement can play a large role in facilitating sympatry and resource partitioning. Although both species exhibited similar net displacement in movement, their use of habitat and patterns of movement varied consistently along the lines of their differing prey detection strategies. The visually-oriented animals positioned themselves in areas where they were able to see prey from a distance, whereas the chemical-sensing species followed pathways indicative of a reliance on close inspection of chemical cues in and around bushes to acquire prey, differences that do not immediately follow traditional classifications of visual and chemosensory prey detection strategies.

In terms of min-to-min spatial advancement, the visually-oriented *G. wislizenii* showed different movement patterns compared to the chemosensory-oriented *A. tigris*. Specifically, the visual predator followed our prediction by moving in more direct paths and spending more time in the open when compared to the sympatric chemosensory species, a result consistent with the notion that visually-oriented predators require a clear visual path to stalk prey. In addition to different movement patterns, we found differences in vegetation use associated with the differing prey detection cues used by our two focal species. For visually-oriented predators, inhabiting more open areas of the habitat allows for visual scans of the edges of many plants, as well as the open area itself, which can increase their prey detection range. Chemosensory predators, by contrast, increase their chances of detecting prey by venturing further into vegetation. We recorded our chemosensory predator spending equal time in sage and greasewood bushes, climbing in brush to pursue insect prey, and digging for insect larva under vegetation, indicating that vegetation can harbor more prey opportunities than open areas but also requires more time to search. The equal occurrence of the chemosensory predator in both shrub types, combined with a consistently less direct trajectory, indicates that the whiptails were detecting chemosensory cues while moving from bush to bush, opportunistically searching each bush for prey. For our visual predator, the choice of vegetation might be related to the types of prey likely to be found therein, coupled with the vantage associated with chosen plants. Greasewood, for example, seemed to have a higher branch ceiling than sage (personal observation), possibly proving less of an impediment to visual scanning from a distance.

Our Alvord Basin study site allowed us to make comparisons of two species and draw novel insights into the interplay between movement and ecology for two sympatric species. While our approach of studying behavioral ecology through an assessment of movement patterns provides a template for examining
differences in behavior attributable to species, sex, body size, or season, we acknowledge that our observations might not generalize to other populations or have larger, species-level implications. Comparing the behavior of the same species in other locations where they occur would promote a more general appraisal of the factors affecting their behavior and sympatry.

**Conclusions**

Path segmentation combined with habitat analyses provides new insight into the ecology of visual and chemosensory prey detection. Individuals of our two species were commonly in proximity to each other, indicating a lack of spatial segregation by species. Yet, they used the same area in very different ways. Landscape features can present a continuum of corridor-barrier patches acting as functional areas whose use varies by species [51]. While *G. wislizenii* moved along a straight path to places that readily could be seen from the previous location (high visibility index), *A. tigris* moved more circuitously through vegetation where visibility was lower, demonstrating how landscape features and movement can interact to promote sympatry. Landscape features that facilitate movement for one species can impede movement for the other [51]. Depending on overlap in diet, differences in sensory priorities might not, by themselves, prevent competition between species. Their movement, however, could represent a form of resource partitioning that facilitates coexistence.

**Abbreviations**

GLM
general linear model

SVL
snout-to-vent length

**Declarations**

**Ethics approval**

All applicable national and institutional guidelines for the care and use of animals were followed under the approval of Erell Institute's Animal Care and Use Committee (IACUC proposal no. F2017-02).

**Consent for publication**

Not applicable

**Availability of data and materials**

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

**Competing interests**
The authors declare that they have no competing interests.

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**Author contributions**

MAE and DAE conceived and designed the study, collected and analyzed data and helped write the manuscript. EMB, KLU, JK, and KMD contributed to project design and all aspects of data collection and manuscript preparation, SGS and GEG collected data, contributed to field logistics, and provided editorial input on the manuscript.

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Figures

![Graph](image)

**Figure 1**
Distribution of turn angles for *G. wislizenii* (visual forager; black) and *A. tigris* (chemosensory forager; grey), separated into bins of 45°.

**Figure 2**

Distribution of step lengths for *G. wislizenii* (visual forager; black) and *A. tigris* (chemosensory forager; grey).
Figure 3

Representative paths from observed G. wislizenii (visual forager; A) and A. tigris (chemosensory forager; B). Grid lines for both plots are 5 m apart
Figure 4

Use of habitat by G. wislizenii (visual forager; A) and A. tigris (chemosensory forager; B)
Figure 5

Use of vegetation by G. wislizenii (visual forager, A) and A. tigris (chemosensory forager, B). Species of plants are in the families Chenopodiaceae (greasewood: Sarcobatus vermiculatus and shadscale: Atriplex confertifolia), Asteraceae (sage: Artemisia tridentate, rabbit brush: Ericameria sp., and horse brush: Tetradymia glabrata), and Poaceae (rye grass: Leymus cincereus).