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Seasonal distribution and routes of pronghorn in the northern Great Basin

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Across their range, pronghorn (*Antilocapra americana*) distribution and migratory movements are affected by human activities and infrastructure. Human barriers (e.g., roads, fences) can restrict pronghorn movements within or between seasonal ranges, limit daily movements, and reduce available habitat (Gates et al. 2012). Pronghorn have also experienced

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behavioral changes and reductions in carrying capacity across their range due to decreased habitat and fragmentation of migration routes as a result of development, fencing, and other land-use practices (e.g., Sawyer et al. 2005, Boccadori et al. 2008, Beckmann et al. 2012, Seidler et al. 2014). The challenge of conserving migratory ungulates is due to their wide-ranging seasonal movements, which occur across both a diversity of ecosystems and varying land management priorities (Copeland et al. 2014). For example, Berger (2004) estimated that 75% of the migration routes for bison (*Bison bison*), elk (*Cervus canadensis*), and pronghorn have been lost in the Greater Yellowstone region. And, although it is an important life history component for many species, migration is often overlooked during conservation planning efforts (Saher and Schmiegelow 2005, Berger et al. 2014).

Strong female site fidelity and social inheritance of learned travel routes appear to be important factors for pronghorn in maintaining access to discrete summer and fawning ranges (Barnowe-Meyer et al. 2013). These factors coupled with both low relative population abundance and disturbance, which contribute to the loss of experienced individuals, may prevent recolonization of unoccupied habitat patches, potentially leading to route abandonment and cessation of migration (Gustafson and Gardner 1996, Avital and Jablonka 2000, Piper 2011, Barnowe-Meyer et al. 2013). Further, although pronghorn do commonly display migratory behavior between distinct summer and winter ranges, and are adapted to moving long distances (O’Gara and Yoakum 2004), not all populations or individuals in those populations migrate annually with consistency; some populations or individuals may at times move only short distances between seasonal ranges (Hoskinson and Teter 1980, Boccadori and Garrott 2002, White et al. 2007, Jacques et al. 2009b). Although limited information is available (e.g., Kindschy et al. 1982, Trainer et al. 1983, Foster 1988), even fewer empirical investigations (e.g., Dalton 2009) have occurred within the western peripheral range of pronghorn occupying the northern Great Basin. This contributes to challenges for long-term conservation of pronghorn, because there is uncertainty about the distribution of seasonal habitats, location of migration routes, impacts of land-use practices, and effects of other potential barriers (Berger et al. 2006, White et al. 2007). There is also uncertainty in how large-scale conservation efforts underway for other species (e.g., Greater Sage-Grouse) could potentially overlap efforts to conserve pronghorn (Copeland et al. 2014). In this study, I documented the movements of adult female pronghorn in southeastern Oregon and northwestern Nevada during 2011–2013 to examine connectivity between 2 summering populations and to develop a baseline for identifying seasonal movement patterns and timing, seasonal distributions, individual fidelity to seasonal use areas, and migration distances and routes in the northern Great Basin.

**STUDY AREA**

Pronghorn were captured on summer ranges within Sheldon National Wildlife
Refuge (SNWR) and Hart Mountain National Antelope Refuge (HMNAR), managed jointly by the U.S. Fish and Wildlife Service, in northwestern Nevada and southeastern Oregon, respectively (Fig. 1A). The 2 national wildlife refuges were created in the early 1930s primarily as conservation areas for a declining pronghorn population. Together the refuges currently protect over 344,000 ha (3440 km²) of some of the last intact examples of the sagebrush-steppe ecosystem in North America. The 2 wildlife refuges jointly supported summer populations of roughly 3600 pronghorn (Collins 2014a, 2014b), and were an inset of the larger study area of approximately 1,525,665 ha (15,250 km²). My study area consisted almost entirely of public federal lands (90%) managed by the U.S. Fish and Wildlife Service (23%) and the Bureau of Land Management (67%). The Summit Lake Indian Reservation (Bureau of Indian Affairs) accounted for 0.3%, and the remaining roughly 10% of the lands within the study area were privately owned or county- or state-managed (Fig. 1A). Controlled, male-only harvest for pronghorn was permitted during the fall on public lands during the study. Season dates in Oregon were early August to mid-September, and in Nevada were early August to early September and late September to late October. Hunting in the larger area has generally been restricted to a relatively low number of limited tags with staggered seasons and dispersed hunting opportunities.

Elevations across the area range from 1320 m to 2640 m (HMNAR: 1448–2443 m; SNWR:...
1326–2183 m). Recent summer temperatures range from 0 °C to 34 °C and winter temperatures range between −29 °C and 14 °C; annual precipitation rarely amounts to more than 30 cm. Dominant vegetation consists of shrubs, particularly sagebrush (Artemisia spp.), and associated sagebrush-steppe habitats. Scattered open woodlands consisting of western juniper (Juniperus occidentalis) or curl-leaf mountain mahogany (Cercocarpus ledifolius) occupy ridgelines and some slopes. Talus and broken rock habitats are found along the edges of table tops and escarpments, and along steep side-slopes.

Methods

In early October 2011, we captured 39 adult (>2 years old) female pronghorn by using accepted capture protocols for net-gunning from a helicopter. Helicopter pursuit time was generally <2 min to limit capture-related mortalities, and pronghorn were processed at the capture site and released without transportation to another location to minimize handling stress (Jacques et al. 2009a). Total handling time (from pursuit to release) averaged 8.4 min per individual. Each captured pronghorn was briefly restrained, blindfolded, and monitored by a veterinarian during handling. Capture crews recorded capture location, sex, body condition, and an age estimate based upon incisor wear and replacement (Dow and Wright 1962). Blood samples were collected by venipuncture of the jugular vein for disease evaluation. Crews then attached to each pronghorn a store-on-board global positioning system (GPS) collar outfitted with a very high frequency (VHF) transmitter and a remote-release mechanism (Model G2110D, Advanced Telemetry Systems, Isanti, MN). Each collar was programmed to remotely detach after 2 years to allow recovery from the field.

The GPS collars were distributed widely across the 2 wildlife refuges in order to capture variation in movements (Fig. 1A). They were programmed to obtain location fixes every 5.5 h; 99% of the GPS fix attempts were successful. Each collar also recorded elevation (m) and temperature (°C) at each location. I also used average daily temperature (°C) and snow depth (cm) data averaged from 5 SNOTEL sites located in the vicinity of the study area (Natural Resources Conservation Service 2014) to compare overall winter severity across the study area between years. We located collared pronghorn via VHF-equipped fixed-wing aircraft within 2 weeks of capture and then approximately every 2–3 months to check their status and general location.

To categorize seasonal use, I used ArcGIS 10.0 (Environmental Systems Research Institute, Redlands, CA) to estimate the migration start and end date for each individual. I defined spring migration as seasonal movement from winter to summer range and fall migration as seasonal movement from summer to winter range. Exploratory movements were defined as movements observed during the fall migration period in which the individuals did not travel directly toward their eventual winter range, but initially traveled away from or bypassed through the wintering area before returning. I defined individual migration start dates as the first location that deviated ≥5 km from a seasonal range and was associated with successive directional movements (Kolar et al. 2011). To omit locations associated with migration or exploratory movements from seasonal range calculations, I defined seasonal ranges (winter and summer) as the time period 1 day after the last marked animal’s arrival date to 1 day prior to the first marked animal’s departure date.

I used Geospatial Modelling Environment (GME; Spatial Ecology 2013) and ArcGIS to calculate 95% and 50% kernel density estimates for individual annual summer and winter seasonal ranges. The 50% kernel density estimates from individual pronghorn by season were combined to estimate population-level seasonal core use areas. ArcGIS was used to determine the geographic center of seasonal ranges (95% kernel), and to measure the distance between geographic center points to estimate migration distances and seasonal range fidelity for individuals surviving 2 consecutive seasons (Jacques et al. 2009b). I noted a break in the overlap of seasonal ranges for pronghorn that migrated and for those that were nonmigratory, therefore I defined migration as seasonal movement between winter and summer ranges that overlapped in area by <15%. Nonmigrating individuals were defined by winter and summer ranges that...
overlapped by >15%. ArcGIS was also used to measure the straight-line distance between an individual’s consecutive locations, and those distances were summed per day to estimate daily travel distances.

I followed the approach of Horne et al. (2007) and Sawyer et al. (2009) to identify both individual and population-level routes used by pronghorn during the migration period. I used the Brownian bridge movement model package (BBMM; Nielson et al. 2013) for R (R Development Core Team 2013) to estimate a utilization distribution (UD) for individual pronghorn for each migration period using data collected from sequential GPS locations from collared animals. I used a grid size of 50 × 50 m and an estimated location error of 20 m because 94% of the GPS locations were 3-dimensional (3-D) fixes, which generally have <20 m of error (Di Orio et al. 2003, Sawyer et al. 2009). The UDs from individual pronghorn were combined to provide an estimate of the relative amount of use across a population-level route, and were categorized into quartiles such that the top 25% represented “high use” and the lowest 25% represented “low use” (Sawyer et al. 2009). Routes estimated by the BBMM provide measures of both time spent in an area and the rate of movement; therefore, high-use areas would be representative of where animals spent more time (e.g., foraging, resting), and moderate-use areas would represent where animals moved through more quickly (e.g., movement corridors) (Sawyer et al. 2009).

I compared pronghorn elevational use, migration distances, and home-range sizes between seasons and years using paired t tests (SYSTAT, Software Inc., Richmond, CA). I considered P ≤ 0.05 as statistically significant.

Results

Of the 39 adult female pronghorn that were collared originally, 5 either shed their collars (n = 3) or died (n = 2) within 45 d of capture and were not included in further analyses. No data were recovered from an additional 2 collars that malfunctioned due to unknown reasons. The 5% failure rate we experienced is at the lower limit of GPS device failures reported by Hebblewhite and Haydon (2010). The remaining 32 GPS collars collected 68,834 locations between October 2011 and October 2013. Individuals were followed for an average of 497 d (SD 218) and yielded an average of 2151 locations (SD 940).

Pronghorn that summered within Hart Mountain NAR and Sheldon NWR demonstrated wide variation in directional movements to winter ranges (Fig. 1A). Pronghorn captured on Hart Mountain NAR in Oregon were the most varied and far-ranging in their movement patterns, and individual animals were observed (1) remaining on HMNAR during winter; (2) moving to winter ranges in Oregon to the east, (3) moving south to winter ranges located in Oregon between the 2 refuges, or (4) continuing to move into Nevada to winter either on Sheldon NWR or farther south. For pronghorn captured on Sheldon NWR, individuals were observed (1) remaining on SNWR during winter, (2) moving north to winter ranges in Oregon, or (3) moving to winter ranges south of SNWR in Nevada. Pronghorn departed their summer ranges between late September (first departure 28 September) and late November, arriving at winter ranges by mid-December (last arrival 15 December). In the spring, pronghorn left their winter ranges between early March (first departure 7 March) and early May, arriving at summer ranges by mid-May (last arrival 13 May). The median departure date for fall migration was 11 October (SD 14.2 d) and for spring migration was 8 April (SD 18.9 d). Very few (≤3) marked individuals had the same initiation date for any given migration period and none were observed traveling together during periodic aerial surveys; therefore, I assumed movement was independent of other collared pronghorn (Saher and Schmiegelow 2005). Previous research efforts had identified the pronghorn fawning season for the study area as 11 May–14 June (U.S. Fish and Wildlife Service unpublished data). Seasonal dates were identified as 16 December–6 March for winter ranges and 11 May–27 September for summer ranges (adjusted to include the start of fawning).

Overall, GPS-collared pronghorn moved an average of 4.7 km · d−1 (SD 3.3). Movement rates for individual pronghorn peaked during fall migration, averaging 5.8 km · d−1 (SD 1.1, n = 32) for fall 2011 and 6.2 km · d−1 (SD 1.8, n = 24) for fall 2012. Female pronghorn moved the least during the fawning season, averaging the same distances for both 2012
To adjacent ranges was 10.3 km (SD 5.1, n = 6), and average distance for those that migrated to nonadjacent ranges was 30.0 km (SD 15.6, n = 20). In contrast, in 2012/2013 all of the remaining collared individuals exhibited migration movements to nonadjacent ranges (x̄ = 39.1 km, SD 29.4, n = 19), including 9 pronghorn who either did not migrate (n = 4) or only migrated short distances to adjacent ranges in the previous year (n = 5). Average distance between summer and winter seasonal ranges was significantly farther in 2012/2013 (x̄ = 44.9 km, SD 27.1, n = 19) than in 2011/2012 (x̄ = 23.1 km, SD 16.0, n = 31) (t = −3.584, df = 48, P = 0.001). The farthest distance documented was 160.2 km for an individual during the fall 2012 migration between summer range on Hart Mountain NAR in Oregon and winter range south of Sheldon NWR in Nevada.

Utilization distributions (UDs) were estimated for 60 individual routes during the 4 migration periods. Individuals who did not migrate or whose distance-moved was too short to identify a discrete route were excluded from this analysis. The average Brownian motion variance (BMV) outputs, which quantify how diffusive or irregular the path of an animal is (Sawyer et al. 2009), were as follows: fall 2011, 13,055 m² (SE 8144, n = 23 migrations); spring 2012, 6558 m² (SE 3800, n = 16 migrations); fall 2012, 10,373 m² (SE 5512, n = 16 migrations); and spring 2013, 8791 m² (SE 3149, n = 5 migrations). These calculations included data from individuals who were either nonmigratory or short-distance migrants in 2011/2012 but who made exploratory movements during the fall migration period before returning to wintering areas either within or adjacent to their respective summer ranges. Indeed, 76% of migrating individuals made exploratory movements—sometimes very long-distance ones (e.g., >260 km)—during the fall migration period before returning.

**Table 1.** Summary of average elevations and seasonal area sizes used by female pronghorn, southeastern Oregon and northwestern Nevada, 2011–2013.

<table>
<thead>
<tr>
<th>Time period</th>
<th>Elevation (m)</th>
<th>95% range (km²)</th>
<th>50% range (km²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>x̄</td>
<td>SD</td>
</tr>
<tr>
<td>Winter 2011/2012</td>
<td>31</td>
<td>1778.3</td>
<td>84.2</td>
</tr>
<tr>
<td>Summer 2012</td>
<td>29</td>
<td>1834.7</td>
<td>133.7</td>
</tr>
<tr>
<td>Winter 2012/2013</td>
<td>20</td>
<td>1565.9</td>
<td>82.4</td>
</tr>
<tr>
<td>Summer 2013</td>
<td>12</td>
<td>1808.2</td>
<td>139.4</td>
</tr>
</tbody>
</table>

(\(\bar{x} = 3.8\) km, SD 1.3, n = 29) and 2013 (\(\bar{x} = 3.8\) km, SD 0.9, n = 12). The greatest distance moved in a single day was 37.9 km by an individual returning from winter to summer ranges in early May 2012. The shortest distance was 0.18 km·d\(^{-1}\) during fawning in early June 2012.

For pronghorn that survived at least through the start of summer 2012 (n = 29), only 2 (7%) did not return to the same summer range on which they were originally captured in October 2011. Individuals that survived 2 consecutive summers (2012 and 2013) demonstrated stronger fidelity to summer ranges than to winter ranges, returning to within 5.7 km, on average, of the previous summer range (SD 5.4 km, n = 12). There was no difference in the size between years for either 95% or 50% summer ranges (95% t = 0.569, df = 36, P = 0.573; 50% t = 0.478, df = 36, P = 0.636) (Table 1). Therefore I pooled the summer range data among years for individuals for subsequent analyses. In contrast, individuals demonstrated weaker fidelity to winter ranges, averaging 39.1 km between consecutive winter ranges (SD 29.4 km, n = 19). The mean 95% winter range size was also larger in 2012/2013 than in 2011/2012 (t = −1.974, df = 48, P = 0.054); there was no difference in 50% winter range sizes between years (t = −1.618, df = 48, P = 0.112; Table 1). In addition to increases in the size of the area used, there was also an overall locational shift in the wintering areas to the east between 2011/2012 and 2012/2013 (Fig. 2A).

During 2011/2012, five of 31 pronghorn did not migrate (16%); 1 individual shed its collar during late fall migration and thus was not included in this analysis. Of the remaining individuals, 20 (65%) made seasonal migrations between summer and winter ranges and 6 (19%) made short-distance (<20 km) migrations to adjacent seasonal areas. Average distance for those who migrated (i.e., seasonal ranges overlapped by <15%) short distances to adjacent ranges was 10.3 km (SD 5.1, n = 6), and average distance for those that migrated to nonadjacent ranges was 30.0 km (SD 15.6, n = 20). In contrast, in 2012/2013 all of the remaining collared individuals exhibited migration movements to nonadjacent ranges (x̄ = 39.1 km, SD 29.4, n = 19), including 9 pronghorn who either did not migrate (n = 4) or only migrated short distances to adjacent ranges in the previous year (n = 5). Average distance between summer and winter seasonal ranges was significantly farther in 2012/2013 (x̄ = 44.9 km, SD 27.1, n = 19) than in 2011/2012 (x̄ = 23.1 km, SD 16.0, n = 31) (t = −3.584, df = 48, P = 0.001). The farthest distance documented was 160.2 km for an individual during the fall 2012 migration between summer range on Hart Mountain NAR in Oregon and winter range south of Sheldon NWR in Nevada.
and settling on their respective winter ranges (e.g., Fig. 2B). Though exploratory movements most often occurred during fall 2011, two individuals also made such movements during fall 2012; similar exploratory movements were not observed during the spring migration period.

At the population level, collared pronghorn used a network of common routes during the migration periods to access and return from their respective winter ranges (Fig. 1B). The population-level routes represent a probabilistic measure of where migration and exploratory movements likely occurred, characterized by high-use areas, where pronghorn spent the majority of their time, connected by moderate-use areas through which pronghorn moved more quickly (Sawyer et al. 2009; Fig. 1B). Overlap of seasonal ranges with population-level routes was high, and influenced by the wide variation in directional movements observed for both individual pronghorn and between years. Depending on the individual and the year, these common areas were utilized for both winter and summer ranges, but also for routes associated with migration and exploratory movements as individuals bypassed one another. As an example, pronghorn #16445 summered on Sheldon NWR in Nevada and wintered in 2011/2012 on an adjacent range in Oregon just to the north of her summer range (Fig. 2B). However, the following year in fall

Fig. 2. (A) Estimated population-level seasonal core use areas (50% kernel) for summer ranges (green) and winter ranges (2011/2012 [purple] and 2012/2013 [blue]). (B) Example exploratory movements made by date between summer range (geographic center, green) and 2012/2013 winter range (geographic center, blue) during the fall 2012 migration period of pronghorn #16445, southeastern Oregon and northwestern Nevada.
2012, she first left her summer range and traveled in the opposite direction from the previous year, to the south of Sheldon NWR in Nevada. After roughly 3 weeks, she returned north and spent a little over a month in the vicinity of her previous winter range in Oregon before continuing farther north, traveling through Hart Mountain NAR and settling on a winter range to the east of the refuge (Fig. 2B).

Collared pronghorn on average occupied similarly high elevations during both summers (Table 1). Pronghorn then moved to lower elevational winter ranges, although individuals occupied significantly lower elevations in winter 2012/2013 compared to winter 2011/2012 ($t = 8.863$, $df = 49$, $P < 0.000$) (Table 1). Recorded ambient temperatures from the collars documented differences between the 2 winters, with temperatures in 2012/2013 being on average slightly cooler ($\bar{x} = 6.6$ °C [43.8 °F], SD 6.7) than temperatures in 2011/2012 ($\bar{x} = 7.5$ °C [45.4 °F], SD 5.7). Temperatures averaged from the SNOTEL sites further support that the winter of 2012/2013 was cooler ($\bar{x} = -2.57$ °C [27.4 °F]) than the previous winter ($\bar{x} = -0.86$ °C [30.4 °F]) (Natural Resource Conservation Service 2014). The winter of 2012/2013 also recorded higher snow depths overall, averaging 38.2 cm compared to 29.1 cm in 2011/2012; all of the 5 SNOTEL sites recorded an increase in snow depth of 17%–45% between the 2 years (Natural Resources Conservation Service 2014). Though snow accumulation during 2012/2013 was greater overall, the first date of measurable snowfall in 2011/2012 occurred nearly a month earlier (6 October 2011 vs. 24 October 2012) (Natural Resources Conservation Service 2014).

**DISCUSSION**

Pronghorn in this study migrated from separate summering areas to a range of wintering areas in late September to mid-December, returning to summer ranges between early March and mid-May. Migration in this study generally coincided with migration timing reported for other pronghorn populations across their range (Oregon—Trainer et al. 1983, Foster 1988; Idaho—Hoskinson and Tester 1980; South Dakota—Jacques 2006; Wyoming—Sheldon and Lindzey 2006; Yellowstone—White et al. 2007). Distances traveled by pronghorn between winter and summer ranges can differ greatly among populations and between years. Pronghorn in this study traveled migratory distances comparable to those reported in South Dakota ($\bar{x} = 23.1$ km; Jacques et al. 2009b), southern Idaho and Montana ($\bar{x} = 44.5$ km; Hoskinson and Tester 1980), and southeastern Oregon (30–40 km; Dalton 2009). One individual during this study traveled >160 km, which is within the range of some of the longest seasonal migrations (116–258 km) reported for pronghorn (Wyoming; Sawyer et al. 2005, Berger and Cain 2014).

Female pronghorn generally demonstrate stronger fidelity to summer ranges and weaker fidelity to winter ranges (Deblinger and All dredge 1984, Kolar et al. 2011, this study). Female pronghorn occupying sagebrush habitats of the northern Great Basin also exhibited a mixture of migration strategies. Not all pronghorn migrated similarly from the same winter ranges, or even with respect to their own previous migrations or travel routes. The individuals in this study that either did not migrate or only migrated a short distance the first year, but migrated long distances the following year support observations by White et al. (2007) and Jacques et al. (2009b) that migration in some pronghorn populations may be a conditional strategy. The proportion of pronghorn that migrated in this study ranged from 65% in 2011/2012 to 100% in 2012/2013 during harsher winter conditions; although the majority also made exploratory movements during fall 2011. Pronghorn may avoid adverse conditions by differing their migration strategy across years (e.g., Bruns 1977, Hoskinson and Tester 1980, Barrett 1984, Kolar et al. 2011). The proportion of migratory individuals within a population can vary greatly among regions, ranging from 8%–19% (Jacques et al. 2009b) to 70%–81% of individuals migrating annually (White et al. 2007). Such partial migration strategies are associated with costs to the individual weighed against increased access to higher-quality resources, and have been documented in several ungulate species including pronghorn, elk, and mule deer (*Odocoileus hemionus*) (Bergerud et al. 1990, Nicholson et al. 1997, White et al. 2007).

Across their range, pronghorn appear to respond to several key landscape features including physiography, vegetation, and movement barriers when selecting seasonal home ranges and migratory routes. Pronghorn...
home-range sizes are also highly variable. Previous estimates for adult pronghorn have ranged from 11.9 to 779.8 km² (Wright and deVos 1986, Clemente et al. 1995, O’Gara and Yoakum 2004). Such variable home-range sizes are likely related to differences in habitat quality, population and group sizes, land-use practices, and seasonal requirements (Kitchen and O’Gara 1982). In general, pronghorn seasonally tend to use higher elevations during summer months, and then descended to lower elevations during the winter (Hoskinson and Tester 1980, Sawyer et al. 2005, White et al. 2007, this study). Pronghorn may make such seasonal elevational shifts to avoid deep snow due to difficulties encountered in both foraging and traveling. In the Great Basin, prolonged snow accumulation over 25–30 cm has been suggested to be detrimental to pronghorn when associated with low quantity and/or low quality of forage and barriers to movement (Sundstrom 1969, Riddle and Oakley 1973, Bruns 1977, Kindschy et al. 1982). When snow is present, pronghorn seek areas that have blown free or areas with low snow cover through which forage is accessible (Bruns 1977). Pronghorn have also been observed to make long-distance movements after a heavy snowfall in search of better forage accessibility (e.g., Guenzel et al. 1982, Sawyer and Lindzey 2000).

Typically, outside of elevational-related migration, pronghorn generally move north in the spring and south in the fall to warmer winter ranges (Sawyer et al. 2005, Sheldon and Lindzey 2006, Kolar et al. 2011). In contrast, pronghorn in the northern Great Basin demonstrated wide variation in directional movements between seasonal ranges and between years. Overall, marked pronghorn did follow a north–south gradient within the study area. However the directionality was not consistent (i.e., some individuals moved south in the fall and north in the spring, others moved south in the spring and north in the fall, and yet others explored in both directions during the fall migration period). As such, there was high overlap of common areas utilized by individual pronghorn across seasons and time periods, which additionally highlights these as important use areas. Individuals using routes during the fall migration period frequently, and sometimes repeatedly, bypassed each other while traveling in different directions, both through movement areas and through summer and winter ranges. Indeed, the majority of individual pronghorn made exploratory movements during the fall 2011 migration period, likely in response to an earlier onset of snow accumulation. Other studies have also identified individuals with extended migration periods (e.g., Sheldon and Lindzey 2006), presumably during which those pronghorn were making similar exploratory movements. In contrast, in 2012/2013, the onset of snow was later, but snow accumulated more quickly. Subsequently, only 2 individuals made long-distance exploratory movements during the fall 2012 migration period before settling on their respective winter ranges. Kolar et al. (2011) hypothesized that milder winters (i.e., warmer temperatures with less snow cover) would result in decreased migration distances. This hypothesis is supported by findings in this study where during 2012/2013, fall migration distances were longer with fewer exploratory movements compared to distances observed during 2011/2012 in more mild winter conditions. Winter ranges were also larger, lower in elevation, and shifted in location during 2012/2013, again corresponding to winter conditions that were on average colder with deeper snow.

Migratory ungulates are declining worldwide, and a broader conservation effort that includes recognizing migration routes as critical habitat is needed (Berger 2004, Bolger et al. 2008, Sawyer et al. 2009, Berger et al. 2014). The cumulative effects of climate change, habitat fragmentation, encroachment of invasive species, woodland expansion, increasing fire frequency and severity, and competition for water and forage resources with livestock and free-roaming horses are all challenging and ongoing conservation issues in the northern Great Basin (Miller et al. 2011, Manier et al. 2013). As such, successful long-term conservation of migratory pronghorn populations, both in this region and across their distribution, should include managing for not only the range of important seasonal use areas, but also maintaining connectivity along routes. Connectivity within this population is high and presently there appear to be no substantial barriers to pronghorn movement in this portion of the northern Great Basin. However, gathering the necessary background information is the first crucial step for
long-term conservation (Berger and Cain 2014), particularly for a species that demonstrates such variability across its range. By identifying the distribution and timing of pronghorn use across the larger landscape, land managers now have a baseline through which a prioritization process can be developed to identify those areas and routes of highest conservation value (Sawyer et al. 2009, Copeland et al. 2014). And although overlap with large, protected lands (e.g., national wildlife refuges) was high in this instance, it is clear that these areas are not large enough to encompass the entirety of this migratory species’ needs. Future obstacles to migration could potentially extirpate routes and contribute to a loss of connectivity (Copeland et al. 2014, Seidler et al. 2014). Partnerships across land management agencies and other stakeholders in applying this knowledge to ongoing landscape-level conservation planning will be necessary to mitigate any future cumulative changes that have the potential to impact important pronghorn habitat or alter connectivity.

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