NITROGEN DYNAMICS IN THE RUMEN AND ABOMASUM OF SHEEP DETERMINED WITH ¹⁵N-LABELLED AMMONIA OR ¹⁵N-LABELLED DUCKWEED

Damry¹ and J.V. Nolan²

¹Department of Animal Science, Tadulako University, Palu 94118 ²School of Environmental and Rural Science, UNE, Armidale, NSW 2351, Australia E-mail: damry_01@yahoo.com

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ABSTRACT

An experiment was carried out to investigate the dynamics of nitrogen (N) in the rumen and abomasum of rumen and abomasum-cannulated sheep using ¹⁵N dilution techniques. The ¹⁵N tracer was administered into the rumen as ¹⁵N-ammonia or ¹⁵N-labelled duckweed and the transfer of the ¹⁵N label to various N pools was followed. Flow of digesta from the rumen into the abomasum was ascertained by double marker technique with cobalt and acid insoluble ash as liquid digesta and particle digesta marker, respectively. Results showed that the average of rumen water volume was $4.5 \ 1 \pm \text{SEM } 0.57$ and the mean water flow through the abomasum (8.6 \pm 0.45 l/d) was higher than outflow from the rumen (7.4 \pm 0.55 l/d). Nitrogen intake tended to be higher, but total-N passing the abomasum tended to be lower when the sheep were infused by ¹⁵N-ammonia than when they were ingesting ¹⁵N-duckweed. The ammonia concentration in abomasal digesta was about 93 mg N/kg and non ammonia N (NAN) was about 1.58 g N/kg. The rates of flow of total-N as ammonia-N and as NAN did not differ (P>0.05) between animals or diets, with means (\pm SEM) of 57.7 \pm 0.96 and 964 \pm 2.13 mmol/d (or 0.81 and 13.5 g N/d), respectively. About 34-59% of the dietary N was removed from the rumen as ammonia (absorbed and in digesta). The enrichments of rumen ammonia N appeared to have reached plateau values after about 10 h of ¹⁵N-ammonia infusion. The percentage of bacterial-N derived from ammonia-N (from the period of ¹⁵N-ammonia infusion) was 53.63 % (ratio of plateau enrichments) and thus 37.47% of bacterial-N was derived from NAN sources in the rumen. The total ¹⁵N flow through the abomasum was higher (P<0.001) when ⁵N duckweed was given rather than ¹⁵Nammonia $(2.40 \pm 0.02 \text{ versus } 1.29 \pm 0.02 \text{ mmol/d})$. The ¹⁵N in NAN flowing to the abomasum (mmol/d) was also significantly higher (P<0.001) when ¹⁵N-duckweed was given rather than ¹⁵N ammonia, with means of 2.31 ± 0.02 and 1.16 ± 0.00 , respectively. The flow of ¹⁵N in ammonia, on the other hand, was lower (P<0.01) when sheep ingested ¹⁵N-duckweed than when they were infused with ¹⁵N-ammonia (0.09 ± 0.00 v. $0.13 \pm 0.09 \text{ mmol/d}$).

Keywords : Nitrogen Dynamics, Rumen, Abomasum, ¹⁵N-Labelled Duckweed, Ammonia-N, Non Ammonia-N, Bacterial-N

INTRODUCTION

Quantitative models of nitrogen (N) transaction in the rumen (*e.g* Nolan, 1975; Oldick *et al.*, 2000) have been made using data obtained from *in vivo* experiments carried out using ¹⁵N dilution techniques. In such experiments, ¹⁵N-labelled ammonia or urea is introduced into the rumen and the appearance of the ¹⁵N label in secondary N pools such as microbial-N can be determined. The ¹⁵N isotope dilution technique has proved to be a powerful means of understanding the dynamics of N in the rumen. For example, the finding that more than 50 % of microbial-N was derived directly from nitrogenous sources more complex than ammonia (*i.e.* non ammonia-N, [NAN]) when sheep were given lucerne diets was obtained from such experiments (Nolan and Leng, 1972). However, the utilisation of plant NAN materials by microorganisms was determined indirectly after ¹⁵N ammonia utilisation by microbes had been determined, and this method gives lower confidence than a direct measure of plant N incorporation. Our current understanding of plant N behaviour and N transactions among various N pools in the rumen may be further enhanced by the use of ¹⁵Nlabelled feed as the source of the tracer. However, there have not been many *in vivo* studies done using this technique, probably because of economical consideration and time that is needed to produce the ¹⁵N-labelled plant at sufficient quantities as required. Ideally, the ¹⁵N should be incorporated by the plant efficiently during the labelling process and significant quantities of ¹⁵N-labelled materials can be obtained in short time.

In the rumen, the ¹⁵N from the labelled plant will appear in rumen N pools at rates depending on the original distribution of the ¹⁵N in plant N and on the rates of plant degradation. Since ammonia is the end product of degradation of N-containing substances, the appearance of ¹⁵N label in this pool can then be used to estimate the degradation of plant N materials (Chapman and Norton, 1984). Under steady state conditions, all the ¹⁵N present in the plant should enter the ammonia pool if the plant N materials is completely degradable in the rumen. In another words, if the amount of ¹⁵N entering the ammonia pool is less than the ¹⁵N amount available from the plant, this means that some of the plant N materials is never degraded to ammonia.

For such studies, it is necessary to perform two experiments with similar conditions in which ¹⁵Nammonia is used in one experiment and ¹⁵N-feed in the other. Some data (such as irreversible loss of ammonia) are obtained from the ¹⁵N-ammonia experiment and assumed to be applicable to the experiment with ¹⁵N-labelled feed as the source of the tracer. If rumen bacterial sample is isolated in these experiments, its enrichment and the ratio of this enrichment to that of rumen ammonia can be compared. This ratio should be similar in both experiments if the bacterial N is only synthesised from ammonia. The bacterial sample isolated should not be contaminated with ¹⁵N from non-bacterial substances and inspection on this matter should be performed.

This experiment was carried out to investigate the dynamics of N in the rumen and abomasum of sheep using ¹⁵N dilution techniques. However, ¹⁵N was administered into the rumen as ¹⁵N-labelled duckweed, a rapidly growing aquatic plant, as well

as ¹⁵N-labelled ammonia. The transfer of the ¹⁵N label to various N pools was then followed. In this way it was possible to determine, by direct means, the fate of plant-N in the rumen and abomasum including the extent of its assimilation in different forms for synthesis of bacterial protein.

MATERIALS AND METHODS

Experimental Animals and Diets

Duckweed (Spirodella punctata) was labelled with ¹⁵N by growing it on a synthetic medium (Damry, 2006). Two ruminally and abomasally cannulated Merino x Border Leicester crossbred sheep (designated as Sheep C and D) were used as experimental animals. They were kept in individual metabolism crates in a temperature-controlled, continuously lit room in Animal House Complex at the University of New England, Australia. The animals were accustomed to the environment and to a daily experimental ration of 400 g oaten chaff (91.2 % DM, 1.14 g N/100 g DM) and 300 g lucerne chaff (89.4 % DM, 3.40 g N/100 g DM). The feed was provided in approximately equal portions at hourly intervals by means of an automatic belt feeder. The sheep had a continuous access to drinking water.

Experimental Procedures

The experiment involved a study of digesta flow using the dual marker technique of Faichney (1975) in which cobalt as cobalt-ethylene diamine tetra acetic acid (Co-EDTA) complex was used as a liquid digesta marker and acid insoluble ash (AIA) as particle digesta marker. The Co-EDTA complex was prepared according to the method of Uden *et al.* (1980).

The experiment was carried out in two periods using a cross-over design. During the two experimental periods, one week apart, the sheep received either an intra-ruminal continuous infusion of ¹⁵N-ammonia or continuous feeding of ¹⁵N duckweed for 72 h. For the sheep receiving ¹⁵N-ammonia, a solution containing 1.48 mg N/ml (2.96 mol ¹⁵N/100 mol total-N above natural abundance) and 116.5 mg Co/ml, was continuously infused into the rumen at a mean rate of 0.82 g/min. For the other sheep, 16 g ¹⁵Nlabelled duckweed (oven-dried and not milled) containing 2.81 g N/100 g DM was included in the diet and delivered hourly by the automatic, overhead belt feeders. To ensure an equal and steady delivery of the ¹⁵N tracer, the ¹⁵N-duckweed was divided into 4 portions of 4 g each, and each 4 g was distributed evenly over the feed allocated for each 8-h period on the delivery belt. A Co solution (115.9 mg Co/ml) was also continuously infused into the rumen while the sheep was being offered the ¹⁵N-duckweed.

During the ¹⁵N-ammonia infusion, rumen fluid samples were collected, processed and analysed for rumen ammonia-N, NAN and bacterial-N. Abomasum digesta samples were also collected and to avoid effects on digesta flow due to sampling and possible disturbance of the animals, only five samples of abomasum digesta were collected.

In Period 2, the routes of administration of ¹⁵N were reversed. After the period of ¹⁵N administration was terminated, five more samples of rumen contents and abomasum digesta were collected.

Sample Preparation

The rumen fluid samples were immediately centrifuged (20,000 x g, 15 min, 4°C) to obtain a supernate which was acidified with 0.25 ml 18 M H_2SO_4 and stored at -18°C until analysed. Bacteria were isolated from the residue by twice resuspending the residue in 20 ml physiological saline (9 g NaCl/l), recentrifuging and removing the top (bacteria-rich) layers of residue. The final bacterial isolate was suspended in about 3 ml saline solution and stored at -18°C. The abomasum digesta sample was separated into liquid-enriched and particle-enriched fractions by allowing the original digesta (as collected) to stand at room temperature. The more liquid fraction was poured into different container leaving the particle fraction in the original container.

Laboratory Analysis

Ammonia-N (NH₃-N) was removed from the rumen fluid supernate (3 ml) and liquid and particleenriched abomasum digesta (2 gr) by steam distillation after the rumen fluid was made alkaline with saturated Na-tetraborate (4 ml). The NH₃-N was collected into 3 ml 0.025 M H₂SO₄ and titrated to pH 5 with 0.025 M NaOH. It was then re-acidified to pH 3 and dried in an analytical oven at 95°C. The non-ammonia-N (NAN) left was subjected to microkjehdahl digestion using concentrated H₂SO₄ (3.5 ml) and a low Se catalyst tablet. The bacterial isolate (0.5 ml) was similarly digested and the digests were distilled after being made alkaline with 40 % NaOH and then dried as for NH₃-N. Ethanol was distilled between all samples to avoid ¹⁵N cross-contamination. To prepare samples for ¹⁵N analysis, the dried (15NH4)2SO4 was re-dissolved with double distilled H₂O and a solution containing 80-100 μ g N was transferred into an 8x5 mm tin capsule (Elemental Microanalysis Ltd.). The solution was then re-dried in a vacuum desiccator and the capsule was closed and folded. The ¹⁵N enrichment was analysed using a mass spectrometer (Tracermass; Europa Scientific) linked to an N analyser (Carlo Erber).

Cobalt-EDTA infusion solution and in liquidenriched and particle-enriched fractions of abomasum digesta were subjected to Co analysis following a perchloric acid (HClO₄) / hydrogen peroxide (H₂O₂) digestion. The Co concentration was analysed on an Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES; Varian, Model MPX). The AIA concentration in ground (1 mm sieve) feed samples and abomasum digesta was determined according to a method described by Choct and Annison (1990) and Vogtmann *et al.* (1975).

Analysis and Calculation

All enrichment versus time curves were normalised to represent a rate of administration of ¹⁵N into the rumen of 2.88 mmol/d above natural abundance (either as ¹⁵N-ammonia or ¹⁵N-duckweed). The resulting data for ammonia and NAN in rumen fluid supernate and abomasum digesta as well as bacterial N were fitted by curves with two exponential functions using GraphPad Prism version 3.02 for Windows (Motulsky, 1999).

The concentrations of AIA and Co were determined on both particle and fluid fractions of the abomasum digesta: the concentrations of dry matter (DM), NAN and ammonia-N were also determined as well as the enrichments of each of the N fractions. The flow of 'true' abomasum digesta was calculated mathematically (Faichney, 1975) using the AIA and Co as the particle and fluid digesta markers, respectively. A similar mathematical reconstitution was done to enable estimation of flows of individual digesta constituents.

The difference between the total ¹⁵N administered and the total ¹⁵N flowing to the abomasum was a measure of the net ¹⁵N absorption from the rumen. It was assumed that this ¹⁵N was removed by diffusion of ammonia through the rumen wall, and thus ammonia-N absorption rate (mmol/d) was determined by dividing net ¹⁵N absorption (mol ¹⁵N/d) by ammonia-N enrichment (mol ¹⁵N/mol total N). The total ¹⁵N flow through the abomasum was calculated as the sum of ¹⁵N in the outflowing ammonia-N and NAN (the latter being bacterial and non-bacterial [dietary and endogenous] ¹⁵N). The bacterial-¹⁵N flow was calculated from the rumen bacterial-N enrichment and the flow of the bacterial-N fraction in the abomasum NAN. The latter was calculated from the enrichment ratio, abomasum NAN : rumen bacterial-N, when the animals were infused with ¹⁵Nammonia. The ¹⁵N found to be flowing out in the non-bacterial NAN fraction when the ¹⁵N-duckweed was being ingested was considered to represent the rate at which duckweed was escaping undegraded from the rumen: the latter rate divided by the rate of ¹⁵N ingestion gave the proportion of duckweed-N that was escaping degradation in the rumen. (For this calculation, it was assumed that ¹⁵N of endogenous origin in the non-bacterial NAN entering the abomasum was negligible). All data were analysed with an analysis of variance on a personal computer using S-Plus 2000.

RESULTS AND DISCUSSION

The DM contents (g/kg air-dry) of the oaten chaff and the lucerne chaff were 912 and 894 g/kg, respectively, while the corresponding N contents (g/ kg DM) were 11.4 and 34.0. The DM intake (g/d) averaged 641 (± SEM 4.57) g DM/d during the periods when ¹⁵N was administered. The mean DM content of the 'true' abomasum digesta was about 50 g/kg digesta. Between 64 and 73 % of the dietary DM intake was apparently recovered in the digesta flowing through the abomasum (Table 1), but the mean true feed DM digestibility in the forestomachs was calculated to be 45-71 %. (after allowance for the contributions of bacterial and endogenous DM to the abomasal DM, made by assuming the concentration of N in microbial materials was 0.8 g/kg DM and the endogenous N flow was 2 g N/d in materials containing 16 g N/kg DM). Rumen water volume averaged $4.51 \pm SEM 0.57$ and the mean water flow through the abomasum $(8.6 \pm 0.45 \text{ l/d})$ was higher than outflow from the rumen $(7.4 \pm 0.55 \text{ l/d})$, indicating there was a net post-ruminal addition of water to the digesta (Table 1)

Nitrogen intake tended to be higher, but total-N passing the abomasum tended to be lower when the sheep were infused with ¹⁵N-ammonia than when they were ingesting ¹⁵N-duckweed (Table 2). The ammonia concentration in abomasal digesta was about 93 mg N/kg and of NAN was about 1.58 g N/kg. The rates of flow of total-N as ammonia-N and as NAN did not differ (P>0.05) between animals or diets, with means (\pm SEM) of 57.7 \pm 0.96 and 964 \pm 2.13 mmol/d (or 0.81 and 13.5 g N/d), respectively. About 34-59% of the dietary N was removed from the rumen as ammonia (absorbed and in digesta).

 Table 1. Intake of DM And Flows Of DM And Water Out of The Rumen and Through The Abomasum of Sheep
 Given A Diet of Oaten Chaff (400 g/d as fed) and Lucerne Chaff (300 g/d as fed)

	¹⁵ N-ammonia		¹⁵ N-duckweed				
	Sheep C	Sheep D	Sheep C	Sheep D	Mean	SEM	Р
DM							
Intake (g/d)	633	633	649	649	641	4.57	
Abomasum flow (g/d)	416	408	472	439	434	14.4	>0.05
Apparent digestibility (%)	34.2	35.6	27.3	32.4	32.4	1.81	>0.05
Water							
Rumen volume (1)	4.7	3.9	3.3	6.0	4.5	0.57	>0.05
Rumen outflow (l/d)	7.1	7.8	6.0	8.6	7.4	0.55	>0.05
Abomasum flow (l/d)	7.4	8.7	8.8	9.5	8.6	0.45	>0.05
Net water addition (l/d)	0.3	0.9	2.8	0.9	1.2	0.56	>0.05

Enrichments versus time in rumen fluid ammonia-N, NAN and bacterial-N, and ammonia-N and NAN in abomasal digesta are presented for Sheep C and D in Figures 1 and 2, respectively. The enrichments of rumen ammonia N appeared to have reached plateau values after about 10 h of ¹⁵N-ammonia infusion, although considerable variation in the enrichments of ammonia was apparent. The percentage of bacterial-N derived from ammonia-N (from the period of ¹⁵N-ammonia infusion) was 53 63 % (ratio of plateau enrichments) and thus 37 47% of bacterial-N was derived from NAN sources in the rumen.



Figure 1 Enrichments (y axis; mol ¹⁵N/100 mol total) versus time (x axis; h) in rumen fluid ammonia (\circ), rumen fluid NAN (\Box) rumen bacterial-N (\triangle), abomasum ammonia (x) and abomasum NAN (\diamond) during a 72 h administration of 2 μ mol ¹⁵N/min as ¹⁵N-ammonia (top) or ¹⁵N-duckweed (bottom) for Sheep C



Figure 2 Enrichments (y axis; mol ¹⁵N/100 mol total) versus time (x axis; h) in rumen fluid ammonia (\circ), rumen fluid NAN (\Box) rumen bacterial-N (Δ), reconstituted abomasum ammonia (x) and abomasum NAN (\diamond) when 2 μ mol ¹⁵N/min as ¹⁵N-ammonia (top) or ¹⁵N-duckweed (bottom) was continually supplied to the rumen (Sheep D).

The total ¹⁵N flow through the abomasum was higher (P<0.001) when ⁵N duckweed was given rather than ¹⁵N-ammonia (2.40 ± 0.02 versus 1.29 ± 0.02 mmol/d). The ¹⁵N in NAN flowing to the abomasum (mmol/d) was also significantly higher (P<0.001) when ¹⁵N-duckweed was given rather than ¹⁵N ammonia, with means of 2.31 ± 0.02 and 1.16 ± 0.00 , respectively. The flow of ¹⁵N in ammonia, on the other hand, was lower (P<0.01) when sheep ingested ¹⁵N-duckweed than when they were infused with ¹⁵N-ammonia ($0.09 \pm 0.00 \nu$, $0.13 \pm 0.09 \text{ mmol/}$ d).

Mean DM intake and DM flow through the abomasum was 641 and 434 g/d, respectively. Thus, about 32.4 % of the ingested DM was apparently digested in the rumen. However, further calculations suggested that the true digestibility of DM in the rumen was about 71 %, a coefficient that is similar to values obtained in comparable published studies (SCA, 1990). The rumen water outflow rate, calculated from the rate of decline in Co concentration after the infusion was terminated, was 7.4 l/d. There was a slightly higher flow rate of water (8.6 l/d) through the abomasum suggesting a *net* post-ruminal water addition, probably as gastric secretions, of about 1.2 1/d: the actual water input may have been higher than the net estimate if water is also absorbed in this part of the gut of the sheep (e.g. in the omasum). The total N flowing through the abomasum was 15.2 g/d which when compared with a mean N intake of 14.3 g N/d indicated that there was a net addition (*i.e.* the excess of endogenous input of N over that absorbed as ammonia or other N) between the mouth and abomasum of about 0.9 g N/d. Harrop (1974) estimated that the daily total N secreted in gastric juice was 0.51-2.86 g N, but this amount would not explain the 0.9 g N/d net gain if, as the current results suggest, there was also an absorption of 5-8 g N/d.

About 95 % of the total N flowing through the abomasum in this experiment was NAN, probably consisting mainly of dietary 'escape' protein and microbial protein formed in the rumen, but there would also have been a small fraction of endogenous N. Because there was a net loss of ammonia from digesta leaving the rumen and passing through the abomasum (about 0.2 g N/d), the endogenous N added to the abomasum may have been mostly in the form of NAN, *e.g.* urea and pepsinogen.

Results obtained in when ¹⁵N-ammonia was infused into the rumen indicated that the rumen ammonia enrichment had reached a plateau value 10 20 h after the start of infusion. There was a considerable between-sample variation, however, in the rumen ammonia enrichments. The most likely reason for this variation is the high fractional rate of turnover of ammonia: this is related to the small compartment size and the high rate of flux of ammonia through the compartment. To some extent, the problem could also have been due to non-ideal mixing of the tracer with unlabelled N, but precautions were taken to minimise this effect, e.g. the sampling probe was positioned at a distance from the infusion line to prevent the samples taken being contaminated by the ¹⁵N in the infusion solution. Variation due to the inaccuracies in the mass spectrometer analyses for determining enrichment can be discounted because the variation was apparent only in the rumen ammonia results (i.e. not in other samples) and, in addition, checks showed that the ¹⁵N analyses were both precise and repeatable.

There was considerable variation between animals in the estimates of the proportions of bacterial-N synthesised from ammonia-N when ¹⁵N-ammonia was infused into the rumen. This was probably mainly due to uncertainties about the plateau ammonia enrichments. Additional variation may have been due to different bacterial populations present in the rumen which can occur even when animals ingest the same diet (Armstead and Ling, 1993) or to possible errors associated with predicting the true plateau enrichment of rumen ammonia-N. The proportion of bacterial-N derived from ammonia-N was also lower when ¹⁵Nammonia was infused than when it was injected into the rumen (Damry and Nolan, 2002). The recycling of the absorbed ¹⁵N-ammonia and ¹⁵N-bacteria during the longer period of infusion may have been responsible for lower estimates of the ammoniaderived bacterial-N for the tracer infusion experiments as compared to the tracer injection experiment.

When the sheep were ingesting ¹⁵N-duckweed, the enrichment of rumen ammonia-N also increased exponentially with time, but the rate of increase in enrichment (*i.e.* from the tangent to the fitted curve) was slower, and the absolute values were much lower than those obtained when the same quantity of ¹⁵N was infused in ammonia. It is, therefore, clear that the ¹⁵N ingested in duckweed was not quantitatively degraded all the way to ammonia-N. The ¹⁵N must have been utilised for microbial protein synthesis in NAN forms, or it escaped either undegraded or partially degraded to the abomasum. The ammonia-N enrichment in the sheep ingesting duckweed would have been due, in part, to the direct contribution of ammonia present in the ingested duckweed at the time of its ingestion (about 8 % of total N). However, the slow rate of increase in enrichment also suggests there was a delayed contribution, probably via intraruminal (and to a lesser extent extra-ruminal) recycling of labelled microbial-N.

The ¹⁵N administered that was not recovered in the abomasum must have been ¹⁵N absorbed from the rumen. The non-recovered fraction was significantly higher (P<0.001) when ¹⁵N-ammonia was infused than when ¹⁵N-duckweed was ingested because ammonia absorption across the rumen wall is an important route of loss of ammonia that is not assimilated by bacteria. Also, because ingested duckweed N was only partially degraded in the rumen, the undegraded portion escaped to the intestines. Assuming that all the non-recovered ¹⁵N (mol ¹⁵N/d) was removed from the rumen as absorbed ammonia, the ammonia absorption rate (mol N/d) could be calculated from knowledge of rumen ammonia-N enrichment at plateau (mol ¹⁵N/mol total N). The estimate of ammonia absorption when ¹⁵N ammonia was administered (0.51 mol/d, 7.1 g N/d) tended to be higher than that obtained when ¹⁵N-duckweed was ingested (4.0 g N/d), but the former value may be more nearly correct because of the earlier and more complete attainment of plateau enrichment in rumen fluid ammonia-N during the ¹⁵N-ammonia infusion.

The flow rate of ammonia-N out of the rumen was about 0.2 g/d higher than the rate of ammonia-N flow through the abomasum (Table 2), indicating there was a net loss of ammonia from digesta passing through the omasum or abomasum, although absorption of ammonia from the abomasum is unlikely (Williams et al., 1968). At the same time, however, the ammonia-N enrichment in the abomasum was lower than the rumen ammonia-N enrichment indicating that some unlabelled ammonia-N was added to digesta in one or both of these organs. The proportion of the abomasal ammonia-N derived from the rumen was about 85% implying that about 15% of the ammonia flowing through the abomasum was added post-ruminally, probably from the degradation of blood urea entering the abomasum in gastric juice secretion (Harrop, 1974).

The proportion of bacterial-N in the abomasum NAN was estimated from the ratio of enrichments in abomasum NAN and bacterial-N isolated from the rumen fluid. This ratio was only applicable when

Table 1.Intake of N and Kinetics of Various N-Containing Fractions in The Rumen and Abomasum of Sheep Given a Diet of Oaten Chaff (400 g/d) and Lucerne Chaff (300 g/d) and Either ¹⁵N-ammonia or ¹⁵N-labelled Duckweed

	¹⁵ N-ammonia	¹⁵ N-duckweed	SEM	Р
N intake (g/d)	14.9	13.6		
Rumen ammonia-N				
Concentration (mg N/l)	131	137	4.53	>0.05
Compartment size (g N)	0.57	0.65	0.08	>0.05
Absorption rate (g N/d)	7.09	3.92	0.82	>0.05
Outflow rate (g N/d)	0.98	0.99	0.06	>0.05
Abomasum ammonia-N				
Concentration (mg N/l)	98.6	91.7	4.57	>0.05
Flow rate (g N/d)	0.81	0.79	0.01	>0.05
Proportion from rumen ammonia-N	0.73	0.97	0.07	>0.05
Bacterial-N				
Proportion from rumen ammonia-N	0.50	0.50		
Proportion of abomasum NAN	0.81	0.81		
Abomasum flow (g N/d)	10.4	9.98	0.02	>0.05

Values Determined when ¹⁵N-ammonia was Infused into The Rumen

	¹⁵ N-ammonia	¹⁵ N-duckweed	SEM	Р
 ¹⁵N administered ¹⁵N flow in abomasum NAN ¹⁵N flow in bacterial-N ¹⁵N flow in non-bacterial-N ¹⁵N flow in abomasum ammonia-N Total ¹⁵N flow through the abomasum ¹⁵N absorbed from the rumen 	2.88 1.16 1.16 0.00 0.13 1.29 1.59	2.88 2.31 1.46 0.85 0.09 2.40 0.48	0.33 0.09 0.01 0.32 0.32	<0.001 <0.01 <0.01 <0.001 <0.05

Table 1. Amounts of ¹⁵N Flowing in Various N Materials Through The Abomasum (mmol/d) During
a Continuous Provision into The Rumen of ¹⁵N-ammonia or ¹⁵N-Labelled Duckweed

¹⁵N-ammonia was infused into the rumen (not when the tracer was provided by the ¹⁵N-duckweed as, in this case, labelled NAN in the abomasum could be non-bacterial in origin). The ratio, which was about 81 %, was assumed to apply throughout the experiment and was also used in calculations for the duckweed-ingesting animals. The ratio was higher than that obtained for sheep with similar intakes of DM by Siddons *et al.* (1982), and the estimates of flow of bacterial-N through the abomasum (10.5 g N/d), and the efficiency of bacterial-N synthesis (39 45 g N/kg DM apparently digested in the rumen, or 22 45 g N/kg DM truly digested in the rumen) also seemed to lie at the high end of the range of published values (SCA, 1990).

The proportion of duckweed-N that escaped rumen fermentation was estimated from the data describing the ¹⁵N balance across the rumen (Table 3). The ¹⁵N flowing through the abomasum would have been present as both ammonia-N and NAN, the latter consisting of microbial-N and non-microbial-N. The non-microbial-N probably also contained relatively small amounts of labelled endogenous NAN. From the knowledge of bacterial-N enrichment and the proportion of bacterial-N in the abomasal NAN, the amount of ¹⁵N in bacterial-N flowing through the abomasum was calculated. Subtracting the result from the total ¹⁵N flow gave an estimate of the non-bacterial ¹⁵N-labelled NAN flow (0.85 mmol/ d). If the non-bacterial-N flow through the abomasum is assumed to be solely from labelled escape duckweed, then about 30 % of the duckweed N ingested escaped digestion in the rumen which also implies that about 70 % of the dietary duckweed-N was degraded in the rumen. About 25 % of the ¹⁵N

in the ruminally degraded component of the duckweed was absorbed through the rumen wall, probably as ammonia as mentioned above. The majority of the ¹⁵N in duckweed digested in the rumen, however, was converted to, and recovered as bacterial-N flowing through the abomasum: this bacterial-N would then be available for intestinal absorption and utilisation by the host.

It may be concluded that the use of ¹⁵N- labelled plants such as duckweed, in combination to that of ¹⁵N-labelled ammonia, in ¹⁵N tracer studies will enhance further our undertandings of N behaviours and transactions in ruminant nutrition.

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