Sheep Spatial Grazing Strategies at the Arid Patagonian Monte, Argentina

Mónica B. Bertiller and Jorge O. Ares

Authors are Scientific Researchers, Centro Nacional Patagónico-CONICET, and Professors, Universidad Nacional de la Patagonia–UNPSJB, 9120 Puerto Madryn, Chubut, Argentina.

Abstract

We asked what vegetation traits influence sheep in selecting foraging paths on the range. We obtained 40 000 records of positions of six ewes (Ovis aries) collared with Global Positioning System receivers during several seasons in a paddock of 1 250 ha at the Patagonian Monte shrublands, Argentina. We classified the vegetation through ground-truth floristic analyses and remotely sensed imagery, and overlaid the ewes’ positions onto a digital map of vegetation units. For each vegetation unit, we assessed the cover of main life forms and preferred plants, the visibility range at ewe’s head height, and several structural/chemical traits of dominant shrubs (leaf mass/area, lignin–phenolics–nitrogen concentration in leaves, presence of thorny stems and spiny leaves). Ewes followed diverse paths across the paddock but always selected among a limited number of vegetation units. Selected vegetation units were those with structural traits allowing wide ewes’ visibility ranges and low structural antiherbivore defenses, irrespective of their local abundance, relative cover of preferred plants, or distance to the watering point. Within preferred vegetation units, ewes further selected those with high cover of preferred plants and/or reduced structural/chemical antiherbivore defenses. We concluded that sheep selectivity at our study paddock resulted from compromises among different structural/visual cues related to visual impairment, antiherbivore physical/chemical defenses, and the offer of preferred plants. In a hierarchy of decisions, the abundance of preferred plants was not a sufficient condition for a high selection of vegetation units. Monitoring animal movements within shrubby paddocks of the Patagonian Monte with high visual impairment can supply criteria to assess the relevance of nonnutritional environmental traits on grazers’ decisions. This information is valuable in identifying and predicting spots of potential land degradation, and planning the distribution of flocks within paddocks in the context of sustainable management practices for shrubby rangelands.

INTRODUCTION

Identifying the rules of decision that herbivores use while foraging in different environments is of relevance to test ecological hypothesis related to plant–animal interactions at different scales (Judson 1994; WallisDeVries et al. 1999; Searle et al. 2006). It is also important in understanding landscape use patterns of livestock and planning for sustainable rangeland use.
(Launchbaugh and Howery 2005; Bailey et al. 2006). Key questions about grazing search mechanisms involve the cues used by grazers at different spatial scales to select grazing locations and preferred plants within them, continuing the use of a selected vegetation patch or moving to a neighbor one (Senft et al. 1987; Bailey et al. 1996; Wallis DeVries et al. 1999; Fortin et al. 2003; Oom et al. 2004).

Senft et al. (1987) recognized two steps in the selection of feeding patches. The “location selection” step implies how herbivores move through plant communities. The “diet selection” step refers to which plants or parts of plants herbivores select to consume. Many studies assumed that grazers encounter food patches in proportion to their abundance, forage biomass, and the distance to a previous selected neighbor patch. According to these, searching would be a simple systematic or random process (Noy-Meir 1973; Charnov 1976; Owen-Smith and Novellie 1982; Stephens and Krebs 1986; Fryxell and Doucet 1993; Fortin et al. 2002; Oom et al. 2004; Searle et al. 2006). Other studies emphasized that search tactics are cognitive processes, or decision sequences that organisms use to locate available resources such as food, mates, refuges, or high-quality habitats (Viswanathan et al. 1999; Zollner and Lima 1999). The location of the watering point or places with drinkable free water has also been repeatedly indicated as major factors influencing the distribution and spatial foraging strategies of free-ranging herbivores (Graetz and Ludwig 1978; Jeltsch et al. 1997; Adler and Hall 2005). There is also evidence that nonnutritional factors such as habitat intricacy and visibility could influence foraging behavior of herbivores at large spatial scales (Fortin 2002; Fortin et al. 2003). Visual scanning in free-grazing large herbivores seems to be relevant for orientation, search, vigilance, and asserting the size of the surrounding kin group. Ungulates live in open habitat form groups, and individuals decrease vigilance and increase bite rate with increasing group size (Jarman 1974; Berger 1978; Risenhoover and Bailey 1983; Penning et al. 1993; Molvar and Bowyer 1994; Barta et al. 2004). Visual contact might be also relevant in maintaining the stability of unherded sheep groups when crossing their trajectories without mixing (Brown 1970).

In arid shrublands, large herbivores must find their preferred food patches while negotiating plant structures of heights nearly at or above their horizontal visual range and with strong chemical and physical defenses (Lauenroth 1998; Adler et al. 2005). How the intricacy of shrubland vegetation affects horizontal visibility of herbivores and their grazing decisions and how antitherbivore defenses influence local grazing choices are issues scarcely explored in arid shrubby ecosystems of the world grazed by free-ranging large herbivores. Monitoring the spatial location of grazing animals in relation to vegetation traits could contribute to identifying rules related to animal decisions about the selection of feeding areas and the permanence on them in shrublands (Skarpe et al. 2007). This information is relevant to plan the distribution of herds within paddocks, avoiding spots of land degradation in a context of adaptive management planning (Ganskopp and Bohnert 2006; Morghen et al. 2006).

We hypothesized that sheep selectivity at shrubby rangelands of the Patagonian Monte results from compromises among different currencies involving structural/visual cues related to visual impairment, antitherbivore physical/chemical defenses, and the offer of preferred plants. We predicted that 1) floristic–physiognomic vegetation units with different visibility range, chemical/structural antitherbivore defenses, and cover of preferred plants can be identified within large paddocks and at paddock subareas differing in their distance to the watering point; 2) sheep could perceive vegetation heterogeneity within the paddock by differentially selecting vegetation units with high visibility range (structural/visual cues); and 3) within vegetation units with high visibility, sheep would further select those with high (although eventually not maximum) forage quality and low antitherbivore defenses.

**METHODS**

**Study Site**

The study site is at an extended flat area dominated by the xeromorphic tall shrub community of jarilla (*Larrea divaricata* Cav.) and coirones (*Stipa* spp.), characteristic of the Patagonian Monte (Ares et al. 1990; León et al. 1998). Perennial plants (mostly shrubs) contribute to more than 99% of the total biomass and form patches covering from 40% to 60% of the soil. The area is arranged in large paddocks (1 000 to 2 500 ha) and has been grazed by unherded sheep since the beginning of the 20th century. Annual grasses, forbs, and perennial grasses are the main preferred plant life forms of sheep (Baldi et al. 2004). Wells to supply water to the foraging animals are usually located at a single corner of the paddocks. Extended ppiospheres (more than 2,500 m) are generated around watering points where structural characteristics of the plant communities are modified by frequent visits of the grazing animals (Ares et al. 2003; Bisigato et al. 2005). Within the area, we selected a triangular paddock of about 1250 ha at Smit’s Ranch (lat 42°38’S, long 65°23’W) with a single watering point at its northeastern corner. The paddock was submitted to the usual stocking rate for the area (approximately 0.10 sheep · ha⁻¹).

**Characterization of the Heterogeneity of Vegetation Within the Paddock (Prediction 1)**

**Vegetation Units and Patchiness.** We identified and described floristic–physiognomic vegetation units (VUs) within the paddock through the combination of ground-truth floristic sampling and analyses of remote imagery using the program IDRISI (v14.02; Clark Labs, Worcester, MA). We performed a preliminary identification of VUs within the paddock through an unsupervised classification procedure performed on all eight bands of a Landsat-7 image of the area obtained in September 2002, geo-referenced with conspicuous ground-truth points. The classification resulted from an iterative cluster analysis using a multivariate histogram peak technique (Richards 1986). We controlled the mapped VU boundaries in the field through ground checks (October–November 2005) and characterized the floristic composition of each VU by registering the species plant cover at 25 randomly selected sampling sites of about 10,000 m² each (three to six sites per VU, depending on VU areas). Plant cover was visually estimated using cover categories with increments of 1% (Bisigato and Bertiller
Ewes' Barkley. Small shrubs were eight directions, at Lorentz, or corresponds to that of an optical telemeter where vegetation 5 Operational definition of ewes' visibility range (EVR). The focal plane c p, j (Cav.) Heimerl in Engler and Prantl., k ~ T (Speg.) Cabrera. Nomenclature follows k 3 t s, C. avellanedae head height. k Gill. ex Mog. u Boungain-Chuquiraga hystrix j is a shrub-dominant species is a sampled site of VU Rangeland Ecology & Management # Schinus johnstonii Dom., is the total number of dominant shrub is the number of sampled sites of VU L.). j [ We identified and characterized structural and chemical n u is a weighed mean value of a shrub-leaf trait Lycium Miers. ex Bertero, three sublocations 48 hours and analyzed for nitrogen (N) concentration by mass per area (LMA) by the procedure described in Bertiller et al. 2005. In five leaves per sampled individual, we assessed the leaf subareas with respect to the watering point in November nine randomly selected individuals at near, mid, and far distance to the watering point: near (0 to 2 500 m), mid (2 500 to 4 500 m), and far (> 4 500 m). We also estimated the area of all patches of each VU at each subarea (IDRISI v14.02; Clark Labs) and calculated their nominal radius (radius of a circle of the same area as the patch). We further calculated the percentage of patches with nominal radius lower than the maximum mean visibility range (see below under Ewes' Visibility Range (EVR) at VUs).

Structural and Chemical Characterization of the Shrubby Cover of VUs. We identified and characterized structural and chemical plant/leaf traits of the dominant shrubby components of VUs related to potential herbivore preference or antiherbivore defenses. Selected (cover) dominant tall shrubs were Bougainvillea spinosa (Cav.) Heimell in Engler and Prantl., Lycium chilense Miers. ex Bertero, Chuquiraga bystrix Dom., L. divaricata, and Schinus johnstonii Barkley. Small shrubs were Atriplex lampa Gill. ex Mog., C. avellanedae Lorentz, Nassauvia fuegiana (Speg.) Cabrera. Nomenclature follows Correa (1971–1999). At all VUs, the sum of the cover of these species represented more than 85% of the total woody cover. We registered at each species the presence/absence of thorny stems and/or spiny leaves and collected fully expanded young to middle-aged green leaves (Bertiller et al. 2006) from three to nine randomly selected individuals at near, mid, and far subareas with respect to the watering point in November 2005. In five leaves per sampled individual, we assessed the leaf mass per area (LMA) by the procedure described in Bertiller et al. (2006). Other leaves of each plant were dried at 60°C for 48 hours and analyzed for nitrogen (N) concentration by semimicro Kjeldahl (Schlesinger and Hasey 1981). Lignin was quantified by the Van Soest (1963) procedure and the concentration of total phenolic compounds by the Folin–Ciocalteu method using 50% methanol as extracting solution and tannic acid as standard (Waterman and Mole 1994).

We further calculated for each sampling site (three to six sites per VU) the relative contribution of each of the eight dominant shrub species to their total plant cover, and we used those values as weights to calculate the VU mean value of each shrub–leaf trait (LMA, lignin, N, and phenolics) as:

\[ T_{tj} = \left( \sum_{j=1}^{p} \sum_{k=1}^{s} \omega_{i,j,k} \right) / p \]  

Where \( T \) is a weighed mean value of a shrub-leaf trait \( t \) at VU, \( j \) is a sampled site of VU, \( k \) is a shrub-dominant species at site \( j \), \( 0 < \omega_{i,j,k} \leq 1 \) is the cover-based weight of species \( k \) at the sampled site, \( s \) is the total number of dominant shrub species at site \( j \), and \( p \) is the number of sampled sites of VU, \( j \). We also calculated the relative cover contribution of species with spiny leaves or thorny stems to the total woody plant cover at each sampled site and their mean cover values at each VU.

Ewes' Visibility Range (EVR) at VUs

We developed an EVR algorithm to assess the relative intricacy of VUs and the potential ability of sheep to acquire visual information across them. To this aim, we randomly selected four sampling sites of each VU across the paddock. At each site, we selected three sublocations and we obtained EVR estimations along eight horizontal sight-views equally spaced at 45° azimuth intervals at ewes’ head height (0.7 m) at each sublocation. Views were obtained with a 200-mm (focal ratio: 3.5) Canon Fb optical telemeter (Canon Inc., Tokyo, Japan) and EVR was defined as the distance (m) from the observer at which the focal plane showed 50% visual obstruction by vegetation (Boyd and Svejcar 2005). For each VU, we further obtained the average EVR (four sites × three sublocations × eight directions, \( n = 96 \); Fig. 1).

Figure 1. Operational definition of ewes' visibility range (EVR). The focal plane corresponds to that of an optical telemeter where vegetation obstructs 50% of the sight field along a horizontal view at a standing ewe’s head height.
**Ewes’ Path Data and Selectivity upon VUs (Predictions 2 and 3)**

We obtained about 40,000 records of positions (one each minute) of six ewes (weight range 35–39 kg) in April 2005 (ewes 1–2), September 2005 (ewes 3–5), and January 2006 (ewe 6), harnessed with standard Global Positioning System (GPS) receivers (e-Trex; Garmin, Olathe, KS) on their forebacks, while foraging at the selected paddock. The spatial accuracy of the GPS receivers was independently tested in simulated sheep trails and was found to be higher than the spatial resolution of the Landsat-7 image used to describe the field structure (data not shown). The two-strap harnesses also carried a solid-state memory and a programmable microcontroller (EEPROM-PIC16F84; Microchips Technology, Inc., Chandler, AR). The total weight of the harnessed system was 1.2 kg, within the usual load range of telemetry equipment in studies of similar free-ranging mammals (Rutter et al. 1997; Hulbert et al. 1998). After being harnessed, the animals were first released in a small handling paddock among 125 other noninstrumented ewes, and their behavior was observed during a period (30 minutes) to check for signs of discomfort or nonacceptance of harnesses. We did not observe any of these, and all animals were released to the paddock. Recording of ewes’ positions continued until battery charge was exhausted (usually 5–8 days). The collars were then recovered and the logged positions downloaded for analysis.

We converted the position estimates to digital vector files into an application for image handling (IDRISI v 14.02; Clark Labs) and overlaid them onto the image files of the classified Landsat-7 image of the area in order to cross-tabulate the frequency of visits $(f_{p_i})$ at each VU. The position data were further used to compute movement velocities $(m \cdot \text{min}^{-1})$ between successive 1-minute records and velocity vector files were similarly overlaid onto the image files of the area for cross-tabulation analysis.

The selectivity $S_i$ exerted by sheep on a given VU$_i$ was defined as the ratio:

$$S_i = \frac{fp_i}{pa_i}$$

where $fp_i$ is the frequency of positions of sheep on VU$_i$, relative to all recorded positions and $pa_i$ is the number of image pixels classified as VU$_i$ with respect to all image pixels of the paddock. Selectivity data of each VU were discriminated by subareas according to distances to the watering point (near, mid, far) and into two speeds: 0.5–3 m · min$^{-1}$ (slow movement, grazing predominant) and > 3 m · min$^{-1}$ (fast movement, exploration predominant). We discarded selectivity values of periods of sustained immobility (0 m · min$^{-1}$) during more than 10 minutes, frequent at early afternoon and late night, which we assumed did not correspond to feeding activities.

**Statistical Analyses**

We used two-way analysis of variance (ANOVA) to analyze the significance of the differences in cover categories (total, tall shrubs, small shrubs, perennial grasses, perennial forbs, and annuals) among VUs at three distances (near, mid, and far) to the watering point, and eventual interactions between VU and distance. The assumptions of ANOVA were previously tested and data on tall shrubs and perennial grass cover were logarithmic-transformed to meet them. Multiple comparisons of cover categories among VUs were performed by LSD (least significant difference) test. Differences in structural and leaf chemical traits among shrub species and VUs were analyzed by one-way ANOVA and Kruskal–Wallis test, respectively. We used two-way ANOVA to analyze the significance of the differences in selectivity ($S$) by VUs (fixed factor) and ewe (random factor). Three-way ANOVA was used to assess the significance of the differences in (square-root transformed) $S$ by VUs, distance to the watering point, and speed (fixed factors). Interactions between factors were also tested. Pearson’s correlation was used to analyze the relationship between structural and chemical shrub traits, and stepwise regression and Principal Component Analysis (PCA) were used to analyze the relationship between $S$ (all ewes together) of VUs and mean VU shrub–leaf traits, life form cover, and EVR. Principal component analysis was performed with the correlation matrix of the mean values of shrub–leaf traits, life form cover, EVR, and $S$ of VUs. For ordination of VUs according to these traits, we further calculated the loading coefficients of each VU with respect to the two first principal components. All statistical analyses were performed with the package SPSS 7.5 (Norusis 1997).

**RESULTS**

**Characterization of the Heterogeneity of Vegetation Within the Paddock (Prediction 1)**

**Vegetation Units and Patchiness.** We identified six floristic–physiognomic vegetation units, which were patchily distributed across the paddock (Fig. 2A). VU6 was the least frequent and VU5 the most frequent on an area basis across the whole paddock (Fig. 2B). The least frequent VU6 was also the unit with the highest percentage of small patches (nominal radius lower than 20 m) whereas the most frequent VU5 was mostly arranged in large patches (Figs. 2A and 2B; Table 1). VU5 and VU2 were highly frequent near the watering point. VU5 and VU3 dominated at mid distances to it, whereas VU2, VU3, VU4, and VU5 were all frequent at the far subarea. At all subareas VU1 and VU6 were rare. VU1 was the unit with the highest proportion of small patches at the subarea near to the watering point.

The VUs differed significantly in the total cover ($F_{5,25} = 4.52$, $P = 0.013$), and tall shrub ($F_{5,25} = 7.19$; $P = 0.002$), small shrub ($F_{5,25} = 10.37$, $P < 0.001$), perennial grass ($F_{5,25} = 9.86$, $P < 0.001$), and preferred plant ($F_{5,25} = 12.48$; $P < 0.001$) covers (Fig. 3). The total cover was high at VU1, coinciding with the highest tall shrub, perennial grass, and total preferred plant cover, and the lowest small shrub cover. In contrast, the total cover was low at VU3 with minimum tall shrub, perennial grass, and total preferred plant cover, and the highest small shrub cover. The total, tall shrub, small shrub, perennial grass, and total preferred plant cover did not significantly ($P > 0.05$) differ among subareas at varying distance to the watering point (near, mid, far). We did not find significant interactions between the cover of life forms at VUs and the distance to the watering point.
The small shrub *C. avellanedae* and the tall shrub *C. hystrix* displayed high LMA values and low N concentration in green leaves (Table 2). We found low LMA and lignin concentration and high concentration of N in green leaves of the deciduous tall shrubs *L. chilense* and *B. spinosa*. The tall shrub *S. johnstonii* and the small shrub *C. avellanedae* had high lignin concentrations in green leaves. *L. divaricata* and *S. johnstonii* had high concentration of phenolics; low values of this trait were in *L. chilense*, *C. hystrix* (deciduous and evergreen tall shrubs, respectively), and the small shrub *A. lampa*. Thorny stems were frequent among tall shrubs, whereas leaves with spines were frequent among small shrubs. High LMA was related to low N concentration, high lignin concentration in green leaves, leaves with spines, and the absence of thorny stems (Table 3).

Table 1. Number of patches and percentage of patches with nominal radius (r) lower than the maximum visibility range (ca. 20 m) per vegetation unit (VU) at the total paddock and at the three subareas (near, mid, and far) from the watering point.

<table>
<thead>
<tr>
<th>VU</th>
<th>No. of patches</th>
<th>Percentage of patches with r &lt; 20 m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Near</td>
</tr>
<tr>
<td>1</td>
<td>180</td>
<td>9</td>
</tr>
<tr>
<td>2</td>
<td>148</td>
<td>40</td>
</tr>
<tr>
<td>3</td>
<td>301</td>
<td>99</td>
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<tr>
<td>4</td>
<td>154</td>
<td>27</td>
</tr>
<tr>
<td>5</td>
<td>218</td>
<td>28</td>
</tr>
<tr>
<td>6</td>
<td>55</td>
<td>7</td>
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</tbody>
</table>

Figure 2. A, Graphical representation of the study paddock showing the spatial distribution of the six defined vegetation units (VU1 to VU6). The study paddock has a single watering point located at its northeastern corner. Curved lines delimit three subareas at varying distances to the watering point (near: 0–2,500 m, mid: 2,500–4,500 m, far: > 4,500 m). B, Polar plots showing the relative frequencies of the vegetation units (VU1 to VU6) at the total paddock and at the three delimited subareas according to their distance to the watering point (near, mid, far as in A).

Figure 3. Polar plots showing the total, tall shrub, small shrub, perennial grass, and preferred plant cover (%) at vegetation units (VU1 to VU6). Different lowercase letters indicate significant differences in cover categories among vegetation units.

Structural and Chemical Characterization of Dominant Shrubs at VUs. The small shrub *C. avellanedae* and the tall shrub *C. hystrix* displayed high LMA values and low N concentration in green leaves (Table 2). We found low LMA and lignin concentration and high concentration of N in green leaves of the deciduous tall shrubs *L. chilense* and *B. spinosa*. The tall shrub *S. johnstonii* and the small shrub *C. avellanedae* had high lignin concentrations in green leaves. *L. divaricata* and *S. johnstonii* had high concentration of phenolics; low values of this trait were in *L. chilense*, *C. hystrix* (deciduous and evergreen tall shrubs, respectively), and the small shrub *A. lampa*. Thorny stems were frequent among tall shrubs, whereas leaves with spines were frequent among small shrubs. High LMA was related to low N concentration, high lignin concentration in green leaves, leaves with spines, and the absence of thorny stems (Table 3).

Mean shrub–leaf traits differed significantly among VUs (LMA: $F_{5,25} = 5.16$, $P = 0.004$; lignin: $F_{5,25} = 5.04$, $P = 0.004$; N: $F_{5,25} = 4.99$, $P = 0.004$; phenolics: $F_{5,25} = 3.81$, $P = 0.015$; thorny stems: $F_{5,25} = 9.64$, $P < 0.001$; leaves with spines $F_{5,25} = 6.17$, $P = 0.001$; Fig. 4). VU3 showed the largest mean...
LMA and lignin and phenolics concentrations in green leaves of shrubs. This VU also showed the highest proportion of woody cover with spiny leaves and the lowest with thorny stems. The other VUs did not differ in the mean values of LMA and lignin concentration in green leaves of shrubs but in their values of N and phenolics. VU1 and VU2 showed high mean N concentration in green leaves of shrubs, and VU1 high concentration of phenolics and the highest proportion of woody cover with thorny stems.

**Ewes' Visibility Range (EVR) at VUs**

The mean EVR was maximum (over 20 m) at VU3 with high values at VU5 and VU6 and low values at VU2 (9.6 m), VU1, and VU4 (Fig. 5). The cover of small shrubs was positively correlated to EVR ($r = 0.765, F_{2,25} = 16.92, P = 0.001$) and the cover of tall shrubs was negatively correlated to EVR ($r = -0.772, F_{2,25} = 17.73, P = 0.001$).

**Ewes' Path Data and Selectivity Upon VUs (Predictions 2 and 3)**

Ewes displayed a variety of paths differing in total length, localization within the paddock, and spatial spread of the visited locations (Fig. 6A). The paths of ewes 1–2 were mostly located at the northwest corner of the paddock, ewe 3 remained close to the watering point at the northeast area, and ewes 4–6 explored wide (but different) regions of the available paddock. There were no differences in $S$ among ewes ($F_{5,96} = 0.86; P = 0.522$) nor significant interactions $VU \times ewe$ ($F_{2,96} = 1.33; P = 0.186$), implying a lack of evidence of seasonal effects. The pattern of selectivity respect to VUs changed at varying distance to the watering point (Fig. 6B; significant interaction $VU \times$ distance $F_{10,192} = 3.77; P < 0.001$) but it did not depend on the speed adopted by ewes to explore the VUs ($F_{1,192} = 1.69; P = 0.194$). Near the watering point, $S$ did not significantly differ among VUs, whereas at mid and far distances ewes selected with higher frequency VU3, VU5, and VU6. At mid-distance to the watering point, VU5 and VU6 were similarly selected, whereas at far subareas VU5 was the most selected. We did not find significant ($P > 0.05$) interactions between speed and other factors (VUs or distance to the watering point).

The PCA (Fig. 7) clustered $S$ at mid and far distances to the watering point along with VU3 and VU6, and dominant structural characteristics of these, such as high EVR. This showed preferential ewes' selectivity for VU3 and VU6 with high EVR at those subareas. The proximity of VU5 to S at all subareas in the PCA analysis indicated that ewes' selectivity for this VU is independent of the distance to the watering point. The PCA clustered $S$ near the watering point along with VU1 and several of its dominant structural characteristics, such as high cover of perennial grasses and preferred plants. This pointed out increased ewes' selectivity for VU1 near to the watering point. VU2 and VU4 were clustered at low scores of PCA 2, unrelated to the previous groups indicating low ewes' selectivity for these VUs (Fig. 7).

The results of the multiple regression analysis also showed that traits explaining the variation in $S$ (all ewes together) varied depending on the distance to the watering point (Table 4). Near the watering point, the cover of perennial grasses was the single variable significantly explaining the highest fraction of the total variance in $S$ (11.1%). At mid and far distances to the watering point, both EVR (positively) and

**Table 2.** Mean ($\pm$ SE) physical and leaf chemical treats of the dominant shrubs at the study area. Different lowercase letters indicate significant differences in leaf traits among species. D indicates deciduous; E, evergreen; and LMA, leaf mass per area.

<table>
<thead>
<tr>
<th>Shrub life form/species</th>
<th>LMA (g·m$^{-2}$)</th>
<th>N (%)</th>
<th>Lignin (%)</th>
<th>Phenolics (%)</th>
<th>Thorny stems</th>
<th>Leaves with spines</th>
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<tbody>
<tr>
<td>Tall shrubs</td>
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<tr>
<td><em>Bougainvillaea spinosa</em> (D)</td>
<td>107.2 ± 7.6 b</td>
<td>2.87 ± 0.08 d</td>
<td>5.78 ± 0.42 ab</td>
<td>2.81 ± 0.12 b</td>
<td>X</td>
<td>—</td>
</tr>
<tr>
<td><em>Lycium chilense</em> (D)</td>
<td>67.3 ± 6.0 a</td>
<td>2.69 ± 0.09 d</td>
<td>5.27 ± 0.45 a</td>
<td>1.68 ± 0.10 a</td>
<td>X</td>
<td>—</td>
</tr>
<tr>
<td><em>Choquiraaga hysterix</em> (E)</td>
<td>318.1 ± 11.4 f</td>
<td>1.25 ± 0.04 a</td>
<td>8.38 ± 0.73 b</td>
<td>1.57 ± 0.18 a</td>
<td>—</td>
<td>X</td>
</tr>
<tr>
<td><em>Larrea divaricata</em> (E)</td>
<td>172.8 ± 6.0 d</td>
<td>1.97 ± 0.04 c</td>
<td>4.98 ± 0.32 a</td>
<td>15.15 ± 0.50 d</td>
<td>—</td>
<td>—</td>
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<tr>
<td><em>Schinus johnstonii</em> (E)</td>
<td>231.8 ± 13.5 e</td>
<td>1.77 ± 0.05 bc</td>
<td>11.24 ± 1.54 d</td>
<td>16.56 ± 0.14 d</td>
<td>X</td>
<td>—</td>
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<tr>
<td>Small shrubs</td>
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<tr>
<td><em>Atriplex lampa</em> (E)</td>
<td>160.6 ± 10.8 cd</td>
<td>2.11 ± 0.06 c</td>
<td>8.70 ± 0.22 c</td>
<td>1.29 ± 0.08 a</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><em>Choquiraaga avellanae</em> (E)</td>
<td>454.3 ± 14.7 g</td>
<td>1.29 ± 0.03 a</td>
<td>10.80 ± 0.93 cd</td>
<td>3.28 ± 0.11 b</td>
<td>—</td>
<td>X</td>
</tr>
<tr>
<td><em>Nassauvia fuegiana</em> (E)</td>
<td>133.8 ± 19.8 bc</td>
<td>1.69 ± 0.05 b</td>
<td>6.99 ± 0.24 b</td>
<td>5.27 ± 0.19 c</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

**Table 3.** Pearson's correlations among leaf shrub traits (LMA indicates leaf mass per area; traits of green leaves include concentration of N, lignin, and phenolics), thorny stems, and leaves with spines. Correlation significance level: * = $P \leq 0.05$, ** = $P \leq 0.01$; $n = 66$.
mean lignin concentration in green leaves (negatively), along with the cover of tall shrubs (negatively) or small shrubs (positively), significantly explained 52.6% and 40.1%, respectively, of the total variance in ewes' selectivity of VUs.

**DISCUSSION**

In accordance with our first prediction, the vegetation was spatially heterogeneous in terms of floristic–physiognomic vegetation units, visibility range, structural/chemical antiherbivore defenses, and cover of preferred plants within the grazing paddock and at subareas differing in their distance to the watering point. In this patchy environment, individual ewes used highly diverse paths, but irrespective of these, they all selected among a limited number of vegetation units. Primary selection was in terms of nonnutritional, structural/visual cues. This is consistent with our second prediction and with previous studies indicating the importance of nonnutritional traits in grazers' decisions (Fortin et al. 2003). Vegetation units with low visual impairment and low levels of structural (thorny stems) plant antiherbivore defenses were primarily selected, irrespective of their relative offer of preferred plant cover or local abundance. The reasons for ewes preferring open vegetation units allowing wide visibility ranges are not clear at this point. It might be speculated that visibility would minimize the time and costs of traveling between feeding stations, which could be substantially high in visually-oriented grazers in shrubby rangelands (Gross et al. 1995). Avoiding visual impairment could also be related to maintaining adequate visual contact among congeners, orientation, search image forming, and predator risk avoidance (Jarman 1974; Berger 1978; Risenhoover and Bailey 1985; Penning et al. 1993; Molvar and Bowyer 1994; Dukas and Kamil 2001). These add further dimensions to views stressing the relevance of the biomass of available forage and the distance to a potentially preferred new neighbor patch as driving variables in grazers' spatial decisions (Noy-Meir 1973; Charnov 1976; Oom et al. 2004; Searle et al. 2006).

We also found that the nature of the structural cues related to visual impairment in the study paddock varied with the distance to the watering point. At subareas at mid and far distances to it, ewes selected vegetation units with low intricacy (high visibility range) resulting from high cover of small shrubs/low cover of tall shrubs (VU3, VU5, and VU6). Near to the watering point, ewes also selected the low visibility unit VU1. At this area, this unit occurred mostly in small patches (diameters < 20 m) that allow interpatch visibility in ranges comparable to those characteristic of intrapatch visibility at the less intricate vegetation units selected at mid and far distances to the watering point. Thus, ewes would profit from a visually open interpatch scenario, while accessing preferred species at the borders of VU1 patches.

In relation to our third prediction, sheep selected vegetation units with larger grazing rewards among those units with low visual impairment. This is consistent with other studies, indicating that predominant grazing options at small spatial scales are related to rewards and costs derived from grazing.
efficiency on preferred plants of high forage quality (Oom et al. 2004; Searle et al. 2006). Also, we identified different grazing-related rewards in relation to the distance to the watering point. High cover of small shrubs with low lignin represented grazing-related rewards at mid and far distances to the watering point, along with low LMA in leaves and reduced physical antitherbivore defenses (absent or rare thorny stems) at VU5 or VU6. Near the watering point, grazing-related rewards included high cover of perennial grasses and preferred plants, and shrub leaves with high N concentration such as in VU1. These results indicate that vegetation selection oriented to maximize food quantity and quality, as predicted by classical foraging models (Senft et al. 1987) was in our study a necessary, but not overriding condition with respect to other nonnutritional foraging cues.

CONCLUSIONS AND MANAGEMENT IMPLICATIONS

Our results indicated that in the study paddock, sheep selectivity occurred in successive decision stages. These were related to visual impairment, antitherbivore physical/chemical defenses, and the offer of preferred plants. In a hierarchy of decisions, the abundance of preferred plants was not a sufficient condition for a high selection of vegetation units. Monitoring animal movements within shrubby paddocks of the Patagonian Monte with high visual impairment might supply criteria to assess the relevance of nonnutritional environmental traits on grazers’ decisions. This information is valuable in identifying and predicting spots of potential land degradation, and planning the distribution of flocks within paddocks in the

Figure 6. A, Graphical representations of the study paddock showing the paths of the six ewes. Ewes 1–2, April 2005; ewes 3–5, September 2005; ewe 6, January 2006. B, Polar graphs showing ewes’ selectivity (S) upon vegetation units (VU1 to VU6) at the three distances to the watering point (WP): near (0–2 500 m), mid (2 500–4 500 m), far (> 4 500 m). See also Fig. 2. Different lowercase letters indicate significant differences in traits among vegetation units.
context of sustainable management practices for shrubby rangelands.

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LITERATURE CITED


