Plant species richness in relation to pasture position, management, and scale

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1. Introduction

A current hypothesis is that pasture ecosystems depend on species diversity. The loss of biodiversity on local, regional, and global scales has prompted scientists to question whether these losses impair ecosystem functions such as biomass production (Hector et al., 1999; Tilman et al., 2001), litter decomposition (Hector et al., 2000; Knops et al., 2001), nutrient cycling (Hooper, 1998; Hooper and Vitousek, 1999), and resistance to invasion by weeds (Crawley et al., 1999; Dukes, 2002). Sanderson et al. (2004) examined implications of this research for temperate forage and grazing land ecosystems and observed that benefits of increased species diversity were extrapolated to forage and grazing land ecosystems with little supporting data and that most information on the potential benefits of plant diversity arose from studies of synthesized grasslands that had not included domestic livestock.

Topographic and soil variation, intrinsic properties of many grazing lands, suggests that a diversity of species may be necessary to optimize production and quality of forage spatially and temporally. Harmoney et al. (2001) examined whether variation associated with landscape position and grazing management affected forage production and legume diversity in temperate pastures. Results suggested that a diverse mixture of legumes sowed on backslope positions in permanent, low-input pastures could enhance the production and quality of forage as a result of positive effects of legumes on soil-
plant nitrogen balances. In a follow up of this research, landscape position and grazing management effects on overall species diversity and functional composition of these pastures were examined (Guretzky et al., 2005). Grazing reduced cover of the dominant, perennial vegetative grasses, smooth brome (Bromus inermis Leyss.) and reed canarygrass (Phalaris arundinacea L.) and encouraged greater diversity and cover of subdominant grasses, legumes, and non-N₂-fixing forbs, effects most evident on backslopes.

The objective of this study was to examine how landscape position, grazing management, and scale affected overall plant species and functional group species richness in temperate pastures located in the central United States. Species richness was determined at 0.18 m² and 4.5 m²-scales from data collected from four experimental pastures that each contained three landscape positions (summit, backslope, and toeslope) and three stocking systems (continuous, rotational, and nongrazed). It was hypothesized that scale would affect the ability of landscape position and grazing management to explain species richness variation and characterize species richness patterns.

2. Materials and methods

Research was initiated in 1995 at the Iowa State University Rhodes Research Farm, USA (41°52′N, 93°10′W). Six cool-season grass pastures, each of which had summit, backslope, toeslope, opposite backslope, and opposite summit landscape positions, were identified and interseeded with a mixture of alfalfa (Medicago sativa L.), biennial yellow sweetclover (Melilotus officinalis (L.) Pall), biennial white sweetclover (M. alba Medic.), birdsfoot trefoil (Lotus corniculatus L.), white clover (T. repens L.), red clover (T. pratense L.), kura clover (T. ambiguus Bieb.), cicer milkvetch (Astragalus cicer L.), berseem clover (T. al exandrinum L.), striate lespedeza (Kummerowia striata (Thunb.) Schindler), and annual white sweetclover (M. alba Medic.). Of the six pastures, four were 1.2 ha and two were 1.0 ha and three had north–south orientations and three had east–west orientations. Slopes ranged from 0% to 5% on summit and toeslope positions and 10% to 24% on backslope positions. Summit and backslope positions consisted of Downs (fine-silty, mixed, superactive, mesic Typic Hapludalf) soils, and toeslope positions consisted of Colo (fine-silty, mixed, superactive, mesic Cumulic Haplauquoll) and Ackmore (fine-silty, mixed, non-acid, mesic Aeric Fluvaquent) soils (Oelmann, 1981).

In spring 1996, each pasture was subdivided into three equal-sized paddocks that crossed in a perpendicular manner the summit, backslope, toeslope, opposite backslope, and opposite summit positions. The three paddocks were randomly assigned a stocking system: continuous, rotational, or nongrazed (control). Stocking rates were similar among the continuous and rotational systems: 9.4–10.1 animal unit months (AUM) ha⁻¹. In most years, the continuously stocked paddocks were grazed by one to two mature, non-lactating beef cows from mid-May through mid-August and during November if forage was available. Paddocks assigned rotational stocking were grazed by eight to nine mature, non-lactating beef cows across a 4 d span, three times annually (May, July, and November). The intent of the rotational system was to minimize selective grazing, remove the majority of forage within a short time period, and increase the period of rest between grazing events relative to the continuous stocking system. Patch grazing appeared to be greater within the continuous system as sward heights ranged from <5 cm in heavily utilized areas to >20 cm in under utilized areas. Sward heights appeared more uniform following rotational stocking: 10–20 cm on summits and backslopes and 20–30 cm on toeslopes. Grazing began on the same date within each paddock and replicate and continued annually through November 2002. Cows were removed from the continuously stocked paddocks when residue height for the majority of the herbage was <13 cm and from the rotationally stocked paddocks after the 4 d period of each grazing event. Nongrazed plots were mowed annually in mid-November.

Data were collected from the four 1.2 ha pastures in May and July of 2000 and 2001 to examine effects of pasture position and management on vegetation species cover. Of the four pastures, two had north–south orientations and two had east–west orientations. Each sampling period ninety to hundred 0.18 m² quadrats were randomly distributed and sampled within each stocking system (paddock) for a total of 270–300 per pasture. After two seasons and 2 years of data collection in the four pastures, a total of 4484 quadrats were sampled. Plant species occurring within each quadrant were identified and classified as perennial grasses, perennial herbs, and annuals–biennials. The perennial grasses group, consisting largely of cool-season grasses considered desirable within these pastures, was subdivided into grasses capable of spreading vegetatively and bunch-types. Perennial herbs were subdivided into legumes and non-N₂-fixing forbs. Most of the legumes still persistent since initial establishment in 1995 were perennial. Few species overall were annuals and biennials. Thus, they were combined into one group consisting of annual and biennial grasses and herbs.

Although 270–300 quadrats were sampled in each pasture during each sampling period, area differences resulted in an unequal number of quadrats sampled per landscape position. Thus, overall species richness and species richness per functional group was determined from of a random subset of 25 samples from each position (3) of each stocking method (3) and pasture replicate (4) combination for a total of 900 samples. Overall species richness and species richness per functional group at the 0.18 m²-scale was determined from the means of the 25 samples. For species richness estimates at the 4.5 m²-scale, means were not calculated, but rather, the same subset of 25 samples were com-
bined to form one sample per landscape position, stocking system, and pasture replicate combination. The limited area of toeslope positions in the pastures restricted the species richness-scale analysis to 25 samples and the scale of 4.5 m². The 4.5 m² samples were subsequently combined into 37.5 m² and 162 m²-scales to model species richness-area relationships across the pastures.

The experiment was a randomized complete block design. Landscape positions and stocking systems were arranged as a split block within each of the four pasture replicates. The experiment was analyzed using the GLM procedure within the Statistical Analysis System software (SAS Institute Inc., Cary, NC). To compare how scale affected the ability of positions and management systems to explain species richness variation, measures of effect size or the degree of association between the effects and the dependent variables were calculated as the ratio of the effect variance to the total variance: \(100 \times \left(\frac{\text{sum of squares}_{\text{effect}}}{\text{sum of squares}_{\text{total}}}\right)\) for each scale. Single degree of freedom contrasts were used to examine mean differences among stocking systems and landscape positions when main effects were significant (\(P \leq 0.05\)). Landscape position and stocking system were considered fixed and pasture replicates random. Because previous analyses showed that effects of aspect, season, and year on species composition and diversity were mostly insignificant (Harmony et al., 2001; Guretzky et al., 2005), these factors were not considered further.

3. Results

3.1. Scale effects

Scale affected the percentage of plant species richness variation explained by landscape position and grazing management. For overall, perennial, perennial grass, perennial vegetative grass, perennial N

\(_\text{2}\)

-fixing herb functional groups, landscape position significantly explained 3–26% of species richness variation at the 4.5 m²-scale compared to 40–63% at 0.18 m². Landscape position did not significantly affect species richness of annual–biennial, perennial bunchgrass, and perennial non-N

\(_\text{2}\)

-fixing herb functional groups at either scale (\(P > 0.05\)). Grazing management, in contrast, significantly explained 22–35% of species richness variation for overall, perennial, annual–biennial, perennial grass, and perennial bunchgrass functional groups at 0.18 m² compared to 30–43% of their variation at 4.5 m². For vegetative-spreading grass, perennial herb, and perennial non-N

\(_\text{2}\)

-fixing herb functional groups, grazing management significantly explained 10–17%, 12–16%, and 14–21% of their variation, respectively, across the two scales. Landscape position and grazing management interacted once, affecting overall species richness at 0.18 m² (\(P \leq 0.05\)). Across all functional groups, landscape position and grazing management interactions accounted for only 1-14% of species richness variation. Together grazing management and landscape position explained 70–81% of species richness variation at 0.18 m² compared to 34–61% at 4.5 m² for overall species, perennial, perennial grass, perennial herb, and perennial N

\(_\text{2}\)

-fixing herb functional groups.

Species richness increased with scale in a logarithmic manner. When averaged across samples combined from the four pasture replicates, three landscape positions, and three stocking systems, 39 species overall were supported at 162 m² (Figure 1). Of these species, 59% were perennials, 33% were annuals–biennials, and 8% were unidentified grasses and herbs (data not shown). Among perennials, the pastures supported 8 grasses and 16 herbs. Five of the grasses were vegetative spreading and three were bunch-type species. Legumes reached a plateau of six species at 40.5 m². Forbs comprised 63% of the perennial herb functional group at 162 m².

3.2. Landscape position effects

Species richness of overall, perennial, perennial grass, perennial bunchgrass, perennial herb, and perennial N

\(_\text{2}\)

-fixing herb functional groups were greater on backslopes than on summits and toeslopes at both 0.18 m² and 4.5 m² (Table 1). Backslopes supported similar species richness of perennial vegetative grasses, perennial non-N

\(_\text{2}\)

-fixing herbs, and annuals–biennials as summits and toeslopes. Summits supported greater species richness of annual–biennial, perennial bunchgrass, and perennial non-N

\(_\text{2}\)

-fixing herb functional groups. Landslopes did not significantly affect species richness of overall species: \(y = 5.0 \ln(x) + 8.4, R^2 = 0.89\); perennials: \(y = 2.8 \ln(x) + 6.4, R^2 = 0.92\); annuals–biennials: \(y = 1.7 \ln(x) + 1.1, R^2 = 0.76\).

Figure 1. Species richness–area relationships derived from four 1.2 ha pastures at the Iowa State University Rhodes Research Farm, USA. For overall species: \(y = 5.0 \ln(x) + 8.4, R^2 = 0.89\); perennials: \(y = 2.8 \ln(x) + 6.4, R^2 = 0.92\); annuals–biennials: \(y = 1.7 \ln(x) + 1.1, R^2 = 0.76\).
Grazing management effects

Continuously and rotationally stocked (grazed) pastures supported more species overall and within each functional group than nongrazed pastures at both scales. Within the continuous, rotational, and nongrazed systems, overall species richness averaged 5.4, 4.4, and 3.3 on backslopes, 3.6, 3.4, and 2.6 on summits, and 2.9, 2.6, and 2.1 on toeslopes, respectively, at 0.18 m² (position × management interaction, \( P \leq 0.05 \)). At 4.5 m², overall species richness averaged 14.8, 12.9, and 8.7 within the continuous, rotational, and nongrazed systems, respectively (Table 2, position × management interaction, \( P > 0.05 \)). Grazing increased species richness of both vegetative and bunch-type perennial grasses, \( \text{N}_2 \)-fixing and non \( \text{N}_2 \)-fixing perennial herbs, and annual–biennial grasses and forbs. Continuous and rotational stocking differed only for species richness of perennials at 0.18 m² and perennial bunchgrasses at 0.18 m² and 4.5 m² (Table 2).

4. Discussion

Overall plant species richness and plant species richness per functional group depended on pasture position, grazing management, and scale. Backslope positions supported more species than summits and toeslopes regardless of scale. Positional differences appeared related to total vegetation production and aboveground competition. Total available herbaceous biomass averaged 1816, 2178, and 2971 kg ha⁻¹ on backslopes, summits, and toeslopes, respectively (Harmon et al., 2001). Grazing likely enhanced species richness through reduced aboveground competition, greater heterogeneity with regards to light and soil nutrient availability, and enhanced seedling recruitment opportunities. Indeed, grazing reduced the percentage cover of smooth brome and reed canarygrass, the dominant grasses in these pastures, from 83% to 69% and enhanced heterogeneity with regards to sward heights through defoliation (Guretzky et al., 2005). An increase of scale enabled detection of a greater number of species per unit area but reduced the ability of pasture position and management to explain pasture species richness variation.

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### Table 1. Landscape position (summit (S), backslope (B), and toeslope (T)) effects on species richness of overall, perennial, annual–biennial, perennial grass, perennial vegetative grass, perennial bunchgrass, perennial herb, perennial \( \text{N}_2 \)-fixing herb (legume), and perennial non-\( \text{N}_2 \)-fixing herb (forb) functional groups at 0.18 and 4.5 m²-scales at Rhodes, IA, USA

<table>
<thead>
<tr>
<th>Functional groups</th>
<th>0.18 m²</th>
<th>1 d.f. contrasts, ( P ) value</th>
<th>4.5 m²</th>
<th>1 d.f. contrasts, ( P ) value</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Summit</td>
<td>Backslope</td>
<td>Toeslope</td>
<td>S vs. B</td>
</tr>
<tr>
<td>Overall</td>
<td>3.2</td>
<td>4.4</td>
<td>2.5</td>
<td>0.000</td>
</tr>
<tr>
<td>Perennials</td>
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<td>4.0</td>
<td>2.3</td>
<td>0.000</td>
</tr>
<tr>
<td>Annuals</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>ns</td>
</tr>
<tr>
<td>Grasses</td>
<td>2.3</td>
<td>2.7</td>
<td>2.0</td>
<td>0.004</td>
</tr>
<tr>
<td>Vegetative</td>
<td>2.2</td>
<td>2.3</td>
<td>1.9</td>
<td>ns</td>
</tr>
<tr>
<td>Bunch</td>
<td>0.1</td>
<td>0.3</td>
<td>0.1</td>
<td>0.010</td>
</tr>
<tr>
<td>Herbs</td>
<td>0.5</td>
<td>1.0</td>
<td>0.2</td>
<td>0.000</td>
</tr>
<tr>
<td>( \text{N}_2 )-fixing</td>
<td>0.4</td>
<td>0.9</td>
<td>0.1</td>
<td>0.000</td>
</tr>
<tr>
<td>Non-( \text{N}_2 )-fixing</td>
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<td>0.1</td>
<td>0.0</td>
<td>ns</td>
</tr>
</tbody>
</table>

Single degree of freedom contrasts indicate significant \( (P < 0.05) \) differences among positions.

### Table 2. Grazing management (continuous (C), rotational (R), and nongrazed (N) stocking) effects on species richness of overall, perennial, annual–biennial, perennial grass, perennial vegetative grass, perennial bunchgrass, perennial herb, perennial \( \text{N}_2 \)-fixing herb (legume), and perennial non-\( \text{N}_2 \)-fixing herb (forb) functional groups at 0.18 and 4.5 m²-scales at Rhodes, IA, USA

<table>
<thead>
<tr>
<th>Functional groups</th>
<th>0.18 m²</th>
<th>1 d.f. contrasts, ( P ) value</th>
<th>4.5 m²</th>
<th>1 d.f. contrasts, ( P ) value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Continuous</td>
<td>Rotational</td>
<td>Nongrazed</td>
<td>C vs. R</td>
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<tr>
<td>Overall</td>
<td>4.0</td>
<td>3.5</td>
<td>2.7</td>
<td>0.005</td>
</tr>
<tr>
<td>Perennials</td>
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<td>3.2</td>
<td>2.5</td>
<td>0.012</td>
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<tr>
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<td>0.1</td>
<td>0.0</td>
<td>ns</td>
</tr>
<tr>
<td>Perennial grasses</td>
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<td>2.4</td>
<td>2.1</td>
<td>ns</td>
</tr>
<tr>
<td>Vegetative</td>
<td>2.1</td>
<td>2.2</td>
<td>2.0</td>
<td>ns</td>
</tr>
<tr>
<td>Bunch</td>
<td>0.3</td>
<td>0.1</td>
<td>0.0</td>
<td>0.004</td>
</tr>
<tr>
<td>Perennial herbs</td>
<td>0.7</td>
<td>0.6</td>
<td>0.4</td>
<td>ns</td>
</tr>
<tr>
<td>Legumes</td>
<td>0.6</td>
<td>0.5</td>
<td>0.3</td>
<td>ns</td>
</tr>
<tr>
<td>Forbs</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>ns</td>
</tr>
</tbody>
</table>

Single degree of freedom contrasts indicate significant \( (P < 0.05) \) differences among systems.
Grazing itself largely accounted for species richness differences among stocking systems. Both continuously and rotationally stocked pastures had greater species richness than nongrazed pastures, effects that extended across scales and functional types. Grazer preference for taller, dominant vegetative grasses or greater tolerance of defoliation may have increased the presence of subdominant vegetative and bunch-type grasses, and disturbance may have created conditions favorable to recruitment of annual–biennials and perennial forbs, functional groups that tend to exhibit good gap colonization abilities (Tracy and Sanderson, 2000). Plant traits that tend to be associated with gap colonization ability include good germination, short-life cycles, early flowering, and heavy allocation to reproduction (Pywell et al., 2003). Litter accumulation, aboveground competition, and accompanied low light availability at soil surfaces have limited seeding recruitment and colonization opportunities and species richness in nongrazed situations (Hartnett et al., 1996; Tilman, 1997; Collins et al., 1998). Grasses that have the ability to tiller and spread rapidly vegetatively have a competitive advantage in nongrazed situations through decreased the risk of plant mortality and the position of buds below or close to the soil surface (Pywell et al., 2003).

Overall species richness compared favorably with values reported in other studies of temperate pastures with variability likely reflecting site, management, and scale differences. Thirty-nine species were found when samples were summed across landscape positions, stocking systems, and pasture replicates to form an area of 162 m² (Figure 1). Tracy and Sanderson (2000) reported that pastures across the northeastern United States supported, on average, 32 species per 1000 m². Non-N2-fixing, perennial, biennial, and annual forbs accounted for 90% of the pasture species in the latter study compared to 60% of the species in this study. Dodd et al. (2004) noted that hill country pasture composition in New Zealand typically comprises 15–25 species m⁻² with current composition largely reflecting effects of fertilizer and grazing management. Like temperate North American pastures, nearly all of the pasture species had European origins (Dodd et al., 2004). White et al. (2004) surmised a need for a functionally diverse plant community over the entire managed area rather than high localized species diversity. Although this experiment was not designed to test the effects of individual species, functional types, or overall species diversity, previous results suggest that enhancing legume content and diversity on sloped positions improves forage production spatially and temporally and enhances the N economy of low-input pasture systems (Harmony, 2001).

References


