Digestion and microbial protein synthesis in sheep as affected by exogenous fibrolytic enzymes

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Abstract
The aim of the study was to determine the effect of an exogenous fibrolytic enzyme (EFE) on the ruminal disappearance of dry matter (DM), neutral detergent fibre (NDF), crude protein (CP) and on the microbial protein synthesis (MPS) in sheep of a milled substrate consisting of a 50 : 50 mixture of lucerne hay (LH) and wheat straw (WS). Four cannulated Döhne-Merino sheep were randomly assigned to two groups in a 2 x 2 cross-over experiment. Additional to the in situ study, the effects of an exogenous fibrolytic enzyme (EFE) on the in vitro gas production (GP) and ANKOM digestion systems on the mixture of milled LH and WS were determined. The substrate was pre-treated with distilled water (control) or EFE (treatment) 12 hours prior to incubation to allow interaction between the substrate and enzyme. Adding EFE significantly improved the in vitro cumulative gas production (GP), but it did not affect the in vitro MPS measured as purine derivates of the GP residues. The enzyme significantly increased the in vitro DM and NDF disappearances after 36 hours. However, the in situ disappearances of DM, NDF and CP of the roughage mixture were not affected by the EFE. Further, the in situ MPS was increased significantly with the EFE treatment of the mixed substrate. It is concluded that the improved ruminal digestion of DM, NDF and CP is not only due to direct enzyme hydrolysis, but also due to the improvement of MPS.

Keywords: Fibre digestion, in vitro and in situ digestion, ruminants

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Introduction
Ruminant production systems are based on the utilization of forages, with fibrous material being important for salivation, rumen buffering and efficient production of rumen end products. However, less than 65% of the fibrous plant cell walls are efficiently digested in the total digestive tract of ruminants (Van Soest, 1994). With 40% – 70% of the forage DM consisting of cell walls, it would be important to improve ruminal fibre digestion. Forage digestibility has been improved by several biotechnological products, i.e. ionophores, direct fed microbial products and a number of enzymes. In the past 10 to 15 years, the use of EFE as feed additives has shown promise at hydrolyzing plant cell walls (Beauchemin et al., 2004). However, their biological response in ruminant systems is difficult to predict because of inconsistent results (Beauchemin et al., 2003). In an effort to improve the nutritive value of ruminant feedstuffs, a novel EFE, produced from a fungal strain isolated from soil in South Africa and cultivated on wheat straw, was evaluated. The EFE was characterised with xylanase as the major fibrolytic activity (Cruywagen & Van Zyl, 2008) and evaluated in a ruminant system using in vitro and in situ techniques. The aim of the study was to determine the EFE effect on the DM, CP and NDF digestion of a mixture of lucerne hay and wheat straw (50 : 50) to improve MPS.
Materials and Methods

The protocol of the study was approved by the Animal Care and Use Committee of Stellenbosch University (SU ACUC, ethical clearance number: 2006B03005). Four cannulated Döhne-Merino sheep were randomly assigned into two groups in a 2 x 2 cross-over experiment. The study was conducted as two main studies: 1) *in situ* digestibility and 2) *in vitro* digestibility based on the *in vitro* gas production (GP) and *in vitro* digestibility (ANKOM Technologies, Fairport, NY, USA) of the pre-treated substrate. The gas production system used was based on the Reading Pressure Technique (RPT) described by Theodorou et al. (1994). Each animal received a basal diet *ad libitum*, supplemented in the morning with 300 g/day of a concentrate providing 880 g DM/kg feed, 10, 25, 1.5 and 0.2% CP, NDF, calcium (Ca) and phosphorus (P) per kg DM, respectively. The basal diet consisted of 298.5 g LH/kg, 298.5 g WS/kg, 149.2 g corn starch/kg, 238.8 g molasses meal/kg and 15 g/kg of a trace mineral and vitamin premix, to contain 830.3 g DM/kg, 75.5 g CP/kg DM and 361.5 g NDF/kg DM. The basal diet and water were offered *ad libitum* daily. Treatments of the milled substrate (a 50 : 50 mixture of LH and WS) consisted of a control (distilled water only) and EFE, a South African fungal enzyme cultivated on wheat straw and developed at the Department of Microbiology at Stellenbosch University. The enzyme was used as either a lyophilised enzyme concentrate (0.5 g enzyme powder + 68 mL of distilled H₂O) or fresh supernatant (one mL of enzyme supernatant + 200 mL of distilled H₂O) to supply the exact same amount of enzyme activity. The EFE was analyzed to contain cellulases, xylanases and mannanases, with xylanase being the major fibrolytic activity (Cruywagen & Van Zyl, 2008). The EFE was applied 12 hours prior to the *in vitro* or *in situ* incubation as pre-treatment of the milled and sieved substrate, to allow for an enzyme-substrate interaction time as recommended by Beauchemin et al. (2003). The application of treatments was at a ratio of one mL of the EFE dilution or distilled water to 0.5 g substrate for the *in vitro* studies. Similarly, the standard diet was treated 12 hours before feeding at a daily level of 5 mL/kg feed to provide either no enzyme (Control: distilled water) or EFE treatment for the *in situ* study.

To ensure good homogeneity, 5 mL enzyme concentrate or distilled water was added to 100 mL of distilled water before being sprayed on one kg of the basal diet. As described by Useni (2011), a series of *in situ* and *in vitro* incubations over 48 hours were simultaneously conducted following a 10 day adaptation period to the basal diet. While the *in vitro* nylon bag technique was executed accordingly to the Daisy® procedure as described by the manufacturer (ANKOM® Technology Corp., Fairport, NY, USA), the *in vitro* GP and the *in situ* digestibility were conducted according to Goosen (2005) (based on the system described by Theodorou et al. (1994)) and Vanzant et al. (1998), respectively. A buffered media solution (Goering & Van Soest, 1970) was used for the *in vitro* incubations while the rumen liquor acquired was collected at 06h00 from the rumen cannulated sheep adapted to the basal diet. The DM, CP, NDF and purine derivatives of original sample and residues were measured according to chemical analyses as described by the AOAC (1995). Data was subjected to a two-way repeated measures of analysis of variance (ANOVA) and treatment means were compared with a Bonferroni test to determine significant differences, using SAS (2006 ). Significance was declared at $P < 0.05$.

Results and Discussion

The effect of EFE on GP is indicated in Figure 1(a) as the cumulative gas produced over a 48 h incubation period. At the end of the 48 h incubation period, MPS was measured on the residual material as purine derivatives of the mixed substrate (Figure 1(b)). Pre-treatment of a mixed substrate with EFE resulted in an improved cumulative GP ($P < 0.01$) but had no effect ($P >0.05$) on the MPS (measured as purine derivate) with EFE treatment was observed (Figure 1).

Additionally, the *in vitro* disappearance of DM and NDF were improved at 36 hours of incubation due to the enzyme treatment, by respectively 6.28% ($P = 0.03$) and 2.85% ($P = 0.05$). Similar to these findings, Goosen (2005) reported a positive EFE effect on the cumulative GP and the *in vitro* DM and NDF degradation of wheat straw. Beauchemin et al. (2003) reported that the addition of EFE to dry feedstuffs before feeding enhances the binding of EFE with the substrate, which can improve the resistance of EFE to proteolysis and prolong their residence in the rumen. This can result in an enhancement of colonization and digestion of slowly degradable plant cell wall fractions by ruminal micro-organisms (Feng et al., 1996).

**Figure 1** The cumulative gas production (GP) (mL/g OM) of the EFE treated or control mixed substrate (a) and the microbial protein synthesis (MPS) measured as purine derivates (RNA equivalent in µg/DM g) on residues of GP after 48 hours (b). Error bars indicate the standard error of the mean.

The *in situ* disappearances of DM, NDF and CP did not differ (*P >0.05*) between the EFE treated substrate and the control (Figure 2). This may be due to the small number of cannulated sheep (*n = 4*) used in the study and the relatively high coefficients of variation associated with measuring ruminal digestion. The *in situ* MPS, measured as purine derivates, was, however, increased with EFE addition to the substrate (*P <0.01*) (Figure 3). Similarly, Nserekho *et al.* (2002) found that EFE indirectly increased the attachment and the number of cellobiose- and glucose-utilizing bacteria in the rumen. Giraldo *et al.* (2008) reported that treating high-forage diets with EFE increased fibrolytic activity and stimulated the *in vitro* numbers of micro-organisms. Likewise, Yang *et al.* (1999) found that EFE significantly improved the supply of metabolizable protein to the small intestine, measured as $^{15}$N. In two other studies evaluating this enzyme at a similar dose, it improved (*P <0.05*) body weight gains and feed conversion efficiency when fattening lambs on forage based-diets (Cruywagen & Goosen, 2004; Cruywagen & Van Zyl, 2008). It is suggested that the EFE increased the hydrolytic capacity in the rumen probably improving fibre digestibility.

**Figure 2** The effects of an exogenous fibrolytic enzyme (EFE) on *in situ* disappearances of (1) CP, (2) DM and (3) NDF of the mixed substrate. Error bars indicate the standard errors of mean.
Figure 3 The effects of an exogenous fibrolytic enzyme (EFE) on the microbial protein synthesis (MPS) measured as purine derivates (RNA equivalent in µg/g DM) on residues of *in situ* nylon bag digestion. Error bars indicate the standard errors of mean.

There are several possible explanations for the increased MPS observed in the *in situ* study. One such explanation is the direct effect of the EFE that can randomly release reducing sugars and possibly make more nutrients available for utilization by the microorganisms (McAllister *et al*., 2001), arising from the partial solubilisation of cell wall components (Krause *et al*., 1998). As stated by Beauchemin *et al*., (2004), the presence of soluble sugars would supply sufficient additional available carbohydrates to shorten the lag time needed for microbial colonization and also enhance the rapid microbial attachment and growth. The EFE effect on nutrient release can also favour an increased production of the glycocalyx, which is produced by bacteria, allowing adhesion between bacteria and between bacteria and substrate. Bala *et al*., (2009) observed an increase in milk yield of lactating goats. The authors ascribed this to the improvement of the energy availability and the utilization of microbial digestible protein, estimated as purine derivatives and creatinine excreted in urine (Bala *et al*., 2009). These authors speculated that the EFE was able to free the trapped nutrients in the cell wall networks of roughages. The increased availability of nutrients to micro-organisms in the rumen was reported by Chakeredza *et al*., (2002) to improve the yield of fermentative end-products due to a change in the ratio of micro-organism biomass to the digestible energy in the rumen.

**Conclusion**

Research on EFE in ruminant diets reports on improvements of forage utilization, production efficiency and reduced nutrient excretion. A South African EFE, produced from a fungal strain isolated from SA soil, shows potential to enhance fibre digestion of forage based-diets fed to ruminants. The current study indicated an increased *in vitro* GP, coupled with increased *in vitro* DM and NDF disappearances at 36 h of digestion. Although *in vitro* digestibility was increased, results did not indicate a significant increase in MPS measured on the residual matter after *in vitro* digestion. Microbial protein synthesis (MPS) was however increased on the residual matter after *in sacco* digestion, especially after 24h digestion periods and up to 48 h. The improvement of feed digestion in this study was attributed to enhanced *in situ* MPS, possibly due to the EFE treatment resulting in an increased availability of nutrients for microbial growth. The net effect of EFE on feed digestion is a combined result of direct hydrolysis of substrate, increased microbial attachment, stimulation of the rumen microbial population and synergistic effects with hydrolyases of ruminal micro-organisms.
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