Supplementation and monensin effects on digesta kinetics II. Cattle grazing winter range

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Abstract

Sixteen ruminally cannulated steers grazing Northern Great Plains native winter range were assigned to 4 treatments in a 2 x 2 factorial arrangement. Main effects were protein and monensin. A soybean meal-barley pellet (P; 26% crude protein) was fed at 0 and .8 kg · head⁻¹ · d⁻¹. Steers either received no monensin (M) or M was released at 101 mg/day via a ruminal delivery device. Forage intake, ruminal fermentation, in vivo organic matter (OM) digestibility, and ruminal fluid passage and particulate digesta kinetics were measured during trials in November and January. Esophageally fistulated steers were used to collect diet samples during each trial. Dietary crude protein was greater (P<0.01) in November (8.3%) than January (4.9%). Forage OM intake was not (P>0.10) influenced by either P or M. In vivo OM digestibility was increased (P<0.05) by P (60.6 vs 57.4%) and not affected (P>0.10) by M. Particulate passage rate increased (P<0.05) when P was combined with M. Ruminal fluid flow characteristics, fluid volume and pH were not affected (P>0.10) by either P or M. Ruminal ammonia-N was increased (P<0.01) by P (2.9 vs .6 mg/lOO ml) and not affected (P>0.10) by M. Total ruminal volatile fatty acid concentrations, along with molar proportions of ruminal propionate and butyrate, were not affected (P>0.10) by P or M. Ruminal acetate was decreased (P<0.10) by P and not influenced (P>0.10) by M. We conclude that supplemental protein, through ruminal modifications, has beneficial effects on OM digestibility, and can thereby provide cattle grazing winter range with additional energy at a time when it is most crucial.

Key Words: particulate passage, fluid passage, digestibility, intake, rumen fermentation, forage, protein

Livestock grazing dormant winter range are often unable to consume sufficient quantities of forage to meet their energy and protein requirements. Supplemental protein and (or) monensin might enable cattle to meet their winter nutrient requirements. Supplemental protein has often increased forage and energy intake (Cook and Harris 1968, Karchner 1980); however, data are inconclusive. Rittenhouse et al. (1970) and Judkins et al. (1987) reported no effect on forage intake as a result of protein supplementation. Monensin can increase nitrogen digestibility, decrease dietary protein requirements, and reduce degradation of dietary protein within the rumen, thereby eliciting a protein sparing effect (Schelling 1984, Bergen and Bates 1984). Clanton et al. (1981) reported that wintering range cows fed monensin had heavier weight calves than unsupplemented cows. The objective of this study was to quantify the effects of supplemental protein, monensin, and their interaction on intake, ruminal fermentation and ruminal fluid, and particulate digesta kinetics in beef steers grazing Northern Great Plains native winter range. Singularly and in combination, supplemental protein and monensin were hypothesized to beneficially alter ruminal fermentation, fluid flow, digesta kinetics, and intake.

Materials and Methods

Sixteen crossbred ruminally cannulated steers with an average initial weight of 390 kg grazed freely on a 48.6-ha, broken upland native range site from 29 October 1987 to 22 January 1988. The stocking rate was considered to be light to moderate for this range (Holscher and Woolfolk 1953) and forage was readily available throughout the study. The range site was located on the Fort Keogh Livestock and Range Research Laboratory, Miles City, Montana. Major forage species were western wheatgrass (Pascopyrum smithii [Rydby.] Löve); blue grama (Bouteloua gracilis [H.B.K.] Lag. ex. Griffiths); needle-and-thread grass (Stipa comata Trin. and Rupr.); buffalograss (Buchloe dactyloides [Nutt.] Engelm.); and threadleaf sedge (Carex filifolia Nutt.). Browse species available included greasewood (Sarcobatus vermiculatus [Hook.] Emory.); shadscale (Artemisia confertifolia [Torr. and Frem.] S. Wats.); Gardner’s salt bush (Atriplex gardneri [Moa.]. D. Dietr.); winterfat (Ceratoides lanata [Pursh.] J.T. Howell); big sagebrush (Artemesia tridentata Nutt.). Perennial forbs were rare.

Steers were assigned randomly to 4 supplementation treatments (4 steers/treatment) in a 2 x 2 factorial arrangement with main effects of protein (P) and monensin (M). A soybean meal-barley pellet (40% soybean meal, 55% rolled barley, and 5% molasses) containing 26.3% crude protein on dry matter (DM) basis was fed at either 0 or .8 kg · head⁻¹ · d⁻¹. Monensin was either not dosed or was released at 101 mg/day via a ruminal delivery device (MRDD) as described by Parrott et al. (1986). Protein was fed to individual steers, at approximately 0800 daily because this time was considered to be least disruptive to winter grazing behavior (Adams et al. 1986). Each MRDD was weighed at the beginning and end of the study and determined to have released an average of 101 mg of monensin/day. Ruminal fermentation variables, fluid passage rates, particulate digesta kinetics, organic matter (OM) intake, and in vivo OM digestibility were measured during 2 trials conducted from: (1) 15 to 20 November, and (2) 17 to 22 January.

Seven days before starting each trial, a mass esophageal extrusa sample was collected from the study pasture from 16 mature esophageally fistulated steers, composited across steers, and labeled with Yb for use as a particulate phase marker. To estimate particulate passage rate and gastrointestinal dry matter (DM) fill, each ruminally cannulated steer received an intraruminal pulse dose (180 g DM in trial 1; 250 g DM in trial 2) of Yb-labeled forage (3.39 g Yb in trial 1; 4.44 g Yb in trial 2) via the rumen cannula at 1800 on day 1 of each trial. To estimate ruminal fluid passage rate and volume, each steer received an intraruminal dose of 1,045 mg of cobalt in a Coethylenediaminetetraacetate solution (EDTA; Uden et al. 1980) 12 h after (0600, day 2) Yb-labeled forage was dosed. Procedures for labeling esophageal extrusa with Yb and introduction of Yb labeled forage and Co-EDTA into the rumen were the same as those described by Ward et al. (1990).

Just before dosing with Co-EDTA, and again at 4, 8, 12, 16, 20,
24, and 36 h after dosing, 100-ml samples of ruminal fluid were withdrawn from the ventral sac of the rumen of each steer. The pH was determined immediately with a combination electrode, after which samples were strained through 4 layers of cheesecloth, acidified with 2 ml 25N H₂SO₄, and frozen for later analysis. Fecal grab samples were taken from each steer at 0, 12, 16, 20, 24, 28, 32, 36, 42, 48, 54, 60, 72, 84, 96, 108, and 120 h after dosing with Yb-labeled forage, dried at 45°C, and ground to pass a 1-mm screen. Adams et al. (1986) reported minimal grazing activity during non-daylight hours for cattle grazing winter native range. Hence, to facilitate ease of sample collection during trials, steers were gathered at dusk and penned in a large corral facility and released to graze at sunrise. Laboratory procedures for determining ammonia-N, volatile fatty acid (VFA) and Co concentration of ruminal fluid samples and Yb concentration of fecal samples and mathematical procedures for determining particulate passage, fluid passage, forage intake, and digestibility have been described by Ward et al. (1990). Five esophageally fistulated steers collected diet samples from the study pasture on the third day of each 6-d trial. They had grazed the study pasture 2-d before each sample collection. Esophageal collections began at about 0700 and lasted for 30 to 45 min. Between trials, the steers grazed an adjacent pasture with similar plant composition. Esophageal steers were not supplemented during the study period. Esophageal samples were dried at 45°C and ground to pass a 1-mm screen. Esophageal samples were analyzed for crude protein, DM, and ash by standard methods (AOAC 1980). Neutral detergent fiber, acid detergent fiber, and acid detergent lignin of esophageal samples were determined by the non-sequential procedures of Goering and Van Soest (1970).

Ruminal fermentation measurements, ruminal fluid volume and fluid flow rates, along with intake, particulate passage, and in vivo digestibility, were analyzed by split-plot analysis of variance with the 4 treatments arranged in a 2 × 2 factorial as the main plot and trial as the sub-plot (Gill and Hafs 1971) using the General Linear Models procedure of SAS (1985). Ruminal fermentation variables were averaged across samples taken at 0, 4, 8, 12, 16, and 20 h, and the mean was analyzed. Chemical composition of esophageal samples was analyzed as a split plot with no main plot and trial as the subplot (repeated measurements). Except where noted, significant differences are reported at the P<0.10 level.

Results and Discussion
The treatment by trial interaction was not significant for any of the variables tested in this study. Significant (P<0.01) trial effects were noted only for intake and digestibility estimates.

Nutrient Composition
Chemical composition of fistula-collected forage samples during the 2 trial periods is presented in Table 1. Dietary crude protein was greater (P<0.01) and OM was less (P<0.05) in trial 1 than trial 2, reflecting nutrient losses that occur from range forages with seasonal progression (Hogan 1984), and potential changes in diet selection. Neutral detergent fiber, acid detergent fiber, and acid detergent lignin did not vary by trial.

Intake and Passage Rate
Forage and total OM intake (Table 2) were not affected by P or

Table 2. Intake, gastrointestinal tract, fill, particulate passage and digestibility as influenced by supplemental protein and monensin in steers grazing native range.

<table>
<thead>
<tr>
<th>Item</th>
<th>Treatment*</th>
<th>EMS²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forage organic matter intake, kg</td>
<td>NR NR+M NR+P NR+P+M</td>
<td>.032</td>
</tr>
<tr>
<td>kg d⁻¹ × 100 kg body wt⁻¹</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total organic matter intake, kg</td>
<td>1.6 1.5 1.5 1.4</td>
<td></td>
</tr>
<tr>
<td>kg d⁻¹ × 100 kg body wt⁻¹</td>
<td>.032</td>
<td></td>
</tr>
<tr>
<td>Gastrointestinal tract fill, kg</td>
<td>5.1 5.7 4.4 4.2</td>
<td>.440</td>
</tr>
<tr>
<td>dry matter</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Particulate passage rate, % h⁻¹</td>
<td>2.6 2.4 2.8 2.8</td>
<td>.050</td>
</tr>
<tr>
<td>Total mean retention time, h</td>
<td>66.5 67.0 59.4 53.6</td>
<td>83.3</td>
</tr>
<tr>
<td>Forage Organic matter digestibility, %</td>
<td>57.3 57.5 60.7 60.6</td>
<td>6.88</td>
</tr>
</tbody>
</table>

*NR=native range, P=protein pellet, M=monensin.
EMS=Error mean squares.
P=protein effect significant (P<0.05).
M=Monensin interaction (P<0.05).

M. Forage and total intake (forage plus supplement) estimates were greater than those reported by Judkins et al. (1987) for steers grazing blue grama winter range in New Mexico. Protein supplementation effects on forage intake have been variable, and explanations regarding variability are not defined clearly in the literature (Cook and Harris 1968, Rittenhouse et al. 1970, Kartchner 1980). Lack of an intake response to protein supplementation in the present study may have resulted from relatively mild weather conditions. The mean minimum daily temperature was −4.9°C in trial 1 and −13.9°C in trial 2. As temperatures decline, grazing time and forage intake decrease (Adams 1987). Consequently, a greater effect of supplemental protein may be elicited during colder winter conditions when intake is expected to be lower. This concept concurs with Kartchner (1980), where supplemental protein had no effect on forage intake during mild winters and a substantial effect during harsh winters. A suppression in intake because of monensin as reported by Lemenager et al. (1978a) for cows grazing winter range contrasts results from the present study. Variation between studies may be the result of nonuniformity in range forage species (i.e., cool vs warm-season), environmental conditions, method of M delivery, and differences in animal type.

Forage OM digestibility was increased (P<0.05) by P, but not affected by M. Protein supplementation would be expected to positively influence OM digestibility because ruminal ammonia-N concentrations (Table 3) of nonsupplemented steers were well below levels suggested (Slyter et al. 1979) as optimal for microbial growth. Improved forage digestibility increases total energy available to the animal and can lead to improved performance of cattle consuming mature winter forage (Kartchner 1980, Ward and Ward 1987). Other researchers (Dinius et al. 1976, Lemenager et al. 1978a) also reported no effects of M on digestibility of winter range grass.

Gastrointestinal tract dry matter fill and mean retention time were decreased (P<0.05) by P while a P × M interaction (P<0.05) occurred for particulate passage rate. Protein combined with M produced an additional increase in particulate passage rate above
particulate digesta kinetics regulated by complex intra-ruminal relationships among intake, digestion, and rate of passage (Ward and Ward 1987). For animals consuming mature forage, feed intake, digestion, and rate of passage (Ward 1985). The effects of protein supplementation on molar proportions of VFA are conflicting and inconclusive. Results from the present study showed that molar proportions of ruminal acetate were decreased by P and not affected by M. Moreover, molar concentrations of propionate and butyrate were not affected by either P or M. McCollum and Galyean (1985) also reported decreases in molar proportions of ruminal acetate resulting from protein supplementation, while Wagner et al. (1983) found no differences in molar proportions of VFA in protein-supplemented cattle grazing rangelands. Results from the present experiment concerning monensin contrast those of Lemenager et al. (1978b), in which monensin decreased molar proportions of ruminal acetate and increased molar proportions of ruminal propionate in steers fed harvested winter range grass. Monensin levels fed by Lemenager et al. (1978b) were twice those of the present study and steers were fed in drylot, which may explain why results varied between studies. In addition, Lemenager et al. (1978b) fed monensin as a pulse dose, as opposed to a continuous release as in the present study. Valerate and isovalerate were increased by P (P<0.10 and P<0.05, respectively) and not affected by M. Isobutyrate was not affected by either P or M. On protein deficient diets, minor acids would be expected to increase as a result of protein supplementation because branched-chain acids are primarily derived from degradation of dietary protein (Ørskov 1982).

### Conclusions

Ruminal ammonia-N concentrations were increased by supplemental protein, which led to a concomitant increase in organic matter digestibility. While forage intake was not increased by protein supplementation, gastrointestinal tract dry matter fill and particulate mean retention time were reduced. When supplemental protein was combined with monensin, an additional increase in particulate passage rate occurred. Increased forage digestibility and particulate passage rate, along with reduced gastrointestinal mean retention time, could combine to provide cattle grazing winter range additional energy at a time when it is most critical to meet nutrient requirements.

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### Table 3. Ruminal fluid flow and volume, rumen pH and rumen ammonia-N concentration as influenced by supplemental protein and monensin in steers grazing native range.

<table>
<thead>
<tr>
<th>Item</th>
<th>Treatment</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th>EMS(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NR</td>
<td>NR+M</td>
<td>NR+P</td>
<td>NR+P+M</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluid passage rate, %/h</td>
<td>10.0</td>
<td>9.8</td>
<td>11.7</td>
<td>10.6</td>
<td>3.14</td>
<td></td>
</tr>
<tr>
<td>Turnover time, h</td>
<td>10.2</td>
<td>10.7</td>
<td>8.8</td>
<td>9.9</td>
<td>3.10</td>
<td></td>
</tr>
<tr>
<td>Flow rate, liters/h</td>
<td>3.7</td>
<td>5.9</td>
<td>4.1</td>
<td>5.9</td>
<td>14.2</td>
<td></td>
</tr>
<tr>
<td>Volume, liters</td>
<td>37.4</td>
<td>56.6</td>
<td>35.0</td>
<td>51.1</td>
<td>887.5</td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>6.54</td>
<td>6.56</td>
<td>6.40</td>
<td>6.41</td>
<td>0.00</td>
<td></td>
</tr>
<tr>
<td>Ammonia-N, mg/100 ml</td>
<td>.62</td>
<td>.58</td>
<td>2.62</td>
<td>3.25</td>
<td>2.70</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)NR=native range, P=protein pellet, M=monensin.

### Table 4. Ruminal concentrations of total volatile fatty acids (VFA) and molar proportions of acetic, propionate, butyrate, valerate, isovalerate and isobutyrate as influenced by supplemental protein and monensin in steers grazing native range.

<table>
<thead>
<tr>
<th>Item</th>
<th>Treatment</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th>EMS(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NR</td>
<td>NR+M</td>
<td>NR+P</td>
<td>NR+P+M</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total VFA</td>
<td>55.2</td>
<td>55.6</td>
<td>67.5</td>
<td>64.4</td>
<td>94.2</td>
<td></td>
</tr>
<tr>
<td>Acetate</td>
<td>75.2(^a)</td>
<td>74.3</td>
<td>72.9</td>
<td>71.8</td>
<td>2.24</td>
<td></td>
</tr>
<tr>
<td>Propionate</td>
<td>17.3</td>
<td>18.7</td>
<td>18.2</td>
<td>18.8</td>
<td>2.82</td>
<td></td>
</tr>
<tr>
<td>Butyrate</td>
<td>6.9</td>
<td>6.4</td>
<td>6.9</td>
<td>7.4</td>
<td>.903</td>
<td></td>
</tr>
<tr>
<td>Isovalerate</td>
<td>.19</td>
<td>.19</td>
<td>.63</td>
<td>.67</td>
<td>.62</td>
<td></td>
</tr>
<tr>
<td>Valerate</td>
<td>.20(^a)</td>
<td>.19</td>
<td>.71</td>
<td>.53</td>
<td>.023</td>
<td></td>
</tr>
<tr>
<td>Isobutyrate</td>
<td>.16(^a)</td>
<td>.20</td>
<td>.59</td>
<td>.76</td>
<td>.042</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)NR=native range, P=protein pellet, M=monensin.

\(^b\)EMS=Error mean squares.

Protein effect significant (P<0.01).

Protein effect significant (P<0.05).
Literature Cited


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