Salinomycin and virginiamycin for lactating cows supplemented on pasture

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ABSTRACT: Animals on pasture generally show higher feed efficiency as a result of the use of antibiotics. This study evaluated the effect of the antimicrobials salinomycin and/or virginiamycin on production and the ruminal parameters of supplemented dairy cows grazing on Panicum maximum cv. Tanzania. Twelve Holstein/Zebu multiparous cows were used, distributed in three Latin squares, one for the evaluation of ruminal parameters, and the others for production parameters. Cows on pasture were fed 50 % of their estimated intake with corn silage and concentrate supplements containing salinomycin, virginiamycin or a combination of additives, in doses of 120 and 150 mg kg⁻¹, respectively. There were no differences in milk production and composition, energy and nitrogen balance, dry matter digestibility and feeding behavior. However, salinomycin and virginiamycin each reduced pasture and total dry matter intake by about 14 % and 10 %, with a consequent improvement in feed efficiency.

Keywords: Panicum maximum, digestibility, feeding behavior, ionophores, nitrogen balance

Introduction

Salinomycin, a polyester antibiotic ionophore produced by Streptomyces albus, has been effective in increasing the production of cattle on high-grain diets (Merchen and Berger, 1985) or on pasture (Bagley et al., 1988). The mechanism of action is related to the transport of high-affinity cations into the cell. This impairs the normal flux of ions through the cell membrane, and reduces the growth rates of susceptible microorganisms as a result of energy loss from the cell.

Produced during fermentation of Streptomyces virginiae, virginiamycin is an antibiotic belonging to the class of streptogramins. Composed of two factors, M and S, with synergistic functions, virginiamycin can be linked specifically and irreversibly to ribosomal units. This inhibits peptidation formation, with a consequent reduction in growth [bacteriostasis effect] or even death of bacteria [bactericidal effect] (Boon and Dewart, 1974).

Gram-negative microorganisms are generally resistant to ionophore and non-ionophore antibiotics, because their outer membrane is impermeable to many macromolecules. The increase of gram-negative bacteria in the rumen improves energy and protein status, due to the change in the ruminal fermentation pattern, which increases propionate production and reduces methane and deamination of amino acids (McGuffey et al., 2001). In dairy herds, reduction of non-esterified fatty acids, ketones and β-hydroxybutyrate, and increases in the availability of glucose and amino acids associated with these antibiotics have resulted in lower body-fat mobilization, and higher milk production, milk-protein content and feed efficiency (Erasmus et al., 2008).

Animals on pasture or fed with higher proportions of forage generally show poor responses to the use of antibiotics (Clayton et al., 1999). However, the use of combinations of antibiotics has increased feed efficiency and milk production, together with the reduction of metabolic problems associated with the use of body reserves (Erasmus et al., 2008).

The small number of experiments which use either salinomycin or virginiamycin, and also a combination of these antibiotics, suggested that useful information could be gained from an evaluation of the effects of these treatments on the physiological and production parameters of dairy cattle supplemented on pasture.

Materials and Methods

The experiment was conducted from February through April 2010 in Santo Antônio do Leverger, in the state of Mato Grosso, Brazil, at 141 m altitude, 15°51'56" S and 56°04'36" W. The climate was Cwa of Köppen, tropical, with two distinct seasons, a rainy summer (Oct through Mar) and a dry winter (Apr through Sep). The mean annual temperature and annual rainfall are 24 °C and 1,300 mm, respectively.

Twelve Holstein/Zebu multiparous dairy cows, after the peak of lactation, were used in three 4 × 4 Latin Square-design experiments, grouped according to the volume of milk production. The first group consisted of rumen-cannulated cows to evaluate nutritional parameters. These cows were producing 9 kg d⁻¹ of milk and averaged 354 ± 35 kg Body Weight [BW]. The other two evaluated production parameters. These cows weighed on average 460 ± 23 and 514 ± 32 kg BW and were producing 13 and 15 kg d⁻¹ of milk, respectively. The cows were adapted to each trial over an 11-day period. During this period, the cows were fed twice daily (06h30 and 15h30) with a total of 3 kg of the same supplement.

The experiment consisted of four experimental periods of 21 days each; the first 14 days were used for diet adaptation and the following seven for data collection. Animals kept on pasture were fed simultaneously at 7h00 and 15h30, after milking, with

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Additives for dairy cows

10.02 L in 83.62 3.59 29.63 53.57, −1 29.81 for experimental diets with less than 3 % of ether extract.

Concentrate supplement was calculated according to the models proposed −1 3.95 28.63 0.50 55.47 2 −1 3 2 −1 6.74 8.39 0.50 2.62 10.10 41.56 61.24 9.43.

The treatments consisted of additives, as follows: i) Control diet (C); ii) salinomycin 120 mg kg−1 of concentrate supplement [S]; iii) virginiamycin 150 mg kg−1 of concentrate supplement [V]; and iv) salinomycin and virginiamycin 120 and 150 mg kg−1 of supplement [SV].

The experimental area consisted of 12 plots of Tanzania grass (Panicum maximum, Jacq. cv. Tanzania); each plot had an area of 2,500 m² and was managed rotationally. Forage availability was estimated when the animals entered each paddock, by measuring the sward height at 20 points. Only paddocks with a mean initial sward height of 75 cm were used. The cows were removed when the sward height was reduced to approximately 40 cm. During the experimental period, the pasture was fertilized with 88 kg ha−1 of nitrogen and 88 kg ha−1 of potassium.

Hand-plucked samples were collected simulating the grazing action. The forage, feed offered and residues were weighed and sampled daily in the last seven days of each period. Forage samples were cut at ground level, in an area defined by quadrats measuring 0.5 × 0.5 m, homogenized and divided to determine fractions of green and dry leaf [leaf blade], green and dry stem [stem + sheath], and forage mass availability (kg DM ha−1) in each experimental paddock [Table 1].

Forage intake and food digestibility were estimated using external and internal markers. Fifteen grams d−1 of chromium oxide (Cr2O3), administered orally to each cow from day 8 through 15 of each experimental period, was used as an external marker to estimate the fecal excretion of individual animals. Fecal samples were collected directly from the rectum [approximately 200 g], on day 14 through 16 of the experimental period, at the following times: day 14 (17h00), 15 (11h00) and 16 (06h00).

The total digestible nutrients (TDN) and digestible (DE), metabolizable (ME) and net energy of lactation (NEL) were calculated according to the models proposed by NRC [2001]. TDN g kg−1 = digestible crude protein (CP) + digestible neutral detergent fiber (NDF) + digestible non-fibrous carbohydrates (NFC) + (2.25 × digestible ether extract [EE]). DE Mcal kg−1 was estimated by multiplying the concentration of each digestible nutrient and its heat of combustion. ME Mcal kg−1 for experimental diets with less than 3 % of ether extract was ME [Mcal kg−1] = [1.01 × DE] + 0.45. NEL for experimental diets with less than 3 % of ether extract was NEL [Mcal kg−1] = 0.703 × ME – 0.19.

Forage intake was estimated with indigestible NDF, using the following model proposed by Detmann et al. [2001]. Forage [DMI] = [(FE × MC) – MCS] MCF−1, where: FE = fecal excretion [kg d−1]; MC = marker content in feces [kg kg−1]; MCS = marker content in the supplement (kg d−1); MCF = marker content in the forage [kg kg−1].

Samples of forage, ingredients, supplements, residues and feces were pre-dried in a forced-air oven at 60 ± 5 °C for 72 h, ground, and sifted through a sieve with a 1 mm mesh size. Each sample was analyzed for dry matter [DM], organic matter [OM], ash, CP, and EE, as described by AOAC [2005] [Table 2].

### Table 2 – Proportion of ingredient of concentrate supplements and chemical composition of feeds.

<table>
<thead>
<tr>
<th>Ingredients (%)</th>
<th>Concentrate supplement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ground corn</td>
<td>71.75</td>
</tr>
<tr>
<td>Sunflower meal</td>
<td>10.10</td>
</tr>
<tr>
<td>Soybean meal</td>
<td>11.50</td>
</tr>
<tr>
<td>Limestone</td>
<td>1.15</td>
</tr>
<tr>
<td>Dicalcium phosphate</td>
<td>1.80</td>
</tr>
<tr>
<td>Urea/Ammonia sulphate</td>
<td>2.50</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>0.50</td>
</tr>
<tr>
<td>Molasses</td>
<td>0.50</td>
</tr>
<tr>
<td>Mineral premix1</td>
<td>0.20</td>
</tr>
</tbody>
</table>

### Table 1 – Morphological components of Tanzania-grass.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Dry Matter (%)</th>
<th>Ash (% in DM)</th>
<th>Crude Protein (% in DM)</th>
<th>Ether Extract (% in DM)</th>
<th>NDFap (%)</th>
<th>NFC (%)</th>
<th>Mineral premix1</th>
<th>Means</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compostion</td>
<td>31.28</td>
<td>6.77</td>
<td>14.32</td>
<td>1.97</td>
<td>61.24</td>
<td>15.68</td>
<td>0.20</td>
<td></td>
</tr>
<tr>
<td>Dry Matter (%)</td>
<td>71.75</td>
<td>10.10</td>
<td>11.50</td>
<td>1.15</td>
<td>1.80</td>
<td>2.50</td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td>Ash (% in DM)</td>
<td>8.39</td>
<td>27.47</td>
<td>6.74</td>
<td>2.66</td>
<td>53.57</td>
<td>28.63</td>
<td>0.27</td>
<td></td>
</tr>
<tr>
<td>Crude Protein (% in DM)</td>
<td>27.81</td>
<td>29.63</td>
<td>27.81</td>
<td>29.63</td>
<td>15.82</td>
<td>47.17</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ether Extract (% in DM)</td>
<td>29.63</td>
<td>26.39</td>
<td>26.39</td>
<td>25.25</td>
<td>15.68</td>
<td>47.17</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NDFap (%)</td>
<td>29.63</td>
<td>41.56</td>
<td>41.56</td>
<td>45.12</td>
<td>47.17</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NFC (%)</td>
<td>29.63</td>
<td>41.56</td>
<td>41.56</td>
<td>45.12</td>
<td>47.17</td>
<td></td>
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</tr>
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<td>Mineral premix1</td>
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<td>45.12</td>
<td>47.17</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 Mineral premix composition: 105 g kg−1 of Calcium; 7,500 mg kg−1 of Magnesium; 230 g kg−1 of sulfur; 330 mg kg−1 of cobalt; 2000 mg kg−1 of Copper; 155 g kg−1 of Iodine; 2,800 mg kg−1 of Magnesium; 220 mg kg−1 of Selenium; 6800 mg kg−1 of Zinc; Control treatment – no antibiotics; S – 120 mg kg−1 of salinomycin; V – 150 mg kg−1 of virginiamycin; NDFap (neutral detergent fiber corrected for ash and protein); NFC (non fibrous carbohydrates) = 100 - (CP % - CP % from urea + urea %) + NDF % + EE % + Ash %.
contents of NDF were determined by using α amylase
without sodium sulfite added, and corrected [NDFap],
discounting ash and neutral detergent-insoluble protein
[Mertens, 2002; Licitra et al., 1996]. Due to the presence
of urea, NFC was calculated as proposed by Hall [2000]:
NFC = 100 - (CP % - CP % derived from urea + urea %)
+ NDF % + EE % + Ash %.

Cows were milked twice daily at 06h00 and 15h00.
Milk production was recorded through a milking device,
from day 15 through 18 of each experimental period. On
days 17 and 18, proportional morning and evening milk
samples of approximately 100 mL were collected and
packed in plastic bottles with preservative. The content
of fat, protein and lactose were analyzed by infrared
spectrophotometry [IDF, 1996]. 3.5 % fat-corrected milk
production (FCM) was estimated [Sklan et al., 1992] by
the following equation: FCM in kg d\(^{-1}\) = [0.432 × kg
milk] + [16.216 × kg milk fat].

Feed efficiency (FE) was calculated as fat-
corrected milk production per total dry matter intake
and energy efficiency (EE) as Mcal of net energy of
lactation excreted on milk per Mcal of net energy of
lactation intake. Energy balance (EB, Mcal d\(^{-1}\)) was
calculated by subtracting the NE\(_{\text{c}}\) consumed from
the required amounts of net energy for maintenance
and lactation. Net energy of maintenance [NE\(_{\text{c}}\); Mcal
d\(^{-1}\)] was calculated as 0.080x BW\(^{0.75}\) and net energy
of lactation [NE\(_{\lambda}\); Mcal d\(^{-1}\)] = [0.0929 × % fat + 0.0547x%
CP + 0.0395 × % lactose] × milk production (kg d\(^{-1}\))
[NRC, 2001].

Animals were weighed every 21 days in each
experimental period, after the morning milking. Blood
samples were collected on day 21, and centrifuged
to separate the serum. Urea was determined in
deproteinized milk and serum using commercial kits
[Labtest\(^{\circledR}\)]. Urea was converted to blood urea nitrogen
by multiplying the observed values by 0.4667, which gives
the total nitrogen in the urea.

Urine spot samples were collected on day 21 of
each period and stored at -20 °C for total nitrogen
analysis. The nitrogen balance was obtained from
the difference between nitrogen intake and nitrogen
excreted in feces, urine and milk.

Ruminal fluid was collected through the ruminal
cannula on day 20, to measure pH and ammonia
concentration before [0] and 2, 4, 6 and 8 h after
the beginning of feeding in the morning. Rumen fluid pH
was immediately determined with the use of a digital
potentiometer. At each sampling, a 50-mL aliquot of
the ruminal fluid from each animal was mixed with 1 mL
of 50 % sulfuric acid and stored at -5 °C for ammonia
analyses.

The feeding behavior was assessed on day 19, for
24 h, by visual observation. Every ten minutes were
recorded activities of grazing, ruminating and idle were
recorded.

Data were statistically analyzed using PROC
MIXED. The statistical model was:

\[
Y_{ijkl} = \mu + A_{ij} + P_{i(l)} + T_{k} + Q_{l} + TQ_{kl} + e_{ijkl},
\]

where: \(Y_{ijkl}\) = observation of cow \(i\) in period \(j\) subject to
supplementation level \(k\), in Latin square \(l\); \(\mu\) = overall
effect of the mean; \(A_{ij}\) = effect of animal \(i\) in Latin
square \(l\), with \(i = 1, 2, 3, 4\); \(P_{i(l)}\) = effect of period
\(j\) in Latin square \(l\), with \(j = 1, 2, 3, 4\); \(T_{k}\) = effect of
supplementation level \(k\), where \(k = 1, 2, 3, 4\); \(Q_{l}\) = effect
of Latin square \(l\), where \(l = 1, 2\); \(TQ_{kl}\) = interaction
effect between treatment \(k\) × Latin square \(l\); \(e_{ijkl}\) = random
error associated with each observation ijkl. \(e_{ijkl} \sim \text{NID}
(0, \sigma^2)\). In the presence of a significant treatment effect,
means were compared using Tukey’s test, considering \(\alpha
= 5\%\) and \(10\%\) for tendency of error type I.

**Results and Discussion**

The intake of supplements was not different \((p >
0.05)\) between the experimental diets \((5.72 \text{ kg d}^{-1})\), and
was close to what was offered \((6 \text{ kg d}^{-1})\). Total DMI of
cows receiving the control diet averaged 11.46 kg d\(^{-1}\),
slightly lower than the level predicted by NRC [2001],
about 12.03 kg d\(^{-1}\). Administration of salinomycin or
virginiamycin reduced \((p = 0.03)\) DMI by about 14 %
and 10 %, respectively, as the nutrients NDFap and
TDN. Thus, the use of antibiotics reduced pasture
intake and, at the same time, fiber intake, below the
values suggested by Mertens [1987]. According to this
author, fiber intake is usually limited by rumen fill
when NDF intake reaches approximately 1.2 ± 0.1 %
of BW [Table 3].

Iparraguirre and Clark [2003] observed that 8
out of 12 studies on monensin for lactating cows found
no differences in DMI. Their findings demonstrate that
ionophore responses may be related to dose and
stage of lactation. In early lactation, the addition of
ionophore was able to reduce losses of body reserves
and increase available energy and animal performance
without changing DMI. However, in the mid- and late
stages of lactation, as well as in the case of beef cattle,
this was able to decrease DMI due to the lower energy
requirement [Erasmus et al., 2008].

Although most studies with virginiamycin have
been conducted on beef animals in feedlots, some
points can be related to dairy cows. Rogers et al.
[1995] analyzed seven experiments on dose response
for virginiamycin and found an improvement in feed
efficiency associated with the mean daily increase in
weight gain. None of these studies found a reduction
in DMI. However, four experiments found a numerical
increase in DMI. Furthermore, no increase in FE was observed in response to doses above
19 mg kg\(^{-1}\) DM, as observed in this study [Table 3].
Salinas-Chavira et al. [2009] reported no differences in
the average daily gain or DMI for confined Holstein steer
calves supplemented with three levels of virginiamycin
\((0, 16, \text{ or } 22.5 \text{ mg kg}^{-1})\).
Few studies conducted with a combination of two antibiotics have found similar results. Núñez et al. (2013) observed a reduction in DMI with virginiamycin (15 mg kg⁻¹) in the diet containing salinomycin (13 mg kg⁻¹), which contributed to greater FE. On the other hand, Silva et al. (2004) obtained differences in mean daily gain with Nellore steers fed 77% concentrate diet, supplemented with salinomycin, virginiamycin, or a combination of the two. However, steers receiving both salinomycin and virginiamycin mobilized the body reserves (Clayton et al., 1999; Duffield et al., 2008), in a review of monensin, noted that the dose and method of administration, in addition to the stage of lactation, could affect milk composition. The use of antibiotics often reduces amino-acid deamination, and thus losses of nitrogen in urine and milk (McGuffey et al., 2001).}

In the present study, FE was about 18% higher with salinomycin, compared to the control treatment.

The small amplitude or even absence of responses in DM and nutrient intake with the use of virginiamycin alone or in combination with salinomycin did not contribute to an improvement in feed and energy efficiency. Providing corn silage and concentrate supplement to low-producing cows in mid- to late lactation limited the response in increasing milk or milk composition production with the use of antibiotics. Virginiamycin used alone tended (p < 0.10) to increase FE by 6%.

The present experimental diets led to a positive energy and nitrogen balance (Table 4). This indicates that energy and protein requirements were satisfied, and the low milk production was related to the productivity of cows and the stage of lactation. Body-weight changes were positive for all animals, averaging 89, 327, 369 and 61 g d⁻¹, respectively, for the control, salinomycin, virginiamycin, and salinomycin with virginiamycin treatments. Measurements of body-weight change help to evaluate the real benefit of additive use, which increases FE without animal weight loss.

Similarly, milk composition (fat, protein and lactose) was not affected (p > 0.05) by salinomycin or virginiamycin. Duffield et al. (2008), in a review of monensin, noted that the dose and method of administration, in addition to the stage of lactation, could affect milk composition. The use of antibiotics often reduces amino-acid deamination, and thus losses of nitrogen in urine and milk (McGuffey et al., 2001). Therefore, the observed reduction in milk urea nitrogen ([MUN] and increase in milk protein [MP] were
expected. However, even cows receiving the control diet showed low levels of both components (9.24 mg dL$^{-1}$ and 3 %).

Milk urea nitrogen is a valuable tool for monitoring dietary protein [Hof et al., 1997]. Levels below 10 mg dL$^{-1}$ with less than 3.2 % of MP, as observed in this study, may indicate that the diet contained low levels of crude protein and rumen-degradable protein (RDP).

Animals were supplemented with corn silage and a supplement concentrate with high energy content that requires higher CP and RDP diets. The commercial concentrate with 220 g kg$^{-1}$ CP used here may have not met the animals’ RDP requirements, which probably influenced the MUN and MP levels and the lack of antibiotic effects.

The inclusion of antimicrobials did not affect DM and nutrient digestibility ($p > 0.05$) of the experimental diets, except for NDF digestibility (Table 5). The effect of virginiamycin on DM digestibility has rarely been investigated [Salinas-Chavira et al., 2009]. These authors observed an improvement in DM digestibility in swine, which was attributed to an increase in intestinal feed retention and a reduction in harmful bacteria [Ravindran et al., 1984].

Ionophores could improve fiber digestibility, mainly because they reduce feed intake and consequently affect the passage rate of solids. However, NDF digestibility was lower ($p = 0.02$) when the combined antibiotics were used, compared to the control diet (C), or to salinomycin alone (S). Lower fiber digestibility in this case may be related to the tendency for ammonia reduction with the use of virginiamycin.

Rumen ammonia nitrogen levels [RAN] were similar ($p > 0.05$) in the experimental diets, above 8 mg dL$^{-1}$ - the minimum needed to maximize fiber degradation. A reduction of ruminal ammonia and blood urea nitrogen (BUN) with antibiotics would be expected, due to the reduction in amino acid deamination.

Although not significant ($p > 0.05$), the addition of virginiamycin decreased RAN by 14 % and reduced it to below 8 mg dL$^{-1}$ for a long period during the day (Figure 1). Moreover, a 9 % BUN reduction was observed, compared to the control diet. In this study, mean levels of BUN were 20.54 mg dL$^{-1}$, i.e., approximately at the limit of 19-20 mg dL$^{-1}$ at which dietary nitrogen losses could occur in dairy cows [Oliveira et al., 2001].

No differences were observed ($p > 0.05$) in ruminal pH between diets [Table 5]. The pH of high-forage diets did not change with the addition of antimicrobials [Clayton et al., 1999]. Grazing, ruminating and idle times did not differ ($p > 0.05$) between experimental diets [Table 6]. Other parameters such as pasture selection and bite size influenced the intake rates, as salinomycin or virginiamycin reduced pasture intake without affecting grazing time.

Grazing time reflects the ease of forage access and removal. The time spent grazing varies between 359 and 711 min d$^{-1}$ [Krysl and Hess, 1993], and grazing times longer than 480-540 min d$^{-1}$ probably indicate limited conditions for forage intake [Hodgson, 1990]. In this study, the mean grazing time was 328 min d$^{-1}$, as a result of good quality forage [Table 2], pasture availability [Table 1] and the provision of part of the diet in feeders.

The addition of salinomycin or virginiamycin for mid-lactation dairy cows improved FE, because it reduced DMI without affecting milk production and milk composition.

**Table 5 – Values of dry matter digestibility, pH, ammonia nitrogen and blood urea.**

<table>
<thead>
<tr>
<th>Items</th>
<th>Treatments$^1$</th>
<th>Means</th>
<th>SEM$^1$</th>
<th>$p^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DDM$^4$</td>
<td>C 65.62 S 66.20 V 65.26 SV 64.19</td>
<td>65.32</td>
<td>2.11</td>
<td>0.15</td>
</tr>
<tr>
<td>DCP$^4$</td>
<td>71.44 S 73.22 V 73.01 SV 70.25</td>
<td>71.98</td>
<td>2.20</td>
<td>0.22</td>
</tr>
<tr>
<td>DNDF$^4$</td>
<td>C 64.20 S 63.46 V 61.48 SV 60.01</td>
<td>62.29</td>
<td>3.93</td>
<td>0.02</td>
</tr>
<tr>
<td>DEE$^4$</td>
<td>80.05 S 77.33 V 78.82 SV 80.53</td>
<td>79.18</td>
<td>7.00</td>
<td>0.91</td>
</tr>
<tr>
<td>DNFC$^4$</td>
<td>C 77.47 S 76.69 V 78.97 SV 79.60</td>
<td>78.18</td>
<td>2.63</td>
<td>0.41</td>
</tr>
<tr>
<td>RAN$^6$</td>
<td>C 11.46 S 11.01 V 9.86 SV 9.54</td>
<td>10.94</td>
<td>1.48</td>
<td>0.09</td>
</tr>
<tr>
<td>pH$^8$</td>
<td>C 6.5 S 6.4 V 6.5 SV 6.6</td>
<td>6.5</td>
<td>0.15</td>
<td>0.11</td>
</tr>
<tr>
<td>BUN$^9$</td>
<td>C 20.78 S 22.17 V 19.85 SV 19.36</td>
<td>20.54</td>
<td>4.96</td>
<td>0.53</td>
</tr>
</tbody>
</table>

$^1$Control (C), salinomycin (S), virginiamycin (V), and salinomycin and virginiamycin (SV); $^2$means followed by different letters in the line statistically differ by Tukey’s test; $^3$SEM = standard error of mean; $^4$DDM = digestibility coefficient of total dry matter; $^5$DCP = digestibility coefficient of crude protein; $^6$DNDF = digestibility coefficient of neutral detergent fiber; $^7$DEE = digestibility coefficient of ether extract; $^8$DNFC = digestibility coefficient of non-fibrous carbohydrates; $^9$RAN = ruminal ammonia nitrogen; $^{10}$pH = ruminal pH; $^{11}$BUN = blood urea nitrogen.

**Figure 1 – Ruminal Ammonia Nitrogen (RAN) before 0, and at 2, 4, 6 and 8 h after feeding in control (C), salinomycin (S), virginiamycin (V), and salinomycin and virginiamycin (SV). Bars show the standard error of mean.**

**Table 6 – Feeding behavior of lactating dairy cows (min d$^{-1}$).**

<table>
<thead>
<tr>
<th>Variables</th>
<th>Treatments$^1$</th>
<th>Means</th>
<th>SEM$^1$</th>
<th>$p^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grazing</td>
<td>C 295 S 316 V 324 SV 301</td>
<td>328</td>
<td>42</td>
<td>0.12</td>
</tr>
<tr>
<td>Rumination</td>
<td>C 539 S 537 V 559 SV 567</td>
<td>550</td>
<td>27</td>
<td>0.33</td>
</tr>
<tr>
<td>Idle</td>
<td>C 549 S 539 V 496 SV 501</td>
<td>519</td>
<td>38</td>
<td>0.13</td>
</tr>
</tbody>
</table>

$^1$Control (C), salinomycin (S), virginiamycin (V), and salinomycin and virginiamycin (SV); $^2$means followed by different letters in the line statistically differ by Tukey’s test; $^3$SEM = standard error of mean.
References


