Quantification of the Efficiency of Rumen Microbial Protein Synthesis in Steers Fed Green Tropical Grass

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ABSTRACT


Tingkat pasokan protein mikroba rumen (MCP) ke usus halus merupakan salah satu unsur kunci dalam mengestimasi respon pertumbuhan ruminan terhadap ransum tertentu. Data MCP hijauan tropis selalu berada di bawah nilai prediksi model rumen yang dipakai saat ini. Dengan demikian, kuantifikasi pasokan MCP rumput tropis diharapkan meningkatkan kemampuan prediksi model rumen untuk pakan daerah tropis. Data MCP pangola (Digitaria erianthe cv. Steudal) dipanen setiap pagi dan langsung diberikan kepada ternak dalam kandang metabolis. Parameter yang diukur adalah produksi MCP dan efisiensi sintesis MCP (EMPS), konsumsi, laju alir digesta, dan konsentrasi ammonia rumen. Kandungan protein kasar dan karbohidrat mudah larut adalah 6,3 dan 7,4% dari DM, sedangkan laju alir digesta cair sebesar 7,84 %/jam dan padat sebesar 6,92%/jam. EMPS rata-rata hanya 72 g MCP/kg bahan organic tercerna. Dikeluarkan bahwa nilai EMPS untuk ruminant tropis yang dipakai untuk hijauan dalam kondisi seperti pada penelitian ini berada jauh di bawah nilai yang dipakai untuk ruminant tropis pada bahan organic tercena.

Kata kunci: Protein Mikroba, Efisiensi Sintesis, Rumput Tropis, Sapi

ABSTRACT


The rate of rumen microbial crude protein (MCP) supply to the intestines is a crucial element in the current rumen models to predict respond of ruminants to a certain diet. Data from tropical pastures are always below predicted results from the existing rumen models. Thus, quantification of the rumen MCP supply from tropical forage will improve predictive rate under tropical feeding conditions. Four Brahman crossbred steers (457 ± 20.1 kg) were used in a metabolism study. Pangola grass (Digitaria erianthe cv. Steudal) was harvested every morning and fed to the animals soon afterward. Parameters measured were EMPS, intake, fractional passage rates, and rumen ammonia concentration. The EMPS was estimated using purine derivative excretion in urine. Mean crude protein and water soluble carbohydrate was 6.3 and 7.4% of dry matter (DM) respectively. Mean DM intake was 1.6% liveweight. Average rumen ammonia concentration was 69 mg/L whilst rumen passage rates were 7.84 and 6.92%/h for fluid and solids respectively. Mean EMPS was only 72 g MCP/kg digestible organic matter. It is concluded that EMPS in steers consuming green pangola grass was at the level below the minimum recommended value for forage diets adopted in the current feeding standards.

Key words: Microbial Protein, Efficiency, Tropical Grass, Cattle

INTRODUCTION

One vital factor in the current rumen models to predict ruminant’s respond to a certain feeding regime is microbial crude protein production (MCP). The amount of MCP available for ruminants is dictated by the efficiency of its synthesis (EMPS) in the rumen. The EMPS is affected by many factors, such as diet quality, level of intake and rumen dilution rate, and varies considerably across diets. The EMPS values for tropical grass hay reported in literatures ranged from 33 to 117 g MCP/kg DM (McMeniman et al., 1986; Poppi et al., 1997; Prior et al., 1998; Bowen, 2003; Mullik, 2006) which is lower than the values adopted in the current feeding standards (SCA, 2007; AFRC, 1992; NRC, 2000). Although a higher efficiency value from high quality pangola grass (176 g MCP/kg DOM) was reported by Mullik (1999) but the value might not
accurate since pasture intake and digestibility were indirectly measured. There are also methodological problems in measuring urinary purine output through spot sample technique as used by MULLIK (1999) since assuming that a constant value in creatinine: purine derivative ratio across diets and animals is still debatable. The present experiment was designed to measure the E_MPS for fertilized tropical grass (pangola) during the wet season and managed to provide high amounts of green leaf, as well as nutrient content.

MATERIALS AND METHODS

Experimental animals

Four Brahman crossbred steers (457 ± 20.1 kg) were used in this study. The steers were vaccinated and drenched against internal and external parasites prior to the commencement of the study. They were held in feedlot pens and housed in metabolism crates on site for the duration of the study.

Experimental design, diet and treatment

There was 1 treatment with 4 replications (steers) to estimate the parameter, efficiency of MCP production, and compare it to the feeding standards. The steers were randomly allocated into metabolic crates. There was a two week preliminary and one week data collection period. Diet was freshly cut pangola grass. The grass was harvested daily and fed at 10% above voluntary intake determined in the last week of the adaptation period, and offered in 3 periods daily. Drinking water and a mineral block were freely available at all times. Approximately 0.5 ha permanent pangola grass pasture was used to provide feed for the steers. The paddock was slashed, approximately 8 cm above ground, and fertilized with 320 kg diammonium phosphate/ha (18% N and 20% Phosphorus) and 130 kg urea/ha 6 weeks before the study commenced. Approximately 17 mm irrigation was applied after slashing and fertilizing. There was no more irrigation because of an adequate rainfall throughout the study (88.3 mm).

The steers were held in individual concrete pens (feedlot pens) during the first 11 days of the preliminary period and were moved onto metabolic crates on day 12. The steers were given 3 days to adapt to the metabolic crates before data collection began.

Experimental procedures

Feed intake

The freshly cut pangola grass was offered at 10% above voluntary intake, based on the intake during the last week in the preliminary period, three times daily at 08.00, 13.00 and 19.00 h. The morning portion was given soon after cutting and two other portions were spread on a large plastic sheet in a cool room at 4°C and fed at 14.00 and 19.00 h. Two samples were taken at morning feeding. One sample was weighed into a plastic bag, sealed and frozen. Another sample was dried in the oven at 55°C for dry matter (DM) and bulked at the end of the collection period. The same procedures applied for the refusal but daily refusals were taken and processed separately between animals. One sample of forage was also taken at each time of feeding (afternoon and evening), weighed and frozen. At the end of the collection period, frozen samples of feed offered at each time and feed refused were bulked within the sample times (without thawing), mixed and one sub-sample was taken, weighed, freeze dried, ground through 1 mm screen and stored for analysis of organic matter (OM), crude protein (CP), water soluble carbohydrates (WSC), and neutral detergent fibre (NDF).

Digestibility

Digestibility of DM, OM, CP, and NDF was calculated from intake and faecal data. Daily faecal output was measured by total collection into individual buckets placed under metabolism crates. The collection was done for 7 days in each treatment period. Each faeces from a 24 h collection was homogenised, and approximately 5% of faeces produced by each animal was taken and bulked individually in plastic containers in a freezer. At the end of the collection period, the bulked samples were thawed at room temperature and 2 sub-samples were taken from each animal. One sub-sample was dried in an oven at 60°C until constant weight (7 days) to obtain DM content, and discarded. Another sub-sample was frozen followed by freeze drying, and grinding prior to Nitrogen, OM and NDF analysis.

Passage rates

Passage rates were estimated during the period in which digestibilities were measured. Fluid and particulate passage rate from the rumen were estimated using chromium-ethylenediamine tetraacetic acid (Cr-EDTA; 2 g Cr/animal) and Ytterbium trichloride hexahydrate (YbCl₃·6H₂O; 1 g Yb/animal) as external markers. A single dose of markers were done at Day 1 of the collection period. Dosing was done a few minutes prior to morning feeding. A faecal sample from each animal was taken before dosing to serve as a blank or base line in marker analysis and calculation. Subsequent faecal sampling (freshly voided faeces) was taken approximately at the following times: 12, 24, 32, 48, 56, 72, 80, 96, 104, 120, 132, and 144 h post dosing. The
samples were oven-dried at 65°C, ground through 1 mm screen, and stored at room temperature prior to processing for marker analysis. The fractional and fluid passage rates were calculated from the slope of natural log of marker concentration against time. Only samples taken from 12 h to 84 h were used in the regression as they did not deviate from linearity determined by visual observation.

**Rumen ammonia-nitrogen concentration**

Two rumen fluid samples, collected on different occasions, were taken from each animal on the last day of the collection period. The first collection was done 3 to 4 h after morning feeding and the second sample was collected before morning feeding the next day (24 h after feeding).

**Urine sampling for predicting microbial protein synthesis**

The MCP production was estimated by reference to purine derivative (PD) i.e allantoin, uric acid, xanthine and hypoxanthine excretion in total urine, and creatinine (Ct) excretion was also measured. Daily urine output of individual animals was measured by total collection into trays covered with a cloth filter to inhibit faecal contamination. The urine pH was kept below 3 by adding approximately 200 mL 10% H2SO4 into individual trays prior to collection. Urine collected over 24 h was mixed and 5% was taken, bulked into a plastic container in a refrigerator over the collection period. Immediately at the end of each treatment period, 5 mL of the acidified sample was measured into a red cap plastic tube containing 1 mL allopurinol (internal standard). The solution was made up to 50 mL using 0.1M NH4H2PO4 buffer. This solution was then transferred into a clean labelled plastic container and frozen prior to analysis for Ct and PD.

**Analytical procedures**

Analytical procedures for DM, OM, CP, NDF, using the method of VAN SOEST and WINE (1967). Ammonia concentration was determined by distillation technique. Concentration of Purine derivatives and Creatinine was analysed using High pressure Liquid chromatography based on the method proposed by BALLCELL et al. (1991). Concentration of WSC was determined by cold water extraction method (THOMAS, 1977).

Concentrations of Cr and Yb in faecal samples were determined using the digestion method. Approximately 0.3 to 0.4 g dried ground sample was transferred into 50 mL individual erlenmeyer flasks. A 15 mL solution of 5:1 nitric: perchloric acid was added and left to stand for 24 h. After standing, the flasks were placed on a preheated frypan (150°C) and were allowed to digest at this temperature until all brown smoke was dissipated. The temperature was then increased to 300°C and the samples were digested at this temperature for 1 h and followed by digestion at 400°C for about 20 min. The flasks were removed and cooled. The residues were then transferred into 25 mL volumetric flasks and diluted to the mark using distilled water and marker concentration was determined using an ICP (Inductively Coupling Plasma Emission Spectrometer, M+P, Spectro Analytical).

**Calculations**

Microbial protein production was estimated from the excretion of PD in the urine based on formula of CHEN and GOMEZ (1995) as described in MULLIK (2006). Fractional passage rate was calculated by regressing the natural log of marker concentration in faecal samples against time and determining the slope which is the fractional passage rate (GROVUM and WILLIAMS, 1973).

**Statistical methods**

There was no statistical analysis as there were no treatments to compare. Rather standard deviation from the mean was calculated and results were compared to the literature. In particular the efficiency of MCP synthesis was compared to that adopted in the SCA (2007).

**RESULTS AND DISCUSSION**

**Herbage composition**

Chemical composition is listed in Table 1. It appears that CP and WSC content of forage used in this experiment was quite low (only 63 and 74 g/kg DM for CP and WSC respectively). It was noticed that soil contamination in the forage occurred during harvesting. The CP content of freshly harvested pangola grass observed here (6.3%) is markedly lower than the values (in a range of 8.1 to 15.8%) reported by MULLIK (1999) for the same grass and location. This is surprising because the pasture was fertilized with DAP and urea after slashing. The grass was harvested only once a day at 0745 h, and the morning portion was fed to the animals within 15 min after harvesting whereas the afternoon portions were stored in a cool room at a temperature of 4°C and fed at 1400 and 1900 h. This feeding method seems to have had no effect on chemical analysis as there was only a small difference in CP content between morning and afternoon feeding.
The WSC content of the grass used here was also low (74 g WSC/kg DM). The low WSC observed here is consistent to values for pangola grass and 2 other tropical grasses (setaria and buffel grass) cut during summer (HUNTER et al., 1970). Among the samples analysed by these authors only green stem of setaria grass contained 95 g WSC/kg DM, which is above the minimum value (90 g WSC/kg DM) suggested to affect net energy value of forage (CORBETT et al., 1966). The WSC content of temperate grasses is usually much higher (DAVIES et al., 1991; FULKERSON and TREVASKIS, 1997).

The WSC content is influenced by solar radiation and balance of photosynthesis and respiration processes within plant, so its level fluctuates markedly within a day with the lowest concentration observed in the early morning due to the respiration process during the night (HUMPREYS, 1991; FULKERSON et al., 1994). This is probably one of the factors contributing to the low WSC observed here because the grass was harvested early in the morning (0745 h). FULKERSON and TREVASKIS (1997) showed that the highest WSC content was around 1800h in the afternoon.

The objective in harvesting fresh pangola grass and feeding in pens was to obtain pasture of high quality which would provide data comparable to that from grazing animals. On the basis of chemical composition this was not successful yet the results are very interesting in that they confirm that very low values of EMPS occur in tropical pastures.

### Intake and digestibility

Data of intake and digestibility is shown in Table 2. Dry matter intake was only 1.57% of the body weight (W). The intake of CP was only 469 g/d equal to 71 g CP/kg OM. Digestibility of DM (60%) and OM (69%) was quite high for this grass.

### Table 1. Chemical composition per kilogram dry matter (DM) of freshly harvested pangola grass fed to steers in metabolism crates over 7 day

<table>
<thead>
<tr>
<th>Nutrients</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry matter (g/kg feed)</td>
<td>247</td>
</tr>
<tr>
<td>Organic matter (g/kg DM)</td>
<td>922</td>
</tr>
<tr>
<td>Crude protein (g/kg DM)</td>
<td>63</td>
</tr>
<tr>
<td>Water soluble carbohydrates (g/kg DM)</td>
<td>74</td>
</tr>
<tr>
<td>Neutral detergent fibre (g/kg DM)</td>
<td>680</td>
</tr>
</tbody>
</table>

The values are the mean. Standard deviation (SD) of 4 animals
The extent of voluntary feed intake is determined by interplay between plant properties, activity of rumen microbes, and passage of particles from the rumen. This interrelationship suggests that using simple general relationships between intake and measures of feed chemical composition, feed digestibility, or feed physical properties are most likely to be less than satisfactory (Wilson and Kennedy, 1996). However, it has been well established that there is a close relationship between intake and chemical and physical characteristics of the forage (Milford and Minson, 1966; Hodgson, 1982; Hodgson, 1984). The mean DM intake of steers in the present study was only 1.6% W, a value similar to that recorded previously with forage of this quality (Minson, 1982; 1990).

Rumen fermentation

Mean concentration of NH₃-N in the rumen fluid of steers in this experiment measured 3 and 24 h after morning feeding were 58.2 and 60.7 mg NH₃-N/L respectively. These values were above the minimum level (50 mg NH₃-N/L) for effective rumen microbial activity as suggested by Satter and Slyter (1974). Since CP content of the grass was quite low, there might be a significant contribution of recycled urea into the rumen. Evidence suggests that for ruminants consuming low quality forages (<6% CP/kg DM) urea recycling plays an important part in meeting requirement of N in the rumen (Norton, 1982; 1984). A stable rumen NH₃-N concentration as found here might be explained by the fact that the steers were fed 3 times a day and the feed refusals were usually greater than 2 kg/d so there appeared to be no times that food was not present.

Fractional passage rate

Mean passage rates of fluid and particulate markers (Cr and Yb respectively) from the rumen estimated from their concentration in faecal samples is presented in Table 3. Estimated fluid passage rate from the rumen was 10.0% /h which was higher than that of particulate passage rates (6.7% /h).

The rate of particulate (6.7%/h) and fluid (10.0%/h) dilution observed here was reasonably high and this is usually associated with a high EMPs (AFRC, 1992) but this did not occur here. Fractional flow rates observed in this study were similar to fast fractional outflow rates observed by De Vega and Poppi (1997) (6.7 and 10.1%/h for particulate and fluid respectively) in sheep fed pangola hay and administered with labelled undigested pangola particles and Cr-EDTA. It appears that the predominant limiting factor for this experiment was rumen degradable protein (RDP) adequacy.

Rumen microbial crude protein

Excretion of Ct and PD, and estimated MCP synthesis are listed in Table 4. Daily Ct excretion was 115.26 mmol/d or 1.17 mmol/kg W⁰.75. Allantoin was the predominant compound (93%) in the total PD excreted. The remaining (7%) was uric acid. The molar ratio of PD/Ct was 0.88. The mean value of EMPs was only 71.8 g MCP/kg DOM.

The EMPs observed here (71.8 g MCP/kg DOM) was only 55% of the minimum value (130 g MCP/kg DOM) suggested for forage based diets (SCA, 2007). The EMPs reported here was similar to those of tropical hays (Prior et al., 1998; Bolam et al., 1998; Bowen, 2003; Mullik, 2006; Marsetyo, 2007). This is surprising because green forages are expected to have a much better EMPs than dried ones. The EMPs value under this experimental condition was even lower than the value (90 g/kg DOM) reported by Marsetyo (2007) for the same breed of cattle given green panic hay (5.7%).

The probable argument for this low EMPs is inadequacy of RDP and energy particularlyWSC. The CP content of the grass used here was only 6.3% (Table 1). It is clear from intake data (Table 2) that CP intake was only 71 g CP/kg OM or 104 g MCP/kg DOM. Assuming that degradability of CP in the rumen is 75% (McLennan et al., 1997) then the RDP availability would be only 53 g RDP/kg OM or 78 g RDP/kg DOM. This calculation clearly shows that RDP supply was far below the recommended level (130 to 170 g RDP/kg DOM) by the current feeding systems (SCA, 2007; NRC, 2000). So, any feeding strategies to provide extra RDP is likely to be effective in improving EMPs under this feeding condition. Predicted EMPs in the current study, according to the above feeding standards, is around 78 g MCP/kg DOM which is close to the actual value (72 g MCP/kg DOM) observed here.
Table 4. Creatinine and purine derivative (PD) excretion, and estimated microbial crude protein (MCP) synthesis in steers given freshly harvested pangola grass

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excreted:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Creatinine (mmol/d)</td>
<td>115.26</td>
<td>6.400</td>
</tr>
<tr>
<td>Creatinine (mmol/kg metabolic weight)</td>
<td>1.17</td>
<td>0.060</td>
</tr>
<tr>
<td>Total purine derivatives (mmol/d)*</td>
<td>101.00</td>
<td>3.900</td>
</tr>
<tr>
<td>Allantoin (mmol/d)</td>
<td>93.00</td>
<td>3.400</td>
</tr>
<tr>
<td>Uric acid (mmol/d)</td>
<td>7.00</td>
<td>0.400</td>
</tr>
<tr>
<td>Molar ratio of PD/Creatinine</td>
<td>0.88</td>
<td>0.102</td>
</tr>
<tr>
<td>Molar ratio of allantoin/creatinine</td>
<td>0.80</td>
<td>0.104</td>
</tr>
<tr>
<td>PD Absorbed (mmol/d)</td>
<td>69.60</td>
<td>24.900</td>
</tr>
</tbody>
</table>

Estimated MCP production:
- g MCP/d: 316.00, 113.500
- g MCP/kg metabolic weight: 3.20, 1.118
- g MCP/kg digestible organic matter: 71.80, 15.440

*Only uric acid and allantoin were used in the total PD since concentration of xanthine and hypoxanthine in urine samples was very small. The values are the mean and standard deviation (SD) of 4 animals.

The importance of WSC in determining microbial growth has been proposed (CORBETT et al., 1966; BEEVER et al., 1978; DOVE and MILNE, 1994). Whilst quantitative aspects of WSC have not been established, particularly the ratio of WSC and RDP, earlier experiments (e.g. CORBETT et al., 1966) indicated that diets containing WSC lower than 90 g/kg DM had a lower net energy value. A study by DOVE and MILNE (1994) showed a two fold increase in E_MPS in sheep grazing spring/summer pasture above those grazing autumn pasture. These authors related this improvement to the WSC of pasture though WSC was not directly measured. The WSC content of the grass used in the present experiment was only 74 g WSC/kg DM. This value agreed with values for fresh pangola grass reported by HUNTER et al. (1970). These researchers showed total sugars in the stem fraction of pangola grass was 70 g/kg DM whereas green leaf contained only 25 g/kg DM.

**CONCLUSION**

The E_MPS in steers given freshly harvested pangola grass used in this study was only 71.8 g MCP/kg DOM which is much lower than the values set for forage diets in the current feeding standards. This low E_MPS most probably stems from deficiency of RDP and WSC in this diet.

**REFERENCES**


