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DO FEMALE ELK SELECT HIDING COVER FOR THEIR CALVES AT BIRTH SITES?

Dillon T. Fogarty¹,² and Chadwick P. Lehman¹

Abstract.—In light of local population declines and poor calf survival, we investigated visual obstruction as potential elk (Cervus elaphus) calf hiding cover at birth sites. We hypothesized that female elk would use hiding cover (i.e., visual obstruction) in order to conceal their calves at the birth site and reduce vulnerability to predation. Selection of hiding cover has been well documented at calf bedding sites subsequent to parturition; however, because of variably reported results from prior studies, it is unclear whether hiding cover is selected at the immediate birth site. We estimated visual obstruction of hiding cover, measured understory vegetation height, and categorized the dominant form of hiding cover at 65 birth sites and 116 random sites within the Black Hills, South Dakota. We found no significant differences in visual obstruction, vegetation height, or selection of dominant hiding cover categories between birth and random sites within forest and grassland systems. We surmise that selection of birth sites with greater visibility was used as a strategy to increase detectability of potential human and puma (Puma concolor) disturbances, which may be hindered by understory vegetation or woody debris that obstructs visibility. This may explain why we did not detect any selection for hiding cover; thus, understory cover for calves may not be a driving factor for selection of birth sites.

Note.—The equivalent of 1 metric ton (t) is 1000 kg or 1 Mg.

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North American elk (Cervus elaphus; hereafter elk) exhibit K-selected life history traits, as adult survival rates tend to be high and stable whereas variable calf recruitment rates can affect herd dynamics (Gailliard et al. 1998, 2000). The majority of neonate mortality is caused by predation (Linnell et al. 1995, Smith and Anderson 1996, Singer et al. 1997), and mortality rates are highest during the first weeks of life and decrease as neonates age (Geist 1982, Singer et al. 1997, Barber-Meyer et al. 2008, Lehman 2015). To mitigate predation risk, elk select habitat resources that benefit their antipredator strategies ( Lima and Dill 1990, Mysterud and Østbye 1999, Mao et al. 2005). At birth sites and nearby areas, risk of predation to the adult female and neonate may be reduced with habitat selection strategies (adult—Barbknecht et al. 2011, Rearden et al. 2011, Lehman et al. 2016; neonate—Pitman et al. 2014).

Primary factors that contribute to birth site selection by ungulates include human disturbance, forage resources, and predator avoidance (Thomas et al. 1979, Bowyer et al. 1999, Barbknecht et al. 2011, Rearden et al. 2011, Bercovitch and Berry 2015, Lehman et al. 2016). Oftentimes fine-scale selection of birth sites reflects antipredation strategies of the adult (Bowyer et al. 1999, Barten et al. 2001,
Poelke et al. 2007, Barbknecht et al. 2011, Rearden et al. 2011, Smith et al. 2015, Lehman et al. 2016. Antipredation strategies that optimize detection of oncoming threats such as humans or predators can be inversely related to forage abundance. In such cases, moose (Alces alces) cope with these situations by selecting areas that provide more visibility but also the necessary forage to meet nutritional needs (Bowyer et al. 1999, Poole et al. 2007), whereas elk select forage resources at larger scales and increased visibility at smaller scales (Rearden et al. 2011, Lehman et al. 2016). Although, elk in Wyoming utilized hiding cover on all scales and, similar to other studies, selected forage at larger scales (Barbknecht et al. 2011). Another factor that can contribute to selection of birth sites is proximity to anthropogenic features such as roads or housing developments. For example, moose in Grand Teton National Park selected birth sites closer to paved roads as risk of predation increased (Berger 2007). Alternatively, bighorn sheep (Ovis canadensis) in the Black Hills selected birth sites that were farther from roads and housing developments to avoid human disturbances (Smith et al. 2015).

Ungulates may select habitat to either optimize visual detection of oncoming threats or increase hiding cover. Both strategies attempt to avoid predation (Lima and Dill 1990, Mysterud and Østbye 1999). During birth and shortly thereafter, any detection of the neonate by a predator drastically increases the risk of predation (Roberts and Rubenstein 2014). Therefore, it is not surprising that during the hiding phase, neonatal young are heavily reliant on hiding cover (Geist 1982, Alldredge et al. 1991, Shallow et al. 2015). Pitman et al. (2014) found that increased cover was important at bedding sites of young elk calves and that adult females selected areas that facilitated the hiding cover needs of the calf. Similarly, Bongi et al. (2008) found that roe deer (Capreolus capreolus) altered their habitat use to provide more cover to neonates. Wallace and Krausman (1990) found that elk calves were usually within a meter of hiding cover and that the type of cover used varied with habitat type. Additionally, Shallow et al. (2015) was able to link concealment cover at mule deer (Odocoileus hemionus) bed sites to increased fawn survival. Given its importance, it is not surprising that hiding cover selection has been widely documented for many species such as white-tailed deer (Odocoileus virginianus; Huegel et al. 1986), mule deer (Gerlach and Vaughan 1991), pronghorn (Antilocapra americana; Canon and Bryant 1997), roe deer (Linnell et al. 1999), and bighorn sheep (Smith et al. 2015).

Our South Dakota study population of elk has been declining since 2006, a trend that is concerning for game managers and hunters (South Dakota Department of Game, Fish and Parks 2015). Our objective was to evaluate resource selection characterizing calf hiding cover at the fine spatial scale at and immediately adjacent to the birth site. Based on previous research that highlights the importance of hiding cover to neonates (Wallace and Krausman 1990, Pitman et al. 2014, Shallow et al. 2015), we hypothesized that to conceal their neonates, female elk would select birth sites with more hiding cover provided by vegetation or woody debris. Further, we hypothesized that even if adult elk desired increased visibility within the 50+ meters surrounding the birth site they would still be able to select hiding cover immediately adjacent to the birth site.

We conducted our research in Custer and Pennington Counties in southwestern South Dakota in the southern Black Hills physiographic region (Flint 1955). Land ownership was mixed private and public land, including Custer State Park (CSP) which encompassed 286 km² in the central part of the study area. Elevations ranged from 1108 to 2208 m. Adult female elk were sedated with tranquilizer darts (butorphanol, azaperone, and medetomidine sedation protocol; Mich et al. 2008) deployed from helicopters (QuickSilver Air Inc., Peyton, CO) during February 2012 and 2013. After elk were sedated, we blindfolded and fitted them with satellite radio collars (Telonics Inc., Mesa, AZ.; Advanced Telemetry Systems Inc., Isanti, MN). We aged elk by extracting an upper canine tooth and evaluating cementum annuli (Matson’s Lab, Milltown, MT; Hamlin et al. 2000). Female elk were inspected for pregnancy using rectal palpation (Greer and Hawkins 1967, Vore and Schmidt 2001). Females suspected of being pregnant were fitted with a vaginal implant transmitter (Advanced Telemetry Systems, Isanti, MN). Vaginal implant transmitters (VITs) were used to determine locations of birth sites (Barbknecht et al. 2009). A veterinarian was
on hand to extract teeth, test for pregnancy, and implant transmitters.

We located female elk using radiotelemetry daily from 1 April to 31 October. From 1 November to 31 March, elk were located 5 days per week. Once adult females isolated themselves from others and began to localize, we checked VIT signals twice daily in morning and evening. After birth, expelled VITs were retrieved and their locations were recorded in UTM coordinates with a handheld GPS unit. We collected data from birth sites with substantial evidence of calving. In most cases we observed a calf with the radio-marked cow immediately adjacent to the birth site. Other evidence of the birthing event typically observed near the expelled VIT included placenta material and/or blood at a cleared spot on the ground, as well as a strong odor, moistened soil, and fresh fecal pellets. If VITs were expelled outside the calving season, or if we did not observe evidence of the birthing event, we did not collect data from those sites.

Within a 100% minimum convex polygon of all female elk locations, we used stratified random sampling (Cochran 1977) to select sites at which to quantify characteristics of the vegetation available to female elk. See Lehman et al. (2016) for detailed methodology on stratified random sampling and vegetation classification. Strata for the random sampling were vegetation structural stages (Buttery and Gillam 1983) described in Lehman et al. (2016). Each year, we randomly selected 5–10 polygons for each stratum of the GIS vegetation coverage without replacement to ensure a comprehensive allocation of random points and sufficient sampling of all the vegetation communities across the study area. We generated one random point in each of these polygons.

We quantified vegetative characteristics at birth sites 3–5 days after the female and calf left the area. We also measured and recorded vegetation characteristics at the random sites during the same period through 31 August each year. We determined understory vegetation characteristics at random points and at birth sites. We estimated percent canopy cover of total herbaceous cover, grasses, forbs, and shrubs in a 0.1-m² quadrat (Daubenmire 1959). We used 4 transects following cardinal directions radiating out from sites and measured percent canopy cover at 2-m intervals (n = 40); we averaged data collected along transects for each site. We interpolated downed woody debris (metric tons per hectare = t·ha⁻¹) using a pictorial guide (Simmons 1982). We used a modified Robel pole marked with alternating colors at 2.54-cm increments (Robel et al. 1970, Benkobi et al. 2000) to characterize visual obstruction. This method requires the observer to kneel to a height of 1 m and record the lowest visible increment on the pole from a distance of 4 m in each of the cardinal directions (n = 4).

Visual obstruction measurements were centered at either the birth site or random point and then taken at 5-m intervals in each cardinal direction (n = 4). We chose to use 5-m intervals to quantify the visual obstruction that was immediately available to the neonate after birth. We did not include data at the center point of birth or random sites because the birth site often had vegetation removed or disturbed by the cow elk during the birthing event. From each point where visual obstruction readings were taken, we also recorded vegetation height. We classified dominant understory cover at sites using percent canopy cover and woody debris data. When vegetation had <20% canopy cover and woody debris was <17.3 t·ha⁻¹ the site was considered bare. Further, when vegetation canopy cover was ≥20% and/or woody debris was ≥17.3 t·ha⁻¹ several dominant cover categories were considered and included: grass/forb, shrub, woody debris, woody debris/grass/forb, and woody/shrub.

We used the multiple-response permutation procedure (MRPP; Mielke and Berry 2001) to test the hypothesis that visual obstruction and vegetation height does not occur randomly across the landscape. MRPP is a distribution test based on Euclidean distance of the hypothesis that the data sets are from the same population. MRPP tests relax the parametric structure requirement of a test statistic and are less affected by an extreme measurement of a single object. We used Pearson chi-square tests to compare dominant cover categories between birth sites and random sites. Significance for all tests was determined at α = 0.05. All statistical analyses were performed using SPSS Statistics (Version 21, IBM Corporation 2012).

We used birth sites from 36 radio-marked female elk (n = 35 adults, n = 1 yearling) in
our analyses, resulting in 65 birth sites measured over 2 years. We also collected data from 116 random sites. We evaluated selection separately in pine forests (n = 29 birth sites, n = 96 random sites) and grasslands (n = 36 birth sites, n = 20 random sites). Visual obstruction did not differ (MRPP test statistic ≤ 0.33, \( P \geq 0.25 \)) between birth sites and random sites in forests (Table 1) or grasslands (Table 2). Vegetation height did not differ (MRPP test statistic ≤ 0.95, \( P \geq 0.13 \)) between birth sites and random sites in forests (Table 1) or grasslands (Table 2). Selection of dominant cover categories did not differ (\( \chi^2 \leq 6.68, df = 4 \) or 5, \( P \geq 0.25 \)) from random sites, and selection ratios indicated no selection for any categories (Tables 1, 2).

We found no support for our hypothesis that elk select more hiding cover for their neonates at birth sites. On the basis of previous research that highlights the importance of hiding cover, our results were unexpected. Young ungulates, whose only defense is to hide, select areas for themselves with more hiding cover (Gerlach and Vaughan 1991, Canon and Bryant 1997, Linnell et al. 1999), including elk (Wallace and Krausman 1990, Pitman et al. 2014). Given the importance of hiding cover away from the birth site, we predicted that female elk would select hiding cover for their calves at the birth site; however, this selection was not found.

We suspect that adult elk are exposed to high risk at the birth site, and due to the

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**Table 1.** Visual obstruction readings (VO), vegetation height (VH), and number of sites by cover categories for birth sites and random sites, as well as selection ratios (95% CIs) for cover categories in the pine forest of the southern Black Hills, South Dakota, 2012–2013.

<table>
<thead>
<tr>
<th>Cover category</th>
<th>Number of birth sites</th>
<th>Number of random sites</th>
<th>Selection ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare(^a)</td>
<td>4</td>
<td>11</td>
<td>1.19</td>
<td>−0.79 to 3.17</td>
</tr>
<tr>
<td>Grass_Forb(^b)</td>
<td>5</td>
<td>32</td>
<td>0.53</td>
<td>−0.09 to 1.14</td>
</tr>
<tr>
<td>Shrub(^c)</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>0.00 to 0.00</td>
</tr>
<tr>
<td>Woody(^d)</td>
<td>4</td>
<td>19</td>
<td>0.69</td>
<td>−1.8 to 1.56</td>
</tr>
<tr>
<td>Woody_Grass_Forb(^e)</td>
<td>13</td>
<td>28</td>
<td>1.52</td>
<td>0.43 to 2.62</td>
</tr>
<tr>
<td>Woody_Shrub(^f)</td>
<td>3</td>
<td>4</td>
<td>2.46</td>
<td>−2.3 to 7.23</td>
</tr>
<tr>
<td>Total</td>
<td>36</td>
<td>96</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\) Bare ground was <20% understory canopy cover of vegetation and <17.3 t·ha\(^{-1}\) of woody debris.

\(^b\) Grass_Forb category was ≥20% grass and forb canopy cover and <17.3 t·ha\(^{-1}\) of woody debris.

\(^c\) Shrub category was ≥20% shrub canopy cover and <17.3 t·ha\(^{-1}\) of woody debris.

\(^d\) Woody category was ≥17.3 t·ha\(^{-1}\) of woody debris and <20% canopy cover of vegetation.

\(^e\) Woody_Grass_Forb category was ≥17.3 t·ha\(^{-1}\) of woody debris and ≥20% canopy cover of grasses and forbs.

\(^f\) Woody_Shrub category was ≥17.3 t·ha\(^{-1}\) of woody debris and ≥20% canopy cover of shrubs.

**Table 2.** Visual obstruction readings (VO), vegetation height (VH), and number of sites by cover categories for birth sites and random sites, as well as selection ratios (95% CIs) for cover categories in grasslands of the southern Black Hills, South Dakota, 2012–2013.

<table>
<thead>
<tr>
<th>Cover category</th>
<th>Number of birth sites</th>
<th>Number of random sites</th>
<th>Selection ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare(^a)</td>
<td>4</td>
<td>3</td>
<td>0.74</td>
<td>−0.59 to 2.07</td>
</tr>
<tr>
<td>Grass_Forb(^b)</td>
<td>17</td>
<td>8</td>
<td>1.18</td>
<td>0.2 to 2.16</td>
</tr>
<tr>
<td>Shrub(^c)</td>
<td>1</td>
<td>3</td>
<td>0.19</td>
<td>−0.36 to 1.73</td>
</tr>
<tr>
<td>Woody(^d)</td>
<td>1</td>
<td>2</td>
<td>0.29</td>
<td>−0.59 to 1.15</td>
</tr>
<tr>
<td>Woody_Grass_Forb(^e)</td>
<td>13</td>
<td>4</td>
<td>1.81</td>
<td>−0.51 to 4.12</td>
</tr>
<tr>
<td>Total</td>
<td>36</td>
<td>20</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\) Bare ground was <20% understory canopy cover of vegetation and <17.3 t·ha\(^{-1}\) of woody debris.

\(^b\) Grass_Forb category was ≥20% grass and forb canopy cover and <17.3 t·ha\(^{-1}\) of woody debris.

\(^c\) Shrub category was ≥20% shrub canopy cover and <17.3 t·ha\(^{-1}\) of woody debris.

\(^d\) Woody category was ≥17.3 t·ha\(^{-1}\) of woody debris and <20% canopy cover of vegetation.

\(^e\) Woody_Grass_Forb category was ≥17.3 t·ha\(^{-1}\) of woody debris and ≥20% canopy cover of grasses and forbs.
hunting tactics of a major predator in our study area, the adult and neonate’s predator avoidance strategies were in opposition to each other. Further, we surmise that this conflict resulted in a compromise of neonate hiding cover in order to maximize the security of the adult. Annual survival rates of calves were $\leq 27\%$ for 3 years of the study, and $26\%$ of predation mortality occurred within 5 days of birth (Lehman 2015), or during the hiding phase (Geist 1982). Most calf mortality was from puma (Puma concolor) predation (Lehman 2015). To mitigate their own predation risk, adult female elk can select for optimal security cover (Thomas et al. 1979, Lehman et al. 2016), which may vary depending on the hunting tactics of major predators. For instance, coursing predators like grizzly bears (Ursus arctos) and wolves (Canis lupus) prefer hunting in open areas where they can perceive prey at long distances; in contrast, puma prefer dense concealment which allows them to stalk and ambush prey in close quarters (Mech 1970, Beier et al. 1995). Avoidance of areas that favor the hunting strategies of major predators is a common antipredation strategy used by ungulates such as moose (Bowyer et al. 1999), caribou (Rangifer tarandus; Barten et al. 2001), elk (Creel et al. 2005), roe deer (Bongi et al. 2008), and mule deer (Long et al. 2009). In the hiding phase, any detection of the neonate by predators drastically impedes visual detection of predators by females in an already risky environment. Therefore, understory cover for calves may not be a driving factor for selection of birthing sites in our case study. Future studies could further explore this relationship between calf hiding cover and adult elk resource selection.

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**Literature Cited**


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