Bite characteristics of wapiti (Cervus elaphus) in seasonal Bromus-Poa swards

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Abstract

We used a cubic sampling quadrat to study the 3-dimensional structure of volunteer Bromus-Poa swards, and explored the relationship of bite depth and sward height as a determinant of bite sizes of wapiti (Cervus elaphus) in the mixed-wood parklands of central Alberta, Canada. The vertical biomass distribution of the sward was pyramidal with leaves dominating the top stratum. Bite depths of yearling and adult wapiti were not significantly different but both were influenced by sward height and season. Wapiti selected bites in both vertical and horizontal dimensions. In spring, wapiti selected vertically, taking green leaves in the top layer of the sward. They selected forbs horizontally in summer and selected leaves vertically in mature autumn swards. Based on the relationships among bite depth and sward height, biomass and sward height, as well as vertical biomass distribution, we calculated expected bite sizes of wapiti on seasonal pasture. We also predicted changes of dietary protein and neutral detergent fiber with increasing bite depth. On spring swards, calculated dietary protein decreased and fiber increased as animals grazed deeper into the swards. In summer and autumn, dietary protein peaked as wapiti cropped about half of the height of the sward whereas dietary fiber was relatively constant. Wapiti adjusted their bite depth to select forage containing at least 14% protein in spring, summer, and autumn. The sacrifice of bite size in tall summer and autumn swards was compensated by diet quality.

Key Words: elk, bite depth, bite size, sward structure, diet quality, foraging ecology, foraging model

Wapiti (Cervus elaphus) were once abundant throughout western Canada, particularly in the aspen parklands and mixed wood forest. Gates and Hudson (1983) explored the foraging behavior of wapiti with a focus on the habitat selection. Hudson and Nietfeld (1985) and Hudson and Watkins (1986) studied feeding rate of wapiti in relation to forage biomass. Both studies indicated that sward structure was an important determinant of foraging behavior of wapiti.

Sward structure influences the forage behavior of herbivores (Milne et al. 1982, Barthram and Grant 1984). Domestic grazers consume more green than cured matter and more leaves than stems (Arnold 1987, Black and Kenney 1984, Dudzinski and Arnold 1973). Grazing sheep adjust their bite stratum according to sward structure (Burlison et al. 1991). Sward height in homogeneous hand-constructed swards has the most influence on bite depth (Ungar et al. 1991). Tiller length is correlated with herbage avail-

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Relative Biomass (%)  

**Fig. 1.** Seasonal sward structure and composition. Vertically, the biomass distribution in swards was a pyramidal structure. In spring, the biomass pyramid had a relatively larger base and there were more green leaves in the top layer. Transposing axes and calculating median RB and RH (Equations 1 and 2), the relationship between biomass and height can be summarized by the following equations: spring, \( RB = 4.5(\pm 1.45) \times 10^{-0.043(\pm 0.006)RH} \) \( (r^2 = 0.98) \); summer, \( RB = 5.3(\pm 1.27) \times 10^{-0.057(\pm 0.004)RH} \) \( (r^2 = 0.96) \); autumn, \( RB = 5.5(\pm 1.06) \times 10^{-0.056(\pm 0.003)RH} \) \( (r^2 = 0.98) \).

**Table 1.** Energy, Protein, NDF, ADF, and lignin in graminoids and dicots.  

<table>
<thead>
<tr>
<th>Season</th>
<th>Energy (kJ/g)</th>
<th>Protein (%)</th>
<th>NDF (%)</th>
<th>ADF (%)</th>
<th>Lignin (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spring</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leaves</td>
<td>15.8 ± 0.4</td>
<td>13 ± 0.2</td>
<td>67 ± 1</td>
<td>35 ± 1</td>
<td>8 ± 0.1</td>
</tr>
<tr>
<td>Culms</td>
<td>16.3 ± 0.3</td>
<td>9 ± 0.5</td>
<td>76 ± 1</td>
<td>42 ± 1</td>
<td>6 ± 0.1</td>
</tr>
<tr>
<td>G. Gr.</td>
<td>16.6 ± 0.2</td>
<td>14 ± 0.4</td>
<td>63 ± 3</td>
<td>35 ± 3</td>
<td>6 ± 0.4</td>
</tr>
<tr>
<td>dicots</td>
<td>17.2 ± 0.0</td>
<td>25 ± 0.4</td>
<td>30 ± 1</td>
<td>14 ± 1</td>
<td>3 ± 0.1</td>
</tr>
<tr>
<td>Summer</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G. Gr.</td>
<td>16.9 ± 0.0</td>
<td>16 ± 0.1</td>
<td>65 ± 2</td>
<td>32 ± 1</td>
<td>6 ± 0.2</td>
</tr>
<tr>
<td>dicots</td>
<td>16.6 ± 0.2</td>
<td>9 ± 0.4</td>
<td>75 ± 1</td>
<td>44 ± 1</td>
<td>7 ± 0.1</td>
</tr>
<tr>
<td>Autumn</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G. Gr.</td>
<td>16.7 ± 0.0</td>
<td>22 ± 0.1</td>
<td>33 ± 1</td>
<td>17 ± 1</td>
<td>4 ± 0.0</td>
</tr>
<tr>
<td>dicots</td>
<td>16.9 ± 0.1</td>
<td>16 ± 0.2</td>
<td>68 ± 0.9</td>
<td>34 ± 1</td>
<td>6 ± 0.1</td>
</tr>
</tbody>
</table>

1. C: cured.  
2. G: green; Gr: grass.  
3. Including heads.
Exponential regression between MRH and MRBi in each season gave a dimensionless biomass density-distribution function along the vertical axis. Bite depth was calculated as an empirical function of sward height. Bite volume is a cylinder with the bite area as the cross-section and bite depth as the height (Ungar and Noy-Meir 1988). Finally, we quantified bite size in relation to bite volume and biomass distribution:

\[ BS_{SH, SN} = BM_{SH, SN} BA \int_{BD_{SH, SN}}^{100} a e^{b H} dH \]  

Where: \( a \) and \( b \) were the exponential regression coefficients, \( BS \) was bite size (g), \( BM \) was biomass (g/cm²), \( BA \) was bite area (cm²), and \( SN \) was season. \( BM_{SH, SN} \) was biomass distribution calculated by logarithmic regression of herbage biomass on sward height in each season. \( BD_{SH, SN} \) was the relative bite depth calculated by linear regression of bite depth on sward height in each season.

In Equation (3), the driving variables were sward height and season. Sward height increased at 10-cm intervals from 10 cm to 30 cm in spring, to 50 cm for sub-adults and 40 cm for yearlings in summer, and to 40 cm in autumn. The 95% confidence limits of the bite size was calculated using 95% confidence limits of the bite depth at specific sward height.

For progressive bite depth, dietary protein and NDF were calculated according to the proportions of green, dead matter, leaf, stem and their respective protein, and NDF contents in each season.

\[ DC(k) = \sum_{i=1}^{n} \sum_{j=1}^{n} sc_{ij} p_{j} \]  

Where: \( DC(k) \) was the dietary component such as protein or NDF when bite depth reached the \( k \)-th layer of the sward. \( L \) was the number of layers cut in the season. \( sc_{ij} \) was the proportion of sward component \( j \) such as cured leaves, cured culms, green leaves, green culms, or forbs in the \( k \)-th layer. \( n \) was the number of sward components found in the layer. \( p_{j} \) was the percentage of protein or NDF in the sward component \( j \). The unidentified portion of the sward sample was pooled with cured stems.

Sod Experiment
We refined our estimates of bite characteristics using six 0.5 × 0.5-m pieces of sod removed with about 5 cm of soil layer. We measured the sward heights and placed 1 piece of sod in a pen with 2 hand-reared sub-adult wapiti. When the wapiti started to eat, we took bites from their mouths to determine bite weight and measured the corresponding bite heights. When we placed a finger on the diastema, the animal egested the bite in attempting to remove the finger with her tongue.

Statistics
Two multivariate analysis of variance (MANOVA) evaluated seasonal and botanical effects and their first order interaction on energy, protein, NDF, ADF, and lignin. A multiple comparison among the means was carried out with the Tukey-Kramer test (Sokal and Rohlf 1981). Because of the different initial sward heights in each season, the impacts of season and cohort on bite depth and their first and second order interactions were explored by 2-way ANCOVA with sward height as a covariate. Data were transformed into their square-roots to achieve homogeneity of variance. Multiple comparisons among the means of sward heights and bite depth were carried out by Fisher’s protected LSD method (Sokal and Rohlf 1981). Differences were considered significant where the probability of a type I error did not not exceed 0.05. Data were summarized as the mean ± 1 se.

Results

Sward Characteristics
Biomass in swards was pyramidal, with the greatest bulk density
at the base of the sward (Fig. 1). Spring swards had more biomass at the bottom \( p < 0.05 \). Biomass and height of layers expressed in relative scale were related exponentially \( p < 0.05 \). The slope \( b \) of the exponential regression in spring was significantly different from those in summer and autumn \( p < 0.05 \). However, summer swards were no different in vertical biomass distribution from autumn ones \( p > 0.05 \).

Protein, NDF, ADF, and lignin differed significantly among seasons \( p < 0.05 \) and forages \( p < 0.05 \) and the interactions between season and forages were significant \( p < 0.05 \). Green matter had more protein and less fiber than cured plant material \( p < 0.05 \); Table 1). Dicots had more protein and less fiber than graminoids \( p < 0.05 \); Table 1). Energy content differed significantly among forages but not among seasons \( p < 0.05 \).

Bite Characteristics

Bite depth was affected by season \( p < 0.05 \), sward height \( p < 0.05 \) and their interaction \( p < 0.05 \), Fig. 2). Wapiti grazed deeper in spring swards than in summer and autumn ones \( p < 0.05 \). Bite area was related to the age of wapiti. The bite area of sub-adults was \( 38 \pm 0.1 \) cm\(^2\) in spring, and \( 49 \pm 0.1 \) cm\(^2\) in summer and autumn of 1990. Bite area of yearlings was \( 27 \pm 0.03 \) cm\(^2\) in spring and \( 38 \pm 0.1 \) cm\(^2\) in summer and autumn of 1991.

In early spring, wapiti selected green leaves above previous year’s growth, which set a constraint on diet selection. When the paddock was first grazed, the heights of green grasses were about double that of dried grasses whereas grazed stubble height was about the same as that of dried grasses. The regressions of bite height \( \text{BH} \), green grass height \( \text{GH} \) and dried grass height were as follows: \( \text{DGH} \text{GH} = 3.3(\pm 0.29) + 1.95(\pm 0.07)\text{DRH} \quad (r^2 = 0.86) \) and \( \text{BH} = 0.081(\pm 0.14) + 1.18(\pm 0.04)\text{DRH} \quad (r^2 = 0.92) \). In summer, wapiti shifted from a solely grass diet to a mixed grass and dicot diet. Nietfeld (1983) reported that dandelions accounted for 18% of the diet of wapiti in late May and 36% in June.

Yearlings and sub-adults had different bite sizes in seasonal swards (Fig. 3). In spring, yearling and sub-adult animals had larger bites in 30-cm tall swards than in 10-cm tall swards \( p < 0.05 \), whereas in summer they took smaller bites in 30-cm tall swards than in 10-cm tall swards \( p < 0.05 \). In autumn, yearlings had smaller bites in swards \( > 20 \) cm high than in 10-cm tall swards \( p < 0.05 \), whereas sub-adults had bites of \( 0.25 \) g in 10-cm tall swards and bites of about \( 0.1 \) g across all swards taller than \( 20 \) cm \( p < 0.05 \). In autumn, the bite size of yearlings decreased as sward height increased. Bite size for sub-adults was \( 0.15 \) g in 10-cm tall swards, \( 0.12 \) g in 20-30-cm tall swards and \( 0.13 \) g in 40-cm tall swards.

In 6 experiments, 1 tame wapiti ate \( 0.28 \pm 0.06 \) g (3 bites) and another ate \( 0.18 \pm 0.03 \) g (2 bites) on sod pieces with plant heights of \( 11.2 \pm 1.2 \) cm. The predicted bite size for this sward was \( 0.25 \) g with \( 0.13 \) g and \( 0.51 \) g as the 95% confidence limits. These results were within the range of predicted bite sizes.
Bite Depth and Diet Quality

The expected dietary protein and NDF are summarized in Figure 4. In spring, as the bite depth increased from the surface to the bottom of sward, calculated dietary protein in diet decreased and NDF increased. In summer, dietary protein increased as bite depth increased to 60% of the sward height. Beyond this point, dietary protein decreased, but NDF only increased slightly. In autumn, dietary protein remain relatively constant until wapiti cropped about half of the sward height. We noted a decrease in dietary protein when wapiti cropped more than 60% of the sward. Dietary NDF increased in autumn as bite depth and slightly decreased because of short forbs in the swards in summer and autumn.

The average bite depth stratum of wapiti shifted upward in swards from spring to autumn (Fig. 4). Subadults grazed deeper in spring swards and they elevated their bite strata in summer and autumn swards. However, the actual bite strata provided higher dietary protein than other cropping heights in summer and autumn. The lower limits of bite depth were adjusted to provide about 14% dietary protein in all seasons.

Discussion

Sward Structure

Hodgson (1981) noted the importance of sward structure in foraging selectivity of domestic grazers and adopted a point quadrat to study the vertical sward profile. On Serengeti grasslands, Coughenour et al. (1984) assessed canopy structure by measuring leaf angle, biomass concentration, and tiller heights. Because of the dense tillers and leaves in our study area, leaf angle measurement was inappropriate and we developed the cubic sampling quadrat to quantify sward structure. In swards less than 5-cm tall, the cubic quadrat sampling appeared suitable. The profile we obtained was similar to the triangle shape obtained by Hodgson (1981) using a point quadrat and that by Stobbs (1973b) with a hedge trimmer mounted on an adjustable frame. Biomass and height expressed in relative scales facilitated calculation of bite size and comparisons among seasons.

Milne et al. (1982) noted the linear relationship between bite depth and sward height allows the prediction of forage intake. Bite depth of wapiti in seasonal swards was very similar to those of sheep and cattle (Milne et al. 1982, Barthram and Grant 1984). Grazers tend to bite a fairly constant proportion of total sward height, because they select tillers with greater leaf length and younger laminae. Barthram and Grant (1984) noticed that reduction of bite depth in tall swards may limit intake per bite and consequently total intake. In tall matured swards, wapiti took smaller bites than in short swards during summer and autumn.

Foraging modes

Wapiti selected forages along horizontal and vertical vectors. Bite depth represents the vertical dimension. In spring, wapiti completely relied on grasses. When green leaves dominated the sward canopy, dried grasses at the bottom of the sward constrained diet selection. Wapiti maximized bite depth by grazing green leaves above dried grasses. In summer and autumn, wapiti preferred short swards with new leaves to tall mature ones. Grass leaves appear more slowly in tall swards and take longer to complete their expansion (Parsons et al. 1991), so grazing can improve pasture quality.

Wapiti selected individual forbs in the horizontal dimension. In early summer dandelion accounted for 20% of the total pasture biomass (Nietfeld 1983) and wapiti shifted their diet from solely grasses to both dandelion and grasses. Pasture became heterogeneous as seed ripened in autumn, wapiti normally nibbled the leaves, avoided tall reproductive culms, and selected feeding stations with short grasses. The horizontal selection in autumn differed from that in summer: the former involved a selection of feeding stations whereas the latter involved selection of individual plants.

Bite size

Bite sizes of sub-adults calculated in this study were in the range of bite sizes measured by other methods with the same wapiti. Jiang and Hudson (1992) reported wapiti had average bites of 0.35 ± 0.08 to 0.41 ± 0.10 g in later spring, compared to 0.13 to 0.51 g in this study. Wilmshurst (1992) measured bite sizes of yearling wapiti as 0.05 to 0.4 g in sod experiments at the same research site.

Bite size is determined by bite dimensions and sward bulk density. Laca et al. (1992) found that bite area increased with sward height and decreased with bulk density in homogeneous swards. Whether this is true for wapiti needs to be tested, because wapiti and cattle differ in foraging mechanism. Wapiti graze by using their lips, dental pad and lower incisors to grasp, and break the forage instead of using their tongues to sweep and prehend forage. Furthermore, tiller density did not vary greatly in our study area, so we assumed that bite area was constant within seasons.

The relationship between bite size and sward height varies with pasture maturity, plant species composition, and grazing system. On homogeneous artificial pastures, animals graze deeper (Milne 1991) and take larger bites from tall swards than short swards (Burilson et al. 1991). On heterogeneous natural grassland, grazing animals take smaller bites in mature than young swards (Van Soest 1982, Stobbs 1973a). In our study, sub-adult wapiti took larger bites in tall swards in spring than in summer, since the green material in spring swards was homogeneous and highly digestible (Jiang and Hudson 1992). As swards matured, wapiti spent more time in short swards, perhaps due to larger bites and high quality of the newly generated leaves. In summer and autumn, wapiti did not crop as deep as they did in tall spring swards, but their bite depth maximized possible dietary protein. Vertical selection is a balance of the tradeoff between dietary protein and bite size. Sacrifice of bite size in summer and autumn might be compensated by diet quality.

There are limitations in our methodology. First, the method of calculating bite size may be more suitable for swards of intermediate heights. In swards less than 5-cm tall, wapiti may prehend and break the grasses at ground level and bite depth may be difficult to measure. Bite depth is only an indicator of average length of tiller removed (Laca et al. 1992). In such a case, treating the bite volume as a cylinder plus a hemisphere may be better. Another shortcoming of Equation 3 may arise when wapiti adjust the angle of mouth to select the leaf stratum in matured swards, but we did not observe this foraging posture frequently. However, the equations provide a theoretical framework although relationships between bite size and sward height may differ due to different sward structures and foraging responses of the grazer.

Our approach integrated bite depth, sward height, and biomass density to predict bite sizes and diet quality of wapiti in different seasons, depicted the foraging responses of wapiti to sward structure, and provided another avenue to study bite sizes and diet selection of grazing herbivores. As a functional response to sward structure, wapiti may select bite depth and bite size according to the distribution of food items, forage quality and sward configuration. Consequently, bite sizes of wapiti may change markedly in response to seasonal changes in sward structure and forage quality.

Literature Cited


