MORPHOLOGICAL RESPONSE OF THE RUMINAL AND OMASAL MUCOSAE TO THE VARIATION IN DIET ENERGY

Resposta morfológica das mucosas do rúmen e omaso à variação da energia na dieta

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ABSTRACT
The absorption capacity of the rumen responds positively to direct and indirect stimulation by VFA; there is also evidence that the wall of the omasum also responds to these stimuli. To further investigate these reports, we compared the tissue morphologies of rumen and omasum biopsy samples. Four cows surgically fitted with ruminal cannulas were sequentially fed two diets. These diets included corn silage (S) and a combination of corn silage and commercial concentrate (CS). The animals were fed S for the first 18 days of the experiment, followed by CS for the next 18 days. They were then fasted for 72 h (F), and then had at least 18 days of re-feeding. Biopsy samples were taken from the blade of the omasum and the ventral sac of the rumen at different time points during each diet: samples were taken from S-diet animals at day 18; CS, at days 4 and 18; F, at the end of the 72 h; and re-fed animals, at days 4, 12, and 18. The mitotic index of the basal layers of the ruminal and omasal epithelia and the VFA concentration in the rumen were higher after 4 days of CS diet. There was a positive correlation between the mitotic indices of the rumen and omasum. The width of the ruminal papillae varied with different diets, and was highest on day 18 of the CS diet. Our results indicate that stimulation of cell division due to increased dietary energy simultaneously affected both compartments of the stomach.

Index terms: Acidosis; morphology; physiology; transition diet.

INTRODUCTION
After calving, dairy cows typically receive high-calorie lactation diets. These diets are rich in carbohydrates that quickly ferment in the reticulorumen, resulting in increased production of volatile fatty acids (VFA). VFA produced in the reticulorumen are absorbed through the rumen wall and, after passage to the omasum, are incorporated in the fluid phase. However, if the VFA are not metabolized in the reticulorumen epithelium at the same rate that they are produced, they accumulate in the ruminal environment and cause rumen acidosis (Plaizier et al., 2009). Dairy cows often receive a high-calorie diet prepartum in order to prepare the ruminal and omasal epithelia to absorb the high postpartum concentrations of VFA. This high-calorie diet induces proliferation of the ruminal epithelium (Dirksen et al., 1984), which is important for control of rumen acidosis. However, there is little information on the morphological response of the omasum to diets of varying caloric content, even though the omasum receives 36% (Volker; Allen, 2003) to 45% (Resende Júnior et al., 2006a) of VFA produced in the rumen. Extension of the absorptive surface of the omasum may play a key role in VFA absorption, thereby decreasing...
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the amount passed to the abomasum and incorporated into the fluid phase. Excess VFA absorbed in the abomasum is correlated to decreased motility that is a precursor to digestive disturbances such as abomasum displacement (Bolton et al., 1976).

The purpose of this study was to compare the morphological responses of the mucosae of the ruminal and omasal epithelia, and the thickness of the keratinized and non-keratinized layers of the ruminal and omasal epithelia.

MATERIAL AND METHODS

All procedures were approved by the Bioethical Committee for Animal Utilization at the Federal University of Lavras, Brazil.

Silicone canulas four inches in diameter were surgically attached to the dorsal rumen sacs of four non-pregnant, non-lactating cows of unspecified breeds and unknown ages (mean body weight [BW], 385 kg); these animals were sequentially fed two diets. These diets consisted of These only corn silage (S) and corn silage plus commercial concentrate (50% dry matter) (CS). The animals were kept in a tie-stall barn on sand bedding with ad libitum access to food and water. They were fed with a total mixed ration once daily at 0700 h. The experiment was conducted over 57 days; in this time, seven sequential ingestion periods, defined as treatments, were carried out in order to compare the morphological behaviors of the rumen and omasum in response to different diets at different time points. The treatments were as follows: 18 days S diet (18S), four days CS diet (4CS), 18 days CS diet (18CS), three days of fasting (F), four days re-feeding with CS (4RCS), 12 days re-feeding with CS (12RCS), and 18 days re-feeding with CS (18RCS). These treatment time points were chosen in accordance with experiments by Goodlad (1981) and Resende Júnior et al. (2006b), who observed higher MI in the ruminal epithelium four days after the start of concentrate feeding and an increase in absorptive surface 12 days after introduction of a concentrate diet, respectively.

Food was weighed before and after feeding to measure consumption. Samples were collected and frozen at -20 °C until further analysis. After all samples had been collected, samples were thawed, grouped by week of collection, dried in a ventilated oven at 60 °C for 72 h, ground in a mill with a 1.0-mm sieve, and dried again in a ventilated oven at 105 °C. Dry matter intake (DMI) was estimated using the average feed consumption over the final three days of each ingestion period.

Crude protein (CP) content was determined by steam digestion with a Microkjeldahl device (Association Of Official Analytical Chemists - Aoac, 1997); ash content was determined by incineration at 550 °C over 8 h in a muffle; NDF was determined as described by Van Soest, Robertson and Lewis (1991); and ether extract was determined as described by AOAC (1997).

Samples of ruminal fluid were collected by suction from the ventral sac 6 h after feeding on the last day of each treatment using a rigid PVC tube (3/4 inch) with several holes in the distal third, coupled to a hose and a plastic bottle. Ruminal fluid was separated into 10-mL aliquots and preserved in 200 µL sulfuric acid solution (50% v/v). The samples were frozen at -20 °C until further VFA analysis. VFA concentrations were determined by gas-liquid chromatography (CP-3800 Gas Chromatograph Varian, Varian Chromatography Systems, California, USA). After thawing, the ruminal fluid samples were centrifuged at 8,855 g for 15 min and the supernatant was collected. Next, 150 µL caproic acid was added to 250 µL of the supernatant as an internal standard (N Caproic Acid, Sigma-Aldrich Chemie GmbH, Steinheim, Germany), and 100 µL of distilled water was added. A solution mixture of acetic, propionic, butyric, and valeric acid was used as an external standard. The samples were analyzed in a composite phase capillary column (length, 25 m; internal diameter, 0.25 mm) and a 2-mm thick film (CP-Wax 58 - FFAP- CB, Varian Analytical Instruments, California USA). A temperature ramp was used in the column oven: the temperature was first increased to 65 °C within 30 s, increased at a rate of 20 °C per minute for the next 3 min to reach 125 °C, and finally increased 50 °C per minute to reach 170 °C. The total analysis time was 4.9 min.

On the last day of each treatment, the reticulorumen was evacuated approximately 10 h after feeding, and biopsies measuring approximately 1 cm² were collected from the rumen ventral sac wall (recessus ruminis) and the omasal lamina (lamina omasi). Biopsies of the rumen were performed using surgical scissors after exteriorization of the rumen mucosa through the cannula. For omasal laminal biopsies, the researcher was required to reach into the ostium reticulo-omasicum through the rumen cannula to obtain the sample with a pliers-like device. The rumen tissue fragment was separated into two aliquots: the first was immediately placed in a vial containing phosphate buffer solution (PBS; pH 7.4) and cooled to 7 °C until macroscopic analysis, while the second ruminal aliquot and omasal tissue fragments were fixed in Bouin’s liquid
for 22 h and stored in 70% alcohol until routine histological processing.

The evaluated variables included the number of papillae per cm² of epithelial wall, mean papillae area, mean papillae height, mean papillae width, and total absorptive area of the rumen wall.

Four reviewers counted the number of papillae on each tissue fragment, and the average calculated for each animal. The area and height of the rumen papillae were measured as described by Resende Júnior et al. (2006b) and modified by Daniel, Resende Júnior and Cruz (2006). Briefly, twelve papillae were cut at their base with surgical scissors, placed in a petri dish, and, together with the tissue fragment without papillae, their images scanned (HP Deskjet F380 All-in-One). A 1-cm scale was placed beside the petri dish and scanned with the fragment for subsequent calibration of the image analysis software. The heights and areas of the scanned images were estimated using the UTHSCSA Image Tool analysis program (http://ddsdx.uthscsa.edu/dig/itdesc.html). Os fragmentos fixados em Boiun foram desidratados em série crescente de álcool etílico a 70º, 80º, 90º GL e absoluto, ficando o fragmento imerso por 30 minutos em cada solução.

The height, width, and area of the rumen papillae and the MI of the ruminal and omasal epithelium basal layers were estimated in 5-µm-thick sections stained with hematoxylin and eosin. To measure the thickness of the rumen and omasum epithelium keratinized layers (TKL) and non-keratinized layers (TNKL), fragments were stained with Masson’s trichrome.

Using a light microscope, at 400× magnification, the nuclei of 2,000 cells each from the ruminal epithelium basal layer and of the omasum lamina were counted for MI determination. All nuclei showing mitotic figures were counted, and the MI was calculated as the percentage of the total nuclei counted. Three evaluators each counted the nuclei; their mean value was the MI.

TKL and TNKL were determined using images from four microscopic fields per slide captured by a camera attached to a light microscope at 400× magnification. One image of a slide with a scale was used for posterior calibration of the image analysis software (UTHSCSA Image Tool). The total area of the epithelium and the area of keratinized epithelium were measured. The length of the epithelium in each photographed field was also measured. The total thickness of the epithelium and TKL were obtained by dividing the area by the respective length of the field. The TNKL was calculated as the difference between these measurements. The mean value of the four fields was used for data analysis, as described by Daniel, Resende Júnior and Cruz (2006).

The total thickness, the thickness of the TKL/ TNKL of the ruminal and omasal epithelia, and the MI of the epithelium basal layer of the rumen and omasum were analyzed as repeated measures using the MIXED tool in the SAS statistical program using the following model: $Y_{ijk} = \mu + \alpha_i + \beta_j + \gamma_k + \delta_{jk} + \epsilon_{ijk}$, where $\mu$: overall mean; $\alpha_i$: fixed effect of the compartment (i = 1 to 4); $\beta_j$: fixed effect of the treatment (j = rumen or omasum); $\gamma_k$: fixed effect of the interaction between cow and compartment (error term used to test the effect of compartment); $\delta_{jk}$: fixed effect of the treatment (k = 1 to 7); $\epsilon_{ijk}$: residual error (error term used to test the effect of treatment and the effect of the interaction between the compartment and treatment). The DMI, VFA concentration, area, height and width of the rumen papillae, the number of rumen papillae per cm², and the entire absorptive surface of the rumen were analyzed with the generalized linear model (GLM) tool in SAS using the following model: $Y_{ij} = \mu + \alpha_i + \beta_j + \epsilon_{ij}$, where $\mu$: overall mean; $\alpha_i$: effect of the cow considered as block (i = 1 to 4); $\beta_j$: effect of the treatment (j = 1 to 7); $\epsilon_{ij}$: residual error, assumed independently and identically distributed with a normal distribution, with a mean of zero and variance $\sigma^2$. The means of these variables were compared by Tukey post-hoc analysis. The MI of the rumen and omasum were compared by linear regression using the REG tool, and the hypothesis that the slope is equal to 1 was tested using the TEST tool within SAS.

**RESULTS AND DISCUSSION**

The experimental diets (Table 1) had non-fibrous carbohydrate values (NFC) comparable to simulations of pre-partum diets based on forage, and high-energy lactation diets based on forage and concentrates (Rabelo et al., 2003). Theoretically, the transition from S to CS diet mirrors the change that occurs when a transition diet is introduced to pre-partum dairy cows in order to prepare the rumen epithelium for higher fractional rates of VFA production. The fasting period simulates the eventual fall of DMI that occurs peripartum, and the re-feed period simulates the postpartum introduction of the high-energy lactation diet. However, DMI (Table 2; Figure 1) was low in the animals in this study, particularly during the re-feeding period. This finding is in contrast to reports of animals in early lactation, likely because the animals in this study had a lower metabolic demand, as they were neither pregnant nor lactating. While the ruminal epithelium is known to respond positively to introduction of high-
energy diets (Dirksen et al., 1984; Melo et al., 2013), the response of the omasal epithelium to high-energy diets has not been evaluated.

Table 1: Nutritional composition of diets consumed by cows during ingestion periods.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Diet</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>S</td>
</tr>
<tr>
<td>DM, % as fed</td>
<td>39.3</td>
</tr>
<tr>
<td>NDF, % of DM</td>
<td>47.0</td>
</tr>
<tr>
<td>Ash, % of DM</td>
<td>3.8</td>
</tr>
<tr>
<td>Fat, % of DM</td>
<td>3.9</td>
</tr>
<tr>
<td>CP, % of DM</td>
<td>9.2</td>
</tr>
<tr>
<td>NFC1, % of DM</td>
<td>36.1</td>
</tr>
<tr>
<td>TDN2, % of DM</td>
<td>48.3</td>
</tr>
</tbody>
</table>

1NFC, Non-fibrous carbohydrates estimated by the equation: 100 – (%CP+%NDF+%Fat+%Ash) 2TDN, total digestible nutrients, estimated according to NRC (2001).

The DMI and total digestible nutrients (TDN) were greater 18 days after introduction of the high-energy diet (CS) ($P < 0.05$), as expected, as there was a higher concentration of nutrients per unit of DMI in the diet concentrates. The fasting period appears to have been detrimental to the restoration of DMI levels; during the 18 days of re-feeding, DMI did not reach the same level as 18 days after introduction of the high-energy diet before fasting. This finding is similar to reports in postpartum dairy cows of a peripartum drop in DMI (Hayirli et al., 2002).

Vazquez-Añon et al. (1994) observed decreased DMI two days before delivery and a 60% reduction in DMI three days earlier in nine dry pregnant female cows. DMI increases gradually postpartum, peaking between 10 and 14 weeks of lactation (National Research Council-Nrc, 2001).

The VFA concentration was higher on day 4CS, most likely because the diet change resulted in increased carbohydrate intake and fermentation in ruminal epithelia not yet adapted to the new rate of VFA production (Table 2). Eighteen days after introduction of the high-energy diet, the lower concentration of VFA in rumen fluid likely indicated that the rumen had adapted to the new conditions (Melo et al., 2013; Resende Júnior et al., 2006b), probably due to higher fractional absorption rate. The absence of a fermentable substrate during the fasting period explains decreased fluid VFA concentrations during this time.

Table 2: Dry matter intake (DMI), rumen of volatile fatty acid (VFA) concentrations, and rumen morphology of cows at different time points.

<table>
<thead>
<tr>
<th></th>
<th>18S</th>
<th>4CS</th>
<th>18CS</th>
<th>F</th>
<th>4RCS</th>
<th>12RCS</th>
<th>18RCS</th>
<th>EPM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMI, kg/d</td>
<td>6.0</td>
<td>6.8</td>
<td>8.4</td>
<td>0.0</td>
<td>5.1</td>
<td>6.6</td>
<td>7.1</td>
<td>0.75</td>
<td>0.04</td>
</tr>
<tr>
<td>TDN, kg/d</td>
<td>2.8</td>
<td>3.5</td>
<td>4.4</td>
<td>0.0</td>
<td>2.9</td>
<td>3.7</td>
<td>4.0</td>
<td>0.41</td>
<td>0.02</td>
</tr>
<tr>
<td>Total VFA, mM</td>
<td>106.4</td>
<td>148.5</td>
<td>115.6</td>
<td>7.0</td>
<td>77.5</td>
<td>64.7c</td>
<td>68.4c</td>
<td>9.62</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Acetate, mM</td>
<td>76.8a</td>
<td>83.8a</td>
<td>78.1a</td>
<td>5.5e</td>
<td>46.5</td>
<td>44.2d</td>
<td>48.7bc</td>
<td>7.92</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Propionate, mM</td>
<td>18.6ab</td>
<td>23.9a</td>
<td>23.3a</td>
<td>1.0c</td>
<td>23.5a</td>
<td>14.1b</td>
<td>12.8b</td>
<td>2.53</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Butyrate, mM</td>
<td>12.3</td>
<td>40.7</td>
<td>14.2</td>
<td>0.5</td>
<td>7.5</td>
<td>40.7</td>
<td>6.8</td>
<td>10.11</td>
<td>0.18</td>
</tr>
<tr>
<td>Acetate/propionate</td>
<td>4.3b</td>
<td>3.5b</td>
<td>3.4b</td>
<td>5.7a</td>
<td>2.0c</td>
<td>3.4b</td>
<td>3.8b</td>
<td>0.45</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Area rumen, m²</td>
<td>5.1</td>
<td>6.6</td>
<td>5.1</td>
<td>6.0</td>
<td>5.6</td>
<td>6.1</td>
<td>6.0</td>
<td>0.70</td>
<td>0.61</td>
</tr>
<tr>
<td>Papillae /cm²</td>
<td>48</td>
<td>75</td>
<td>53</td>
<td>51</td>
<td>55</td>
<td>62</td>
<td>61</td>
<td>16.0</td>
<td>0.89</td>
</tr>
<tr>
<td>Area papillae, cm²</td>
<td>0.15</td>
<td>0.19</td>
<td>0.15</td>
<td>0.22</td>
<td>0.16</td>
<td>0.18</td>
<td>0.18</td>
<td>0.021</td>
<td>0.22</td>
</tr>
<tr>
<td>Length papillae, cm</td>
<td>0.56</td>
<td>0.66</td>
<td>0.48</td>
<td>0.66</td>
<td>0.49</td>
<td>0.55</td>
<td>0.54</td>
<td>0.073</td>
<td>0.12</td>
</tr>
<tr>
<td>Area/Length papillae</td>
<td>0.27d</td>
<td>0.51bc</td>
<td>1.40a</td>
<td>0.72bc</td>
<td>0.84b</td>
<td>0.33d</td>
<td>0.33cd</td>
<td>0.142</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

18S = 18 days eating corn silage; 4CS = 4 days eating 50% corn silage, 50% concentrate; 18CS = 18 days eating 50% corn silage, 50% concentrate; F = 3 days of fasting; 4RCS = 4 days re-feeding with 50% corn silage, 50% concentrate; 12RCS = 12 days re-feeding with 50% corn silage, 50% concentrate; 18RCS = 18 days re-feeding with 50% corn silage, 50% concentrate. Means with the same letter do not differ significantly, Tukey test ($\alpha = 0.05\%$).
With re-feeding, fluid VFA concentrations were lower than in the first high-energy diet period, while DMI and TDN were similar. It is possible that 72 h of fasting was not sufficient for the rumen lose its absorptive capacity. Therefore, during the re-feeding period, the ruminal epithelium was still able to tolerate the increased rate of VFA production. This lends support to the hypothesis that although there is drop in the DMI peripartum, the transition diet introduced before delivery is sufficient to promote adaption of the rumen epithelium to provide better absorptive capacity postpartum.

Teófilo et al. (2009) studied dairy cows in the transition period and found increased absorptive surface in the rumen of postpartum cows fed a high-NFC diet for three weeks prior to delivery. Anderson, Sehasted and Invartsen (1999) and Reynolds et al. (2004) discouraged use of transition diets based on the understanding that peripartum fasting would result in the rumen epithelium returning to the absorptive capacity it had before introduction of the transition diet, an understanding that appears to be incorrect.

Propionate concentration increased after the diet switch from S to CS. Silage-based diets have higher NDF content (Table 1). However, NDF are slowly degradable carbohydrates, resulting in a higher rumen pH that favors cellulolytic bacteria and a metabolic pathway more focused on the production of acetate. Introduction of a high-energy diet containing rapidly fermentable carbohydrates results in increased ruminal VFA concentration that decrease that ruminal pH, which favors the proliferation of amylolytic bacteria, and a metabolic pathway more geared towards production of propionate (Gregorio et al., 1982). This dynamic is shown with the change of production focus from acetate to propionate (Table 2), which is directly proportional to pH and is a standard indicator of fermentation and balance in the rumen. Although we did not measure the ruminal pH, VFA concentration is known to be the major determinate of pH (Pereira; Armentano, 2000). Therefore, changes in VFA concentration in this study can be considered inversely proportional to ruminal pH changes and thus the acetate: propionate production ratio.

There was no difference between the ingestion periods in the total estimated area of the rumen. This estimate considers the area and the number of buds on tissue fragments obtained by biopsy, and there were no statistical differences between experimental periods. The values of macroscopic measurements of the extent of absorptive surface were not consistent with the microscopic estimation of MI (Figure 2), as they

Figure 1: Average daily dry matter intake (DMI, black bars) and NDT (white bars), during the different ingestion periods of non-pregnant, non-lactating cows. They were fed for 18 days with corn silage (18S); 4 (4CS) and 18 (18CS) days with 50% corn silage and 50% concentrate with dry matter basis; 72 h fasting (F); and 4 (4RCS), 12 (12RCS), and 18 (RCS) days of re-feeding 50% silage and 50% concentrates. P < 0.04 for overall effect of ingestion period.
differed between ingestion periods and were similar to the epithelial proliferation log function of direct and indirect stimulation (Sakata; Tamate, 1976) of VFA adsorption on the developing epithelium. Resende Junior et al. (2006b) also detected differences in the MI of rumen papillae of cows subjected to two concentrated feed meal frequencies. However, they detected no differences in the area and papillae length. The only variable for which the authors reported a difference was the area/length relationship, which reflects the width of the papillae. Similarly, in this experiment, we found differences between the ingestion periods in the area-length relationship of papillae in the rumen. As reported by Resende Júnior, et al. (2006b), the inaccuracies in the macroscopic measurement technique may decrease accurate detection of behavior differences in the extension of the rumen’s absorptive surface. However, other authors have reported accurate measurements of the extent of the absorptive surface using the same technique described in the current experiment (Melo et al., 2013).

The average width of the papillae of the rumen of cows, measured as the area/length ratio, increased with increasing dietary energy level. The average papillae width was highest after 18 days of feeding with CS, indicating stimulation to increase the absorptive area of these buds, similar to that reported by Resende Júnior et al. (2006b). They observed increased papillae width on the 12th day of feeding a high-energy diet. After fasting, the buds decreased in width, but not to the S-diet width. During re-feeding, the width of the papillae recovered, but unlike the MI, the increase was not continuous, as was expected, which may be due to measurement inaccuracies, as discussed above.

The thickness of the keratin layer was higher in the ruminal epithelium than in the omasal epithelium (Table 3). However, there was no difference between the two compartments in the total thickness of the epithelium and non-keratinized layers. It is expected that the omasal epithelium has a thinner keratin layer, since its relative digestive surface is much larger than in the rumen (Daniel; Resende Júnior; Cruz, 2006), resulting in increased abrasiveness of the omasum’s absorptive surface and on the mucosa of the blades, resulting in higher keratin peeling and a thinner keratin layer.

The known dehydration of the contents of the omasum in relation to the contents of the rumen (Becker; Marshall; Dixarnold, 1963) may also favor the abrasiveness and the narrow space of the interlaminar recesses as well as omasum blade movement. The similarity between the non-keratinized epithelium layers may indicate that the cell proliferation response can be triggered by stimuli common to both the rumen and the omasum.

![Figure 2: Correlation of rumen and omasum MI in non-pregnant, non-lactating cows. Omasum MI = 0.8705X+0.0281*MI of rumen; r²=0.66; P<0.01.](image)

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The MI of the basal layer of the epithelium of the rumen was higher than the basal layer of the epithelium of the omasum (Table 3). Daniel and Resende Júnior (2012) studied the stomachs of slaughterhouse animals and reported a higher average IM in the omasum than in the rumen. However, the authors did not know and had no control of the diet and the period during which the animals consumed the diet. Therefore, the animals may have been ingesting diets with different energy levels during different times, which could lead to inconsistent values, since the average values were obtained from animals with different eating patterns.

There was a positive correlation between MI of the omasum and rumen (Figure 2). This finding suggests that the factors that stimulate cell proliferation in the omasum are the same as those that stimulate cell proliferation in the rumen. This relationship has also been reported in slaughterhouse animals (Daniel; Resende Júnior, 2012).

The behavior of MI in both compartments during the ingestion periods was as expected, with values relative to the energy level of the diet (Figure 3). The MI in both the rumen and omasum increased after four days of feeding the high-energy diet, both before and after fasting. Goodlad (1981) observed an increase in the basal cell layer of the ruminal epithelium, from 0.58% with a forage-only diet, to 1.20% after the fourth day of a high-energy diet. After this peak MI, we observed a decrease after 18 days of feeding the high-energy diet, similar to a report by Goodlad (1981), who found that, after 62 days of concentrated diet, the IM returned to levels similar to those measured when the animals were fed only forage.

The significant interaction between diet and compartment may indicate that the proliferative response to stimulation differs between the rumen and the omasum; it appears that the omasum responds more quickly to positive and negative stimulation of cell proliferation. Therefore, the proliferative response is quicker in the omasum after introduction of the high-energy diet, and a quick decrease in cell division is seen during fasting. We did not find any literature comparing omasal and ruminal responses to proliferative stimulation of dietary energy that also included a fasting period.

### Table 3: Mitotic index, keratin thickness, total epithelium, and epithelium without keratin in the rumen and omasum.

<table>
<thead>
<tr>
<th>Compartment</th>
<th>Rumen</th>
<th>Omasum</th>
<th>SEM&lt;sup&gt;1&lt;/sup&gt;</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mitotic index (%)</td>
<td>0.55</td>
<td>0.51</td>
<td>0.014</td>
<td>0.02</td>
</tr>
<tr>
<td>Thickness of the keratin layer (µm)</td>
<td>76.43</td>
<td>56.23</td>
<td>4.330</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Thickness of the epithelium (µm)</td>
<td>352.94</td>
<td>342.71</td>
<td>17.431</td>
<td>0.674</td>
</tr>
<tr>
<td>Thickness of the non-keratin layers (µm)</td>
<td>276.51</td>
<td>286.48</td>
<td>16.812</td>
<td>0.67</td>
</tr>
</tbody>
</table>

<sup>1</sup>Standard error of the mean.
CONCLUSIONS

The similarity in TNKL and the positive correlation between MI of the epithelial basal layers of the rumen and omasum indicate that stimulation of cell proliferation triggered by dietary energy content operate simultaneously in both compartments. Compared to the rumen, however, the omasum appears to respond more quickly to both dietary energy proliferative stimulation and fasting.

REFERENCES


