In vitro gas production from cassava peels supplemented with unconventional nitrogen sources and forages by small ruminants

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ABSTRACT: A study was conducted to ascertain the nutritional value of N-source treated and forage supplemented cassava peels (CSP) using in vitro gas production technique. Cassava peels were treated and supplemented with materials rich in nitrogen: fertilizer grade urea (T1 = CSP + U), broiler litter (T2 = CSP + BL), cassava forage (T3 = CSP + CSF) and sweet potato forage (T4 = CSP + SPF). Results revealed that, CSP supplemented with CSF (T3) and SPF (T4) recorded better gas production (31.90 and 32.20 mL/200 mg DM), metabolizable energy (ME) (7.54 and 7.67 MJ/kg DM), and organic matter digestibility (OMD) (52.02 and 52.80%), respectively. Similarly, the estimated short chain fatty acids (SCFA) revealed higher values of 0.70 and 0.71 µM, respectively, for cassava and sweet potato forage supplementation compared to those fed urea and broiler litter-treated cassava peels. The superiority exhibited by the forage supplemented cassava peel diets for meeting the ME levels, SCFAs and OMD reveals that, cassava peels supplemented with either cassava leaves or sweet potato forage are valuable sources of nutrients for sheep and goats in terms of meeting their energy requirements for maintenance and part or all their milk production needs depending on the level of productivity.

Keywords: Cassava peels, forage, goats, nutritional value, sheep, supplementation, sweet potato.

INTRODUCTION

Rapid growth in human and livestock population in Sub-Saharan Africa is creating unprecedented increases in food and feed demands. Such population pressure on a fixed land base is, likely, to promote stern competition for resources and, progressively, force agriculture in the direction of intensification (Smith et al., 1997). For such intensification to work, effectively, under smallholder livestock farming, research is required to develop alternatives that will promote better resource use, less environmental impacts, increase farm income, and promote farmers’ livelihoods in Nigeria and other parts of Sub-Saharan Africa.

The in vitro method is a laboratory technique for feed evaluation, which has numerous advantages over the in vivo method since it utilizes small amounts of test feeds suitable for screening large samples of feed (Njidda et al., 2010). Other benefits of the in vitro technique are that, it is less expensive, less time consuming, accurate, allows more precise maintenance of incubation conditions and simulates the rumen fermentation process making it useful for evaluating the potentials of feedstuffs to supply nutrients to ruminants (Akinfemi et al., 2009). It helps to simulate the digestive processes generated by microbial activities and to understand feed fermentation and degradability as a function of nutritional quality and nutrient availability for the rumen bacteria (Murillo et al., 2011). It is less animal dependent; more appropriate for characterizing soluble and small particulate feeds and can be automated thus reducing the labour input (Adesogan, 2002). It has been widely used to estimate the nutritive quality of cereal straws (Valizadeh et al., 2010). Similarly, it has also been successfully used to evaluate compounded feeds as well as for determining the effects of anti-nutritive factors in African browse plants on rumen
fermentation and for predicting feed intake, digestibility, microbial nitrogen supply and animal performance (Getachew et al., 2004a; Akinfemi et al., 2009; Boga, 2014).

A survey conducted in cross River State of Nigeria indicated that, crop by-products predominantly cassava peels are available in large quantities in the rural communities that may serve as valuable feed resources for small ruminants (Kalio et al., 2013). Consequently, there is also the need to upgrade the feeding value of these crop by-products by interfering with the protective effect of lignin on the availability of the substrate to the rumen bacteria or to hydrolytic enzymes (Preston and Leng, 1986). This process can be enhanced by using simple inexpensive processes such as treatment with non-protein nitrogenous (NPN) sources like urea and broiler litter or supplementation with nitrogenous forages such as cassava and sweet potato leaves that could increase the utilization of these crop by-products through digestibility trials (Kalio et al., 2014). However, there is little information on in vitro ruminal gas production by small ruminants fed unconventional nitrogen treated or forage supplemented cassava peels. Therefore, the aim of this study was to evaluate in vitro gas production from cassava peels supplemented with unconventional nitrogen sources and forages by small ruminants.

MATERIALS AND METHODS

Experimental site and feed preparation

The study was conducted at the Ruminant Nutrition Laboratory of the University of Ibadan, Ibadan, Nigeria while the feed samples were collected from the Teaching and Research Farm of the University of Calabar, Calabar, Nigeria. Dried cassava peels (CSP) were treated and supplemented with different unconventional nitrogen sources: Urea (U), broilers litter (BL) and forages (cassava forage- CSF and sweet potato forage – SPF) to give four treatments namely:

1. T1 = cassava peels + urea (CSP + U);
2. T2 = cassava peels + broiler litter (CSP + BL);
3. T3 = cassava peels + cassava forage (CSP + CSF), and
4. T4 = cassava peels + sweet potato forage (CSP + SPF).

Experimental animal management and feeding

Prior to the commencement of the in vitro gas production studies, a ram weighing 11 kg was kept for an adaptation period of 5 days in a metabolic cage measuring 3 m x 2 m x 4 m and used for rumen fluid collection. Feed, water and mineral-salt licks were provided ad libitum. Feed was provided, daily, at the rate of 5% of the body weight of the ram. The feed fed to the ram comprised a basal diet of 60% Panicum maximum and 40% concentrate (40% corn, 10% wheat offal, 10% palm kernel cake, 20% groundnut cake, 5% soya bean meal, 10% dried brewers grain, 10% common salt, 3.75% oyster shell, and 0.25% fish meal).

In vitro gas production

Rumen fluid was obtained from the ram through a suction tube before the morning feed and put into a thermo flask that had been pre-warmed to a temperature of 39°C (Babayemi et al., 2009). The incubation was conducted following standard procedures using 120 ml calibrated syringes in three batches maintained at a temperature of 39 to 40°C (Menke and Steingass, 1988). About 200 mg of the feed samples were then weighed into the syringe and added to 30 mL of inoculums containing cheese cloth strained rumen liquor and buffer (comprising 9.8 g NaHCO3 + 2.77 g, Na2HPO4 + 0.57 g KCl + 0.47 g NaCl + 0.12 g MgSO4.7H2O + 16 g CaCl2; 2H2O) per litre.

Rumen liquor and buffer were mixed together in the ratio 1:4 (v/v) under continuous flushing with CO2. Using 50 mL calibrated plastic syringe, 30 mL of inoculums was dispensed into the substrate through the silicon tube. The plunger was pushed upward by pushing the inoculums to the tip of the syringe to completely eliminate air. The silicon tube in the syringe was then tightened by a metal clip so as to prevent escape of gas. The volume of gas produced was measured at 3 hourly bases (3, 6, 9, 12, 15, 18, 21, 24, 27, 30, 33, 36, 39, 42, 45, and 48 h). The volume of gas produced was then read by measuring the space formed between the top of the piston and the liquid in the syringe. At the termination of the 24, 36 and 48 h incubation periods, 4 mL of NaOH (10 M) was introduced to each of the syringes containing the samples for methane (CH4) gas determination. The gas volume was determined by difference, by subtracting the average gas production values of the blank from the individual feed samples. The volume of CO2 was determined by subtracting the CH4 gas production from the gas volume.

Chemical analysis

Chemical analysis of the feed samples for this study were similar to the procedures and results obtained by Kalio et al. (2014) in their study while investigating the performance of West African Dwarf (WAD) goats fed N-treated source and forage supplemented cassava peels in humid Cross River State, Nigeria.

Metabolizable energy, organic matter digestibility and short chain fatty acid estimation

Metabolizable energy was estimated as (ME, MJ/kg DM) = 2.20 + 0.136*Gv + 0.057*CP + 0.0029*CF. Similarly,
organic matter digestibility (OMD) was estimated as
OMD\% = 14.88 + 0.889∗Gv + 0.45∗CP + 0.0651∗XA, while
short chain fatty acid (SCFA) was estimated as SCFA =
0.0239∗Gv – 0.0601; where Gv = net gas production
(mL/200 mg DM), CP = crude protein, CF = crude fibre,
and XA = ash contents, respectively, of the incubated
samples (Menke and Steingass, 1988).

Statistical analysis

Data obtained from the in vitro studies were analysed
using analysis of variance (ANOVA) as described by SAS
(1999). Where significant differences were detected,
means were separated using the New Duncan’s Multiple
Range Test (Steel and Torrie, 1980).

RESULTS AND DISCUSSION

Proximate composition of experimental diets

The proximate composition of forage supplemented and
N-treated cassava peels for the study is similar to those
reported by Kalio et al. (2014) in their study while
investigating the performance of West African Dwarf
(WAD) goats fed N-treated source and forage
supplemented cassava peels in humid Cross River State,
Nigeria.

Estimation of the nutritional value by in vitro gas
production techniques

The results of the in vitro gas production, estimated
metabolizable energy (ME), short chain fatty acid (SCFA)
and organic matter digestibility (OMD) of urea and broiler
litter treated and forage supplemented cassava peels via
the in vitro gas production technique are presented in
Table 1. There were significant (p < 0.05) variations in
the gas volume (Gv), metabolizable energy (ME), short chain
fatty acids (SCFAs) and organic matter digestibility (OMD)
among the cassava peels treated with urea (T1: CSP + U)
and broiler litter (T2: CSP + BL) and those cassava peels
supplemented with cassava forage (T3: CSP + CSF) and
sweet potato forage (T4: CSP + SPF).

The gas volume (Gv) produced in order of decreasing
volumes for the different feedstuffs after 48 hours of
incubation were 32.20, 31.90, 21.17 and 18.17 mL/200 mg
DM for T4 (CSP + SPF), T3 (CSP + CSF), T2 (CSP + BL)
and T1 (CSP + U), respectively. However, the gas volumes
produced by T4 (CSP + SPF) and T3 (CSP + CSF) were
not significantly (p > 0.05) different. This study revealed
that, the cassava peels supplemented with the forages
from cassava and sweet potato plants recorded the
highest gas volumes after 48 hours of incubation. This may
be attributed to the fact that, the forages have fermentable
and readily degradable cell wall fractions, which would
increase the substrates available to cellulytic microbes
with a consequent increase in the population of these
microorganisms (Van Soest, 1982). Similarly, the
presence of these microbes influences the extent and rate
of substrate degradation, which is related to the gas
volumes produced (Blümmel et al., 1997). The gas
produced by the different feedstuffs increased with
increasing hours of incubation, but gradually stabilized
with no additional gas volumes produced between 42 and
48 hours of incubation. The gas production trends of the
different feedstuffs are presented in Figure 1.

The metabolizable energy (ME) of the different
feedstuffs in order of decreasing values after 48 hours of
incubation were 7.67, 7.54, 6.56 and 6.42 MJ/kg DM for T4
(CSP + SPF), T3 (CSP + CSF), T2 (CSP + BL) and T1(CSP + U),
respectively. This is slightly higher than the range of
ME values (4.46 to 7.42 MJ/kg DM) reported by Songsak
et al. (2007) for some energy feeds in Thailand. However,
the metabolizable energy of the feedstuffs: T4 (CSP + SPF)
and T3 (CSP + CSF) were not significantly (p > 0.05)
different and were higher. The results revealed that, the
cassava peels supplemented with cassava forage (T3:
CSP + CSF) and sweet potato forage (T4: CSP + SPF)
possess a higher ME values than the ME values for T1
(CSP + U) and T2 (CSP + BL). However, it is important to
note that urea has no energy value of its own (Maynard et
al., 1984). Consequently, the energy value recorded in T1
(CSP + U) is simply due to the associative effect of
cassava peels (CSP) with urea (U) in the feed mixture.
This is in agreement with the findings of Getachew et al.
(2004b), who explained that, feeding straw alone to a
ruminant reduces its digestibility, but by adding nitrogen in
the form of urea or protein, the digestibility of the straw will
be increased and in turn, the energy derived from straw
organic matter in the diet will also be increased. The ME
values recorded in this study revealed that, the ME of the
different diets were within the recommended ME values for
an average diet (6 to 13 MJ/kg DM) (Steele, 2006) hence,
can fulfil the energy requirements for the West African
Dwarf (WAD) small ruminants (sheep and goats).
Furthermore, the derivation of these ME values for these
feeds could be valuable for purposes of ration formulation
and to set the economic value of the feeds for trading
(Njidda et al., 2010).

The short chain fatty acids (SCFAs) or volatile fatty acids
(VFAs) of the different diets in order of decreasing values
after 48 hours of incubation were 0.71, 0.70, 0.45 and 0.37
µM for T2 (CSP + SPF), T3 (CSP + CSF), T2 (CSP + BL)
and T1 (CSP + U), respectively. The SCFA value of the T4
(CSP + SPF) treatment was higher although, they were not
significantly (p > 0.05) different from the T3 (CSP + CSF)
treatment group. The T4 (CSP + SPF) and T3 (CSP + CSF)
treatment groups recorded the highest values of SCFAs.
The SCFAs or VFAs such as acetic, propionic, butyric,
isobutyric, valeric, isovaleric, 2-methylbutyric, hexanoic
and heptanoic acids have been reported as major sources
Table 1. *In vitro* gas production, estimated metabolizable energy (ME), short chain fatty acid (SCFA) and organic matter digestibility (OMD) of treated and forage supplemented cassava peels.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Feed materials</th>
<th>T1 (CSP + U)</th>
<th>T2 (CSP + BL)</th>
<th>T3 (CSP + CSF)</th>
<th>T4 (CSP + SPF)</th>
<th>Mean ±SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas volume, Gv (mL/200 mg DM)</td>
<td></td>
<td>18.17bc</td>
<td>21.17ab</td>
<td>31.90a</td>
<td>32.20a</td>
<td>25.86 ± 5.97</td>
</tr>
<tr>
<td>Metabolizable energy, ME (MJ/kg DM)</td>
<td></td>
<td>6.42b</td>
<td>6.56b</td>
<td>7.54a</td>
<td>7.67a</td>
<td>7.05 ± 0.31</td>
</tr>
<tr>
<td>Short chain fatty acids, SCFA (µM)</td>
<td></td>
<td>0.37c</td>
<td>0.45b</td>
<td>0.70a</td>
<td>0.71a</td>
<td>0.56 ± 0.05</td>
</tr>
<tr>
<td>Organic matter digestibility, OMD (%)</td>
<td></td>
<td>45.59b</td>
<td>46.21b</td>
<td>52.07a</td>
<td>52.80a</td>
<td>49.17 ± 1.97</td>
</tr>
</tbody>
</table>

Means bearing different superscripts along the same row are significantly different (P < 0.05). Metabolizable energy, ME (MJ/kg DM) = 2.20 + 0.136*Gv + 0.057*CP + 0.0029*CF. Organic matter digestibility, OMD (%) = 14.88 + 0.889*Gv + 0.45*CP + 0.0651*XA. Short chain fatty acids, SCFA (µM) = 0.0239*Gv – 0.0601. Gv = Gas volume; CP = Crude protein; CF = Crude fibre; XA = Ash.

Figure 1. Gas production trends of treated and forage supplemented Cassava peels.

of energy as well as building blocks for milk synthesis (Bergman, 1990). Thus, the T4 (CSP + SPF) and T3 (CSP + CSF) diets having recorded the highest value of SCFAs have better potentials to fulfil this role. Similarly, the results of the SCFAs for these diets have also been corroborated by the earlier ME values for these diets. This will, therefore, give a strong assurance for the provision of up to 80% of maintenance energy for WAD goats or any other ruminant when fed with this diet (Bergman, 1990).

The organic matter digestibility (OMD) of the different diets in order of decreasing values after 48 hours of incubation were 52.80, 52.07, 46.59 and 45.59% for T4 (CSP + SPF), T3 (CSP + CSF), T2 (CSP + BL) and T1 (CSP + U), respectively. The OMD value of the T4 (CSP + SPF) treatment was higher, although they were not significantly (p > 0.05) different from the T3 (CSP + CSF) treatment group. The T4 (CSP + SPF) and T3 (CSP + CSF) treatment groups recorded the highest values of OMD. The OMD values recorded in this study were higher than those recorded by Alasa et al. (2010) in their studies while determining the chemical composition and *in vitro* gas production of *Panicum maximum* intercropped with two cultivars of *Lablab purpureus*, but lower than the OMD values reported by Adewumi and Ajayi (2010) in their studies conducted to determine the replacement value of full fat neem fruit for corn bran using *in vitro* gas production techniques. Consequently, this study revealed that the cassava peels supplemented with the forages from cassava (T3) and sweet potato (T4) plants recorded the highest OMD after 48 hours of incubation. This may be attributed to the fact that, the forages have fermentable nitrogen and readily degradable cell wall fractions which would increase the substrates available to cellulolytic microbes with a consequent increase in digestibility (Van...
Soest, 1982; Adebowale, 1994). In addition, the estimated higher and better OMD values recorded in the present study corroborate the results of the dry matter digestibility (DMD) values earlier reported in the in vivo studies by Kalio et al. (2014) for the various treatment groups. The trend of the results revealed that better OMD and DMD values were recorded for the in vitro and in vivo digestibility trials, respectively, for T₄ (CSP + SPF) and T₃ (CSP + CSF) treatment groups compared to those of T₁ (CSP + U) and T₂ (CSP + BL) in the two trials. This supports the reports of Akinfemi et al. (2009) and Pashaei et al. (2010) who gave a positive relationship between results of in vivo and in vitro digestibility studies and further explained that, efficient laboratory methods should be reproducible and should correlate well with actually measured in vivo parameters. This is important because it can provide enough information on the nutritive value of the feedstuffs as opposed to the results of chemical analysis alone (Sallam, 2005).

Conclusion and recommendation

This study revealed that the cassava peels (CSP) supplemented with the forages from cassava and sweet potato plants recorded the highest gas volumes after 48 hours of incubation. Furthermore, these diets possess higher metabolizable energy (ME) values. The ME values recorded in the study were within the recommended ME values (range: 6 to 13 MJ/kg/DM) for an average diet, which can meet the daily energy requirements for small ruminants. Similarly, the cassava and sweet potato forage supplemented CSP recorded higher values for short chain fatty acids (SCFAs) compared to the urea and broiler litter supplemented CSP. This implies that these diets may be potential sources of energy as well as a building block for milk synthesis. The results of SCFAs of these diets corroborate the earlier ME values for these diets and can give a strong assurance for the provision of up to 80% of maintenance energy for sheep and goats. Also, the organic matter digestibility (OMD) values recorded for the cassava and sweet potato forage supplemented CSP suggest that they were better utilized by sheep and goats and therefore recommended for utilization in small ruminant animal feeding regimen.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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