2,6-Diaminopimelic acid (DAPA) in microbial protein quantification of heifers fed different forage sources

Cristovão Colombo de Carvalho Couto Filho¹, Eloisa de Oliveira Simões Saliba², Norberto Mario Rodriguez², Geraldo Sérgio Senra Carneiro Barbosa³, Regane Martins de Freitas², Matheus Pinheiro Diniz Resende²

¹ Universidade Federal de Minas Gerais, Escola de Veterinária, Programa de Pós-graduação em Zootecnia, Belo Horizonte, MG, Brasil. 
² Universidade Federal de Minas Gerais, Escola de Veterinária, Departamento de Zootecnia, Belo Horizonte, MG, Brasil. 
³ Universidade Federal de Viçosa, Departamento de Zootecnia, Campus Florestal, Florestal, MG, Brasil.

ABSTRACT - The objective of this study was to evaluate the flow of nitrogenous compounds, protein degradability, rumen degradability of total carbohydrate and organic matter and microbial efficiency in heifers subjected to diets containing corn silage, sugarcane or Tifton. For this purpose, the 2,6-diaminopimelic acid (DAPA) technique was adopted and analytical procedures for amino acids by HPLC were adapted. Six rumen-fistulated Holstein-Zebu heifers with 480 kg of initial BW and at 24 months of age kept in individual tie stalls were assigned to two 3 × 3 Latin squares. Omasal digesta dry matter and microbial dry matter flows were determined using the isolated, purified and enriched lignin (LIPE®) and DAPA markers, respectively. Isolated bacteria from rumen showed on average 5.84 g/100 g microbial N, 0.25 g/100 g DAPA in dry matter and 44.61 DAPA: N ratio. The forage sources did not influence the flows of nitrogen compounds, except for total omasal flow and non-ammonia N in relation to N intake for the corn silage diet, for which there was an upward trend compared with the other diets. The degradation of the organic matter and total carbohydrates did not differ, averaging 6.1 kg/day and 5.2 kg/day, respectively. The studied forage sources do not influence the flows of nitrogen compounds, except for total omasal flow and non-ammonia N in relation N intake for the corn silage diet, for which there is an upward trend compared with the other diets. Protein degradability and microbial efficiency are similar between evaluated diets.

Key Words: corn silage, flow of nitrogen compounds, HPLC, microbial efficiency, protein degradability, sugarcane

Introduction

The protein requirements of ruminants are met through the absorption of amino acids in the intestine, and 60% of these amino acids come from microbial protein synthesized from the symbiosis between animal and ruminal microbiota. The microbial protein, whose intestinal digestibility is close to the ideal, regardless of the protein composition ingested by the animal.

Because of the importance of microbial protein to protein metabolism of ruminants, the quantification of its flow under different physiological and dietary conditions is essential for the requirements of amino acids to be met.

In this study we proposed to evaluate different supplementary forages regularly used aiming to minimize forage seasonality in Central Brazil in order to verify a possible difference in protein synthesis between sources. The sugarcane (Saccharum officinarum, L.) was used because of its peak production and nutritive value coinciding with the period of scarcity of forage. Of the conserved fodder, those selected were corn silage, due to its high content of soluble carbohydrates, which provide a good fermentation pattern, thereby placing it on the level of ideal grass and most used in ensiling processes, and Tifton hay, whose morphophysiological characteristics allow for a more uniform drying, generating a hay that maintains the quality and nutritional value of the fresh forage.

Several microbial markers were used for the abovementioned type of evaluation, each with its advantages and limitations. A noteworthy marker is the 2,6-diaminopimelic acid (DAPA), an amino acid present in the bacterial cell wall, suggested as microbial marker by Synge in 1953, and widely used in experiments in the past for this purpose (Broderick and Merchen, 1992). However, its laboratory quantification by classic chromatography demanded extensive labor time, thus falling into disuse. With the evolution of the technique came the high-performance liquid chromatography (HPLC), recognized by rapid analysis, automation, precision and sensitivity.

Received May 2, 2013 and accepted June 23, 2014.
Corresponding author: cristovao.couto@ifma.edu.br
*Current address: Instituto Federal do Maranhão, São Luís, MA, Brasil.
http://dx.doi.org/10.1590/S1806-92902015000500003
Copyright © 2015 Sociedade Brasileira de Zootecnia. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License, which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.
Therefore, this study was undertaken to assess the flow of nitrogen compounds, protein degradability, rumen degradation of total carbohydrates and organic matter and microbial efficiency in heifers subjected to diets containing corn silage, sugarcane or Tifton hay. For this purpose, DAPA technique was used and analytical procedures for amino acids by HPLC were adapted. The technique used for microbial quantification was not the focus of this study, but only a means to achieve the objective.

Material and Methods

Six rumen-cannulated crossbred Holstein-Zebu heifers with 480 kg of initial live weight and at 24 months of age were kept in individual tie stalls sheltered and equipped with individual water troughs and troughs for the feed supply.

The experiment lasted 57 days, divided into 3 periods of 19 days. The animals were weighed, identified and dewormed at the beginning of the experiment and kept for 13 days in a period of adaptation to experimental diets and facilities.

Three treatments consisting of different forages in diets were evaluated: corn silage (*Zea mays* L.), Tifton hay (*Cynodon spp.*) and sugarcane (*Saccharum officinarum*, L.). The diets were supplemented with concentrate based on ground corn grain, soybean meal, mineral and vitamin premix, and formulated according to NRC (2001) to be isonitrogenous (14% CP), with a forage:concentrate ratio of 75:25, on a dry matter basis (Table 1).

The dietary ingredients were weighed and mixed manually twice daily to be delivered at 8.00 h and 14.00 h daily as total mixed ration, in an amount sufficient to allow for 5 to 10% of the offered as daily orts.

The dry matter intake was calculated by multiplying the daily intake of fresh matter of each feed, between days 14 and 19 of each period, by their respective dry matter content, followed by subtracting the daily orts of dry matter. The daily nutrient intake per animal was calculated by multiplying the offered dry matter of each ingredient by its respective nutritional content. Daily orts from the same nutrient were subtracted from the total of nutrients offered per animal.

During the sampling period, samples of forages, concentrates and orts from each animal were collected daily, placed in plastic bags and stored at −20 °C. At the end of each period, composite samples were made for each feed and also for the orts from the mixture of equal amounts of fresh matter. In the case of the orts, composite samples were made for each heifer per period.

To quantify omasal flow, the external marker LIPE® was supplied in capsule form at a dosage of 500 mg per heifer daily with two days for adaptation plus the provision during the entire sampling period. To collect samples of omasal digesta a set of devices consisting of kitassato, collecting tube and a vacuum pump was used, according to the technique described by Leão (2002). The omasal digesta collections were performed between the days 14 and 16 of each period, according to the following protocol: on the first day collections were made prior to the morning feeding (0 h) and 6 hours after; on the second day, at the 2 and 8 hours and, on the third day, 4 and 10 hours after it. Omasal digesta samples were frozen at −20 °C immediately after collection and kept this way until processing. Composite samples were made for each animal and experimental period. The samples of omasal digesta were dried in a forced air oven at 55 °C for 96 hours and processed in a mill with a 1 mm mesh sieve. To estimate the flow of the omasal digesta, LIPE® was used as a marker. The analysis of the LIPE® marker was performed in a spectrophotometer with light detector in the spectrum of infrared (Saliba et al., 2005). The flows of dry matter and other nutrients in omasal digesta samples were estimated according to the equation of France and Siddons (1986) for external markers:

\[
\text{Omasal flow} = \frac{\text{Marker concentration (mg/g digesta)}}{\text{Marker dosage mg/day}}
\]

On the 19th day of each period, ruminal digesta was collected four hours after feeding for the isolation of bacteria according to technique described by Cecava et al. (1990). The ruminal content was then centrifuged at 170 x g for 5 minutes at 4 °C for the separation of solids. The precipitate was

### Table 1 - Ingredients and nutrient composition of the experimental diets

<table>
<thead>
<tr>
<th>Item</th>
<th>Corn silage</th>
<th>Sugarcane</th>
<th>Tifton hay</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nutrients</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crude protein</td>
<td>139</td>
<td>135</td>
<td>138</td>
</tr>
<tr>
<td>Neutral detergent fiber</td>
<td>513</td>
<td>508</td>
<td>618</td>
</tr>
<tr>
<td>Ash</td>
<td>52</td>
<td>54</td>
<td>64</td>
</tr>
<tr>
<td>Ether extract</td>
<td>27</td>
<td>11</td>
<td>23</td>
</tr>
<tr>
<td>Non-fibrous carbohydrates</td>
<td>269</td>
<td>292</td>
<td>157</td>
</tr>
<tr>
<td>Total digestible nutrients</td>
<td>706</td>
<td>683</td>
<td>680</td>
</tr>
<tr>
<td><strong>Ingredients</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corn silage</td>
<td>741</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Sugarcane</td>
<td>-</td>
<td>738</td>
<td>-</td>
</tr>
<tr>
<td>Tifton hay</td>
<td>-</td>
<td>-</td>
<td>745</td>
</tr>
<tr>
<td>Ground corn grain</td>
<td>163</td>
<td>-</td>
<td>215</td>
</tr>
<tr>
<td>Soybean meal</td>
<td>89</td>
<td>255</td>
<td>33</td>
</tr>
<tr>
<td>Minerals and vitamins&lt;sup&gt;1&lt;/sup&gt;</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
</tbody>
</table>

<sup>1</sup> Minerals and vitamins (per kg): 150 g of Ca; 90 g of P; 17.78 g of Mg; 15.40 g of S; 114 g of Na; 1,500 mg of Mn; 4,000 mg of Zn; 1,200 mg of Cu; 145 mg of Co; 90 mg of I; 38 mg of Se; 900 mg of F; 200,000 IU of vit. A; 50,000 IU of vit. D; 1,500 mg of vit. E.
discarded and the supernatant was subjected to a second centrifugation at 17,000 x g for 15 minutes at 4 °C, for precipitating the bacteria. Immediately after, the supernatant was removed and the bacteria were resuspended with 1M NaCl. A third centrifugation at 17,000 x g for 15 minutes at 4 °C was performed. The precipitate was then collected with a spatula and washed with acetone. Finally, the precipitate was subjected to oven-drying at a temperature of 45 °C and subsequently ground in a mortar pestle to turn the material into powder form.

The flow of microbial nitrogen (mic N) in omasal digesta was obtained from the ratio between the amount of DAPA in omasal digesta and DAPA:N on the microorganisms isolated. The microbial efficiency was expressed in different ways: g microbial nitrogen/kg of rumen-degradable organic matter (g mic N/kg RDOM), g microbial nitrogen/kg of rumen-degradable total carbohydrates (g mic N/kg RDTC) and g microbial crude protein/kg total digestible nutrients intake (g mic CP/kg TDN).

The separation and quantification of 2,6-diaminopimelic acid was carried out in a liquid chromatograph (HPLC) model Shimadzu LC-10AD (Shimadzu Corporation, Kyoto, Japan) equipped with an ion-exchange column and a fluorescence detector (equipment specifications: communication module CBM - 10A, LC-10AD pumps, injector: SIL-10A, oven, CTO-10A and detector: RF-535).

The analyses of DAPA on the bacterial pellet samples and on the omasal digesta were performed using ion-exchange chromatography system with post-column derivatization, using o-phthalaldehyde (OPA) and fluorimetric detection. The standard utilized contained 5.0 µg/L of DL-2,6-diaminopimelic (Fluka AG Buchs SG, Switzerland) in sodium citrate buffer pH 2.2. The software CLASS-LC10 - LC Workstation allowed access to the communication module CBM - 10A, LC-10AD pumps, injector: SIL-10A, oven, CTO-10A and detector: RF-535).

The analyses of DAPA on the bacterial pellet samples and on the omasal digesta were performed using ion-exchange chromatography system with post-column derivatization, using o-phthalaldehyde (OPA) and fluorimetric detection. The standard utilized contained 5.0 µg/L of DL-2,6-diaminopimelic acid (Fluka AG Buchs SG, Switzerland) in sodium citrate buffer pH 2.2. The software CLASS-LC10 - LC Workstation allowed access to the chromatograms obtained after analysis of the standard and of the samples and also the determination of the areas of peaks related to the DAPA. The final concentration of DAPA on the samples was obtained from a simple ratio between the area of the standard and area of the sample, as can be seen in the following expression:

\[
\text{Sample concentration} = \frac{\text{Standard concentration} \times \text{Area of the sample}}{\text{Area of the standard}}
\]

The analytical conditions in the system were: Column Shim-pack Amino-Na (4.6 x 100 mm), Trap Shim-pack ISC-30/S0504 Na, mobile phases A - Citrate buffer pH 3.2, B - Citrate buffer pH and 10 C - NaOH (0.2 N), gradient elution flow 0.4 ml/min, temperature 60 °C and detection RF-20Axs Ex 350 nm, 450 nm.

For the hydrolysis procedure, 20 mg of crude protein of the bacteria samples and omasum digesta samples were weighed in 30 ml test tubes with screw caps. Then, 10 mL of 6N HCl were added to the samples. The tubes were sealed off and dried in the oven, where they remained for a period of 22 hours at a temperature of 110±5°C (Bernardi, 2000). After this period, the samples were removed from the oven and left at room temperature until cooling. They were subsequently filtered through a qualitative filter paper and subjected to evaporation. After this procedure, they were resuspended in citrate buffer at pH 2.2 in a 10 mL volumetric flask, completing the volume. One milliliter of this sample was pipetted into a graduated test tube, and citrate buffer was added to a level high enough to immerse the electrode of the potentiometer, and the pH of the sample was adjusted to 2.2, using solutions of 4N NaOH and 0.1N HCl for this purpose. After this adjustment, the sample was transferred to a 10 mL volumetric flask, and its volume was completed. Finally, the sample was subjected to microfiltration using Millex filters type 25 mm HV 0.45 µM and then transferred to the standard vials used in the automatic injector of the liquid chromatograph.

The samples of food, orts and omasal digesta were pre-dried at 55 °C, ground in a Thomas-Wiley mill with a 1 mm mesh sieve and stored in plastic containers. The analyses of dry matter (DM), mineral matter (MM), crude protein (CP) and ether extract (EE) contents were performed according to recommendations of the Association of Official Analytical Chemists (AOAC, 1995). The contents of neutral detergent fiber (NDF) were determined according to Van Soest et al. (1991).

The percentages of non-fibrous carbohydrates (NFC) and total carbohydrates (TC) were obtained using the equations proposed by Sniffen et al. (1992):

\[
\text{NFC} = 100 - (% \text{NDF} + % \text{CP} + % \text{EE} + % \text{Ash})
\]

\[
\text{TC} = 100 - (% \text{CP} + % \text{EE} + % \text{Ash})
\]

The content of total digestible nutrients (TDN) of diets were calculated according to the equation described by the NRC (2001), as follows:

\[
\%\text{TDN} = \% \text{digestible CP} + \% \text{digestible NDF} + \% \text{digestible NFC} + 2.25\% \times \% \text{digestible EE}
\]

Regarding the samples of bacterial pellets, not including the 2,6-diaminopimelic acid, they also underwent analysis of DM and CP according to the procedures described above.

For the study of the flows of nitrogen compounds, protein degradability, rumen degradation of total carbohydrates and organic matter and microbial efficiency through the 2,6-diaminopimelic acid (DAPA), two 3 x 3 Latin squares were used in the experiment, according to the following model:

\[
Y_{ijkl} = \mu + Q_{i} + D_{j} + P_{k} + N(i)l + e_{ijkl}
\]

in which: \( \mu \) = overall mean; \( Q_{i} \) = effect of Latin square i; \( D_{j} \) = effect of diet j; \( P_{k} \) = effect of period k; \( N(i)l \) = effect...
of heifer l, nested in Latin square i; and eijkl = independent experimental error and with normal distribution, zero mean and $\sigma^2$ variance. LIPE was adopted as the marker of the flow of digesta, due to the good results reported by Couto Filho et al. (2012).

For statistical analysis, all data obtained was subjected to analysis of variance and the means of the diets were compared by SNK test at 5% probability, on the SISVAR statistical software (Ferreira, 2000).

**Results and Discussion**

The microbial N contents of 6.01, 5.55 and 5.97 g/100 g DM for corn silage, sugarcane and Tifton hay (Table 2), respectively, are within the range of 4.83 to 10.58 g/100 g DM reported by Clark et al. (1992) from the compilation of literature data. Another compilation made by Valadares Filho (1995) with results of studies conducted in Brazil showed variations from 81.1 to 95.7 g/100 g as fed for dry matter and 5.2 to 8.7 g/100 g DM for total nitrogen compounds, which are also consistent with the values obtained in this study, mainly because these values were obtained in tropical conditions.

In another study, Silva et al. (2007) evaluated diets with Tifton hay and different protein sources for cattle, reporting mean values ranging from 5.14 to 5.55 g/100 g DM.

The concentrations of DAPA are very similar for the different forage sources (Table 2). The overall mean percentage of 0.25% DAPA in bacterial DM and 44.61 for the DAPA:N ratio is within the range of 0.14 to 0.44% reported by Clark et al. (1992), and very close to the average 0.31% cited by the same authors. In an experiment evaluating different techniques for quantification of microbial synthesis in different breeds of cattle, Valadares Filho et al. (1990) also reported an average value of 0.3% of DAPA in DM, DAPA:N ratio of 49 mg/g of N in ruminal bacteria, and microbial N values very similar to those reported here.

For Clark et al. (1992), the origin and amount of crude protein in the diet and the roughage:concentrate ratio of the diet significantly alter this relationship. According to Craig et al. (1987) and Cecava et al. (1990), this relationship varies in the post-feeding period of high-concentrate diets. However, they do not vary when diets with a high roughage proportion are used (Dufva et al., 1982; Olubobokun et al., 1988), a fact that was also observed in the present study.

For Nocek and Russell (1988), the chemical composition of isolated bacteria may be altered according to the conditions of the limiting environment. These authors suggested that the estimations of microbial yield were expressed based on the bacterial N content, instead of the dry weight of cells. According Clark et al. (1992), the variations in composition of the bacteria can be attributed to the different techniques used to isolate and determine such composition and also to the very variations in their composition.

Ruminal fermentation and flow of microbial and dietary protein to the intestine are affected by feed intake and by the amount and source of energy and protein in the diet. Dietary protein and carbohydrates that are not degraded in the rumen increase the amount of dietary protein that passes to the intestine and may decrease the amount of microbial protein that is synthesized in the rumen, thus leading to variations in the microbial composition.

In the present study there was a trend towards superiority of the silage diet on the omasal flows of total N (P = 0.08) and non-ammonia nitrogen (NAN; P = 0.08) in relation to N intake (Table 3). These results can be explained by the differences in the chemical composition (Table 1) of the diets, since the diet of corn silage has a low NDF content, as well as a high content of non-fibrous carbohydrates. Therefore, these factors may have contributed to the increase in these flows to the omasum. These results can also be explained by the lack of synchronism in the rumen degradation of protein and carbohydrates. In the case of the hay, the fiber fraction represents the main source of energy in the rumen and it has slow degradation. As for the sugarcane diet, the lack of synchrony may be justified by the absence of non-protein nitrogen, whose source is important when this type of roughage is used in a diet; however, it was not used in this study as we sought to standardize the protein source used in the different treatments.

Martins et al. (2006), in a comparative study between corn silage and Tifton hay, observed differences in the evaluated parameters. The levels of rumen-degradable organic matter (RDOM), as well as total N, NAN and microbial N flows were higher in the diets containing corn silage. Several factors can influence the flow of microbial protein, mainly the content and solubility of dietary protein, the N sources of endogenous origin, the amount of

<table>
<thead>
<tr>
<th>Item</th>
<th>Diets</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CS</td>
<td>SC</td>
</tr>
<tr>
<td>Dry matter (g/100 g as fed)</td>
<td>85.63</td>
<td>80.60</td>
</tr>
<tr>
<td>Microbial N (g/100 g DM)</td>
<td>6.01</td>
<td>5.55</td>
</tr>
<tr>
<td>DAPA (g/100 g DM)</td>
<td>0.26</td>
<td>0.26</td>
</tr>
<tr>
<td>DAPA (mg/g DM)</td>
<td>2.57</td>
<td>2.56</td>
</tr>
<tr>
<td>DAPA:N (mg/g mic N)</td>
<td>47.15</td>
<td>46.23</td>
</tr>
</tbody>
</table>

DAPA - 2,6-diaminopimelic acid.

Table 2 - Chemical composition of the bacteria isolated from the rumen of heifers fed diets based on corn silage (CS), sugarcane (SC) and Tifton hay (TH).
digestible organic matter in the diet, the treatment to which dietary protein was subjected, and the absorption of N by the rumen, particularly in the form of ammonia (Hoover and Stokes, 1991).

The flow of mic N represented, on average, 70.33 g/100 g of omasal flow of NAN and 69.85 g/100 g of total N flow in the different treatments. No effect was observed also on the degradability of the diets (Table 3). Similar results were reported by Berchielli et al. (1995b), who studied diets with different roughage:concentrate ratios and, among them, the 80:20, similar to that used in this study (75:25). According to Clark et al. (1992), a reduction in the flow of microbial protein to the small intestine may occur when diets with a high roughage inclusion are used, which may be attributed to the deficiency of readily available energy and to the increase of nitrogen-compounds recycling by rumen microorganisms. The average value of mic N, expressed compared with the NAN in the omasum of the present study, was close to the 75.57 and 73.25 g/100 g reported by Rode et al. (1985) and Klusmeyer et al. (1990), respectively.

The omasal flow of microbial N of the diets containing hay averaged 75.66 g/day (Table 3), which is lower than the 100.6 g/day obtained by Guimarães et al. (2001) in a study with a roughage:concentrate ratio of 70:30, consisting of Tifton hay and cassava waste. This difference may be related to the starch source in the diet. According to Zeoula et al. (2002), the microbial N flow may increase when corn is replaced by a starch source that is more degradable in the rumen, such as cassava.

The quantities of RDOM and RDTC were not influenced by forage sources, averaging 6.1 and 5.2 kg/day, respectively (Table 4). Berchielli et al. (1995a) also found no effect of treatments on the amount of RDOM and RDTC. No effect of the experimental diets was observed on microbial efficiency either, when it was expressed as g mic N/kg RDOM, g mic N/kg RDTC and g mic CP/kg TDN, whose overall mean values were 16.51, 18.83 and 80.70, respectively (Table 4).

In the CNCPS system (Fox et al., 2004), microbial yield is estimated at 40 g of cells per kg of rumen-degradable total carbohydrates, a value obtained from the maintenance requirements of microorganisms and their respective growth rates (Russell et al., 1992). These values are satisfactorily accurate for higher dry matter omasal flows, which are a result of nutrition plans superior to those of the present study.

<p>| Table 3 - Omasal flow of nitrogen compounds (g/day) and degradability of protein in diets based on corn silage (CS), sugarcane (SC) and Tifton hay (TH) |</p>
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Diets</th>
<th>SEM</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total N</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flows</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>g/day</td>
<td>139.24</td>
<td>128.21</td>
<td>114.71</td>
</tr>
<tr>
<td>g/100 g N intake</td>
<td>64.28</td>
<td>46.47</td>
<td>53.97</td>
</tr>
<tr>
<td>Non-ammonia nitrogen</td>
<td>138.46</td>
<td>127.22</td>
<td>113.98</td>
</tr>
<tr>
<td>g/day</td>
<td>63.92</td>
<td>46.11</td>
<td>53.65</td>
</tr>
<tr>
<td>g/100 g N intake</td>
<td>87.20</td>
<td>99.78</td>
<td>75.66</td>
</tr>
<tr>
<td>Microbial N</td>
<td>65.24</td>
<td>80.40</td>
<td>65.36</td>
</tr>
<tr>
<td>g/day</td>
<td>64.88</td>
<td>79.73</td>
<td>64.95</td>
</tr>
<tr>
<td>RDP (g/100 g)</td>
<td>74.45</td>
<td>90.25</td>
<td>82.74</td>
</tr>
<tr>
<td>RUDP (g/100 g)</td>
<td>25.55</td>
<td>9.75</td>
<td>17.26</td>
</tr>
</tbody>
</table>

SEM - standard error of the mean; P-value - probability value for the effect of diet; NAN - non-ammonia nitrogen; RDP - rumen-degradable protein; RUDP - rumen-undegradable protein.

<p>| Table 4 - Rumen-degradable organic matter and total carbohydrates (RDOM and RDTC) and microbial efficiency in heifers fed diets based on corn silage (CS), sugarcane (SC) and Tifton hay (TH) |</p>
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Diets</th>
<th>SEM</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDOM (kg)</td>
<td>6.67</td>
<td>6.01</td>
<td>5.58</td>
</tr>
<tr>
<td>RDTC (kg)</td>
<td>5.90</td>
<td>4.90</td>
<td>4.85</td>
</tr>
<tr>
<td>g mic N/kg RDOM</td>
<td>13.62</td>
<td>17.35</td>
<td>18.55</td>
</tr>
<tr>
<td>g mic N/kg RDTC</td>
<td>15.22</td>
<td>21.36</td>
<td>19.92</td>
</tr>
<tr>
<td>g mic CP/kg TDN</td>
<td>72.25</td>
<td>90.32</td>
<td>79.49</td>
</tr>
</tbody>
</table>

SEM - standard error of the mean; P-value - probability value for the effect of diet.
In the case of low levels of intake, overestimated values of the microbial N flow are predicted. The mean values found for the efficiency of microbial synthesis in g mic CP/kg TDN were lower than the 130 recommended by NRC (2001), irrespective of diet. For Dewhurst et al. (2000) and NRC (2001), the availability of energy in the rumen and N values are the factors that most limit microbial growth. According to these authors, to enhance fermentation and microbial protein synthesis in the rumen, it is necessary to synchronize the availability of fermentable energy and rumen-degradable N.

The estimated values of microbial efficiency expressed in relation to RDOM and RDTC for the hay diet were greater than those obtained by Silva et al. (2007), working with Tifton hay supplemented with soybean meal, among the different nitrogen sources evaluated, but lower than the results reported by Berchielli et al. (1995a), who evaluated different markers, including DAPA, and different roughage:concentrate ratios, and Cabral et al. (2008), who evaluated different forages. These same authors, utilizing corn silage, reported slightly greater efficiencies than the results obtained in the current study.

Taking into account the ruminal availability of energy and nitrogen as the determinant factors of microbial growth, high roughage proportions such as that used herein (75:25) can lead to reduction of energy availability and increase in the recycling of nitrogen compounds by rumen microorganisms. Thus, the decrease in the passage of microbial protein to the small intestine may have contributed to the lower values of microbial protein synthesis efficiency in this experiment. Therefore, the high roughage:concentrate ratio used in this study may have influenced the microbial growth by limiting it. Furthermore, the similarity between the roughage:concentrate ratio (75:25) of all experimental diets may have resulted in the absence of significant effect on the microbial efficiency among the many roughage sources used in this study. For Clark et al. (1992), faster rates of microbial growth are attributed to the greater availability of nutrients, associated with a faster passage of microorganisms to the abomasum, leading to the reduction of maintenance requirement of rumen macrobiotic and, consequently, an increased microbial efficiency.

An important criticism about the studies with microbial protein synthesis is regarding the calculations of the microbial compounds flows, taking into account the percentage of nitrogen in relation to organic matter rather than dry weight. Obviously, potential contamination of bacteria with saline solution during the isolation procedures can occur, and when that happens the N content is normally used in relation to organic matter. Therefore, this emphasizes the need for standardizing these calculations, because as they are based on the dry matter, they provide significantly different results for both the nitrogen-compound flow and the microbial efficiencies as compared with those derived from organic matter. The consequence of this is the difficulty in comparing results of different studies.

Conclusions

The studied forage sources do not influence the flows of nitrogen compounds, except for total omasal flow and non-ammonia nitrogen in relation to nitrogen intake for the corn silage diet, for which there is an upward trend compared with the other diets.

The protein degradability, the degradation of organic matter and total carbohydrates in the rumen, and the efficiency of microbial synthesis are similar among the evaluated diets.

Acknowledgments

The authors would like to thank Fundação de Amparo à Pesquisa do Estado de Minas Gerais (FAPEMIG) for the financial support to this research, and Fundação de Amparo à Pesquisa e ao Desenvolvimento Científico e Tecnológico do Maranhão (FAPEMA) for the partial support to the publication of this research.

References


