Effects of ensiling cassava peels on some fermentation characteristics and growth performance of sheep on-farm

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ABSTRACT
This study determined the effect of drying or ensiling cassava peels on some conservation characteristics and growth performance of sheep. Fresh peels were either sun-dried to a DM of 904 g/kg or ensiled for 45 days for determination of some chemical and microbial characteristics, and growth performance of sheep. 45 Djallonké sheep were randomly assigned to three supplementary dietary treatments (Control and dried or ensiled) and fed for 70 days. Ensiling reduced the pH from 5.65 in the fresh peel to 4.15 compared to 6.15 in the dried peel. Crude protein (CP) increased from 45±0.44 g/kg DM in the fresh peel to 46±0.48 and 52±0.88 g/kg DM in the dried and ensiled peel, respectively. Reduction in neutral detergent fibre concentration was greater by ensiling than by drying. However, a greater (P = 0.001) reduction in HCN concentration was achieved by drying than by ensiling. Moulds were greater (P = 0.011) in the ensiled than dry peels. Average daily weight gain was higher (P = 0.031) for sheep offered the ensiled than the dried or Control diet. In conclusion, sun-drying was more effective at reducing HCN concentration whereas ensiling improved the CP content of cassava peels and growth performance of sheep.

Keywords: cassava peels; crop residues sheep; ensiling; growth performance; sun-drying

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Cassava is produced in almost all the ecological zones of Ghana. A survey by Seidu et al. (2012) showed that about five to six tonnes of cassava are processed in a day in each processing centre in Techiman and Wenchi in the Bono East and Bono regions of Ghana, respectively. These peels usually become an environmental nuisance in communities where they are generated in large quantities and are often left to rot or burnt (Akinfala & Tewe, 2002).

Cassava peels are however a rich source of readily fermentable carbohydrates and can be used to improve the energy concentration of tropical grass silage diets (Onua & Okeke, 1999). However, the utilization of cassava peels as animal feed is constrained by its high perishability (Tewe, 1992), high HCN concentration (Cardoso et al., 2005), and low protein content (Konlan et al., 2016). Hydrogen cyanide makes the peels bitter thus reducing their palatability and intake. It also interferes with the utilization of essential amino acids and the function of some enzymes (Enneking & Wink, 2000).

Apart from the reduction in palatability, high consumption of HCN can also result in respiratory and cardiovascular weaknesses or even death (Bolarinwa et al., 2016; Obadoni & Ochuko, 2001). Toxicity of HCN to ruminants is believed to be uncommon because of microbial detoxification of HCN in the rumen (Smith et al., 1991). This is because when cyanogenic glucoside is exposed to linamarase enzyme extraruminally during cassava processing, HCN is released and can be detoxified in the rumen but when unprocessed cassava peels are fed to ruminants, microbial fermentation may rather enhance HCN formation in the rumen thereby increasing the level of toxicity (Llewellyn, 2014).

Various methods have been used including sun-drying, soaking, grating and fermenting in an attempt to reduce cyanogenic glycoside, improve the CP content and increase their shelf-life. Sun-drying reduces HCN concentration by 86% whereas fermentation (ensiling) reduces it by 96% (Tewe, 1992). Ensiling may also increase the digestibility of the peels. Ensiling improved the in situ DM degradability of dried cassava peels at 24 h by 3% points from 70% to 73% (Asaolu, 1988).

It is believed that the use of appropriate methods of conserving cassava peels will help to reduce feed costs and improve the efficiency of smallholder farmers. This, therefore, informed the decision to situate the study on-farm with the direct involvement of smallholder sheep farmers to enable them learn first-hand the conservation of cassava peels into animal feed. This study, therefore, sought to investigate the effects of drying or ensiling on some conservation characteristics of cassava peels and the growth performance of Djallonké sheep on-farm.

**Materials and Methods**

**Study location**

Botingli (9.61 N 0.79 W) is a farming community located in the guinea savannah zone of Ghana. It is bordered to the east by Savulugu, the administrative district capital of the Savulugu-Nanton (9°24′N 0°28′W) district and to the South-east by Nanton. Botingli has a unimodal rainfall pattern (950-1200 mm) which starts in late April and ends in October. The dry season spans from November to April. Temperatures range from 33 to 39°C (day) to 20 to 25°C (night). During the harmattan, wild bushfires are rampant, destroying native pasture and crop residues left on the farm fields after harvest. Farmers keep cattle, sheep, goats and local poultry in an integrated management system with crop production. The commonest crops grown in the area include maize, groundnuts, cassava, yam and rice.
Livestock production is mainly through the semi-intensive system of management.

Experiment I: Fermentation characteristics of cassava peels conserved by ensiling or drying

Collection and processing of cassava peels

Only cassava farmers who also owned sheep were enlisted for this study. Immediately after harvesting, cassava tubers were transported home and peeled for the production of dried cassava chips. The fresh peels were collected and allowed to wilt (250-280 g/kg DM) for two days. The peels were then thoroughly mixed on a clean surface to ensure uniformity and then ensiled in mini and bag silos.

For ensiling cassava peels in mini silos, approximately 2.5 to 3.0 kg of cassava peels were packed into each of twenty duplicate labeled mini silos (10.4 cm diameter × 35.6 cm height) by manual pressing. Each labeled silo was weighed with its cap prior to being filled and immediately after sealing. The silos were stored at ambient temperature (20°C) and opened on 1, 3, 10, 15 and 45 days of ensiling as a means of monitoring the trajectory of fermentation in the bag silos. Triplicate silos were prepared and opened for each sampling day. The procedure for filling the silos involved randomly selecting one of three sets of triplicate silos (1, 2, or 3) for each sampling day and filling them with a 25-kg lot of fresh cassava peels. Details of the mini silo experimental procedure have previously been described (Addah et al., 2012). On each opening day (1, 3, 10, 15 and 45), the silage was sampled for determination of pH, lactic acid, HCN, and NH₃-N, and the enumeration of lactic acid bacteria, yeasts and moulds.

For ensiling, a subsample (day 0) of fresh cassava peels was also collected for the determination of some nutritional qualities of the peels prior to being conserved. On opening day (day 45), the ensiled peels were sampled for sensory evaluation and graded for their sensory quality characteristics such as colour, smell and texture, and nutritional quality such as neutral detergent fibre (NDF), acid detergent fibre (ADF), NH₃-N and crude protein.

The ensiled residue was subsequently dried for three days to a DM of 940 g/kg prior to being used to prepare two supplementary diets (Olafadehan et al., 2012). The diets were manually mixed every week and fed to sheep on-farm for 70 days. The supplementary diets consisted of either the 750 g/kg DM of the ensiled (silage diet) or dried (dried diet) cassava peels plus 25% DM each of whole cotton seed. The ingredient and nutrient composition of the diets are shown in Table 1. The cotton seed was treated with 1% salt solution and rubbed vigorously between the two palms to reduce the lint and to expose the seed coat and then sun-dried (816 g/kg DM). These were used to formulate the diet consisting of either dried or ensiled peel.

Experiment II: Growth performance

Management and feeding of experimental animals

A total of 45 sheep (4–6 months) with an initial weight of 11.9 kg (SD = 2.6) were sampled from nine communal pens at Botinligi in the Savelugu-Nanton Municipal Assembly in the Northern Region of Ghana. Five sheep (three males and two females) were allocated
to each pen, ear-tagged and dewormed with Albendazole (Hubei Guangren Pharmaceutical Technology Co. Ltd., Mainland, China) before the commencement of the experiment.

Each pen contained both experimental and non-experimental sheep. The sheep were provided with water continually. Every morning, the non-experimental flock in each pen was let out of the pen to graze on natural pasture whereas the experimental animals sampled for the study were detained in the pen and offered measured quantities of the supplementary diet from 08:30 am to 12:30 pm. After 12:30 pm, the animals were also let out to join their counterparts to graze on the field of natural pasture and the supplementary feed withdrawn. The animals returned to their pens at about 06:00 pm. A Control group of animals were given similar management regimes except that salt lick was provided in each pen and sheep were let out to graze from 08:00 am to 06:00 pm and did not receive any supplementary diet. The sheep were adapted to this feeding regime for seven days prior to data collection.

The pens were made of mud and wooden rails and were owned by a group of two to four farmers. The animals were provided with water continually. Salt lick was also provided in each pen. The dried and ensiled peels were used to formulate two diets that were manually mixed every week. The diets consisted of either the 750 g/kg DM of the ensiled or dried cassava peels plus 250 g/kg DM of whole cottonseed. The cottonseed was obtained from Wienco Cotton Industry Ltd., Tamale, Ghana. The seed was treated with 1% salt solution and rubbed vigorously between the two palms to reduce the lint and to expose the seed coat and then sun-dried (816 g/kg DM). This was then used to formulate a diet of three parts ensiled or dried peel to one-part cotton seed (3:1).

Each morning, the animals were offered measured quantities of the supplementary diet from 08:30 am to 12:30 pm. After 12:30 pm, the animals were then let out to graze in the open field. The animals returned to the pen to sleep at about 06:30 pm. A Control group of animals were given similar management regimes except that they were let out to graze from 08:00 am to 06:00 pm and did not receive any supplementary diet.

**Growth performance**

Each animal was weighed using a scale (Avery digital scale; Avery Weigh-Tronix, Minnesota, USA) at the beginning of the experiment and at the end of every two weeks. The initial liveweight (kg) per animal was subtracted from the final live weight (kg) per animal at the end of the experiment and the weight multiplied by 1000 to obtain final liveweight gain per animal in grams. This was then divided by the number of days (d) the experiment lasted to obtain the average daily weight gain in grams. The feed on offer and leftovers were weighed and sampled every 14 d until the end of the experiment which lasted for 70 d. Feed offered and leftover samples were used for determination of DM content and daily DM intake (DMI) per pen as feed offered on DM basis minus leftover feed on DM basis. Feed intake expressed as dry matter intake (DMI) was thus calculated as:

\[
\text{DMI (g/d)} = [\text{Fresh feed weight (kg)} \times \text{DM content of feed (g/kg)}] - [\text{Leftover feed (kg)} \times \text{DM content of leftover (g/kg)}] \text{[1]}
\]

**Laboratory analyses**

**Measurement of pH**

pH of duplicate samples of the fresh peel and those obtained on day 1, 3, 10, 15 and 45 of opening the mini silos in Experiment I was measured. About 10 g of the sample was
weighed (Sartorius Gottingen, Germany) into a clean beaker. About 135 mL of distilled water was added and blended for 40 seconds using a kitchen blender (Kenwood, China). The blended sample was then poured into another clean beaker and a pre-calibrated pH meter (Sartorius, Gottingen, Germany) was used to measure the pH of the blended sample.

**Proximate and fibre analysis**

Feed samples were collected every seven days and pooled together into a single sample every 14 d (n = 5) for subsequent laboratory analyses. Proximate analysis of the diets and ingredients was carried out according to the official methods of analysis described by AOAC (2005). All analyses were done in duplicates. Neutral Detergent and ADF were determined using procedures of Van Soest et al. (1991) with the aid of an Ankom 200 fibre analyzer (Ankom Technology Corp., Fairport, NY). The concentrations of NDF or ADF were determined using the formula:

\[
\text{% NDF or ADF} = \frac{100 \times W_1 \times C_1}{W_2} \quad \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots [2]
\]

where: \( W_1 \) = bag tare weight, \( W_2 \) = sample weight, \( W_3 \) = dried weight of bag with fibre after the extraction process, \( C_1 \) = blank bag correction (running average of final oven-dried weight divided by original blank bag weight)

**Determination of ammonia nitrogen**

A composite portion of the fresh, dry, or ensiled cassava peel obtained after 45 d of ensiling in mini silos was blended and filtered through two layers of cheesecloth and the filtrate was centrifuged for 15 min at 10,000 \( \times g \) (4 \( ^\circ \)C). The supernatant was collected for analysis of \( \text{NH}_3 \)N concentration. The supernatant (1.6 mL) was combined with 0.15 mL of 65% (wt/vol) trichloroacetic acid and analyzed by the phenol-hypochlorite method described by Broderick and Kang (1980). Colorimetric calibration and quantification of \( \text{NH}_3 \)-N were then done on a Spectrotroquuant Pharo 300 (J.P. Selecta, Spain) spectrophotometer at a wavelength of 630 nm at a sample concentration range of 0.00-3.00 mg/L.

**Hydrogen cyanide concentrations**

Total HCN (ppm) in the cassava peels was analyzed using the alkaline titration method as described by AOAC (1990) and Famurewa and Emuekele (2014). The peels (10 g) ball-ground to pass through No. 20 mm sieve was soaked in an equal volume (100 ml each) of a mixture of distilled water and orthophosphoric acid. The samples were each thoroughly mixed and stored at room temperature overnight. This was done to set free all bounded hydrocyanic acid. The resulting sample (mixture) was then transferred into a distillation flask and a drop of paraffin (antifoaming agent) was added. The flask was then fitted to another distillation apparatus and distilled. About 112.5 ml of the distillate was collected in the receiving flask containing 100 ml of distilled water and 0.25 g of sodium hydroxide pellets. The distillate was then transferred into a 125 mL volumetric flask and made up to mark with distilled water. About 4.0 mL of 5% potassium iodide was then added and titrated against 0.01M Ag \((\text{NO}_3)_2\). End-point was indicated by faint but permanent turbidity. The total HCN content in mg/kg was calculated as:

\[
\text{Total HCN content} = \frac{12.5 \times \text{TV}}{M} \quad \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots [3]
\]

where: \( \text{TV} \) = titre value, \( M \) = mass of sample

**Microbial analysis**

On each day of opening, each duplicate mini-silo was opened and subsampled and prepared for use in enumeration of lactic acid-producing bacteria (LAB) using lactobacilli MRS agar and in enumeration of...
yeasts and moulds using Sabouraud’s Dextrose Agar (SDA). For enumeration of LAB, 62 g of the MRS agar powder was measured into a clean flask and 1L of distilled water added and dissolved by boil using a magnetic rod on a magnetic stirrer (J.P. Selecta, Spain) and then autoclaved (Microclave, J.P. Selecta, Spain) at 121°C for 15 minutes. The agar was then allowed to cool in a water bath (50°C). The agar was poured into sterilized Petri dishes to about half-full on a laminar flow hood (Envair, Haslingden, UK) under aseptic conditions. The plates were allowed to cool and set to room temperature. A similar protocol was used for the enumeration of yeasts and moulds except that 65 g of the SDA was added to 1 L of distilled water.

About 10 g of the sub-sampled dried or ensiled cassava peel was weighed and added to 90 mL of distilled water in a ziplock bag (17.7 cm x 18.8 cm) and vigorously agitated to detach the microbes from the sample. Serial dilutions of -2 to -5 were made and 1 µL of each dilution (-2, -3, -4 and -5) was used to inoculate each prepared MRS and SDA agar plates. For the MRS, the plates were placed upside-down in an incubator (Incubator-Co2, J.P. Selecta, S.A., Barcelona, Spain) at 32°C. Counting of bacteria or fungal colony-forming units (CFU) was done after 24 h for MRS media plates and 48 h for SDA plates using a digital colony counter (J.P. Selecta, S.A., Barcelona, Spain).

Sensory evaluation of silages

On the day of opening, 12 farmers were randomly selected and trained on the sensory evaluation criteria before administering a questionnaire to each. Sensory evaluation procedure was based on methods previously described by Jianxin (2002) and Jian et al. (2015). The farmers scored the silages for smell (1–15), colour (1–15) and texture (1–10). The silages were graded by summing the scores for each trait as excellent grade (40–31), good grade (30–21), general grade (20–11) and low-grade (≤10).

Statistical analysis

Data were analyzed using the PROC Mixed procedure (SAS Inst. Inc., Cary, NC). Data on microbial populations were transformed to log10 colony-forming units prior to statistical analysis with each mini silo as the experimental unit and the bulk storage container as the experimental unit for the dried peel. For the growth performance study, data on DMI and growth performance of sheep (body weight gain and ADG) were analyzed for the effects of ensiling as a completely randomized design with initial body weight as a covariate in the model and pen as the experimental unit. Differences in least squares means of all fixed effects were declared statistically significant at $P \leq 0.05$.

Results

Fermentation characteristics of cassava peels

There was an indication of a good trajectory of fermentation as the pH declined rapidly (Fig. 1) from 6.2 in the unensiled peel to 4.1 in the ensiled peel and 6.1 in the dry peel after 45 d of ensiling (Table 1). The populations of LAB and moulds were greater ($P \leq 0.04$) in the ensiled compared to dried cassava peels (Table 1). However, the yeast population was similar ($P = 0.474$) between the dried and ensiled peel. Ammonia N concentration was 13% higher in the ensiled than the dried peels even though the difference was not significant ($P = 0.229$). Generally, ammonia N concentration increased from 0.25 mg/kg in the fresh peel to 1.12 mg/kg after 45 d of ensiling and 0.48 mg/kg after drying (Table 1). The concentration of HCN was reduced from 27.4 mg/kg in the fresh peel to 25.0 mg/kg in the ensiled and 18.0 mg/kg after drying.
TABLE 1
Effects of conserving cassava peels by ensiling or sun-drying on some conservation characteristics after 45 d of ensiling in bag silos

| Item                              | Fresh peel | Dried    | Ensiled   | SEM   | P value       |
|-----------------------------------|------------|----------|-----------|-------|---------------|
| pH                                | 6.1        | 6.2<sup>a</sup> | 4.2<sup>b</sup> | 0.029 | <.0001        |
| Ammonia N (mg/kg DM)              | 0.25       | 0.48     | 1.12      | 0.365 | 0.229         |
| Hydrogen cyanide (mg/kg DM)       | 27.37      | 18.0<sup>a</sup> | 25.0<sup>b</sup> | 0.145 | 0.001         |
| Lactic acid bacteria (CFU/kg DM)  | nd         | 5.4<sup>a</sup>  | 6.4<sup>b</sup>  | 0.256 | 0.014         |
| Yeasts (CFU/kg DM)                | nd         | 6.5      | 6.7       | 0.272 | 0.474         |
| Moulds (CFU/kg DM)                | nd         | 5.4<sup>a</sup>  | 6.4<sup>b</sup>  | 0.164 | 0.011         |

CFU: Colony-forming units; nd: not detectable (at dilution -1).
Least square means with different superscripts within rows differ significantly (P < 0.05).

Fig. 1: Effect of conserving cassava peels by ensiling or drying on the trajectory of pH decline

Sensory characteristics of cassava peels
From responses collated from the questionnaires administered to selected farmers on sensory evaluation of ensiled cassava peels, the colour of the silage ranged from light brown to deep brown, the silage had a pleasant smell and a very firm texture generally rated as good quality (Table 2). No great difference in sensory characteristics among silages from the three bulk silos was observed. The sensory quality of the silage indicated that it was of good quality grade (27.7±0.72; Table 2).

TABLE 2
Sensory characteristics of cassava peel silage after 45 d of ensiling in large bag silos

| Item      | Description          | Score<sup>1</sup> |
|-----------|----------------------|--------------------|
| Colour    | Light brown          | 9.22±2.24          |
| Smell     | Very pleasant        | 10.50±2.46         |
| Texture   | Firm                 | 8.00±2.10          |
| Grade<sup>2</sup> | Good               | 27.72±0.72         |

<sup>1</sup>Values are means of three large silos (n = 3)
<sup>2</sup>The silages were graded by summing the scores for colour (1–15); texture (1–10); smell (1–15) as excellent grade (40–31); good grade (30–21); general grade (20–11) and low-grade (≤10) according to Jianxin (2002) and Jian et al., (2015).

Proximate composition of cassava peels and diets
The NDF concentration of cassava peels decreased from 448±11.0 g/kg DM in the fresh peel to 418±15.1 g/kg DM and 364.1±7.6 g/kg DM in the dried and ensiled peel, respectively. On the contrary, ADF increased from 133±1.2 g/kg DM in the fresh to 139±0.65 g/kg DM and 190±1.5 g/kg DM in the dried and ensiled peels respectively. Ensiling also increased the CP concentration from 45±0.44 g/kg DM in the fresh peel to 46±0.48 g/kg DM in the dried peel and 52±0.88 g/kg DM in the ensiled peel. The trend of the CP and NDF concentrations of the ensiled and dried peel was consistent with
the concentration of these constituents in the respective diets (Table 3).

TABLE 3

| Item (g/kg DM)          | 1Cassava Peels | 2Diet |
|------------------------|----------------|-------|
|                        | Fresh | Dried | Ensiled | Dried | Ensiled |
| Dry matter             | 945±5.6 | 941±7.6 | 945±6.7 | 944±8.5 | 950±6.1 |
| Crude protein          | 45±0.44 | 46±0.48 | 52±0.88 | 103±5.0 | 126±4.2 |
| Ether extract          | -     | -     | - | 42±4.8 | 74±5.5 |
| NFE                    | -     | -     | - | 646±9.4 | 559±9.6 |
| Ash                    | 43.6±1.3 | 46±1.8 | 45.3±0.1 | 44±1.5 | 44±2.4 |
| NDF                    | 448±11.0 | 418±15.1 | 364.1±7.6 | 463±16.5 | 325±4.5 |
| ADF                    | 133±1.2 | 139±0.65 | 190±1.5 | 216±7.1 | 232±15.4 |
| NH₃-N (mg/kg DM)       | 0.25 | 0.48±0.01 | 1.12±0.51 | 0.58±0.01 | 0.35±0.06 |
| ³ME (MJ/kg DM)         | - | - | - | 12.5 | 12.7 |

1Cassava peels were either analyzed as fresh, dried (6 d) or ensiled (45 d) in 3 large polyethylene-lined bag silos
2Diet contained cassava peels (dried or ensiled) and whole cotton seed in a ratio of 3:1.
3ME: Metabolizable Energy (MJ/kg DM; Pauzenga, 1985).

Animal growth performance

There was no significant difference ($P = 0.637$) in the initial weight of sheep among the three treatments. There was a significant difference between animals fed the peel-based diets (ensiled or dried) compared to those on the Control group in terms of total weight gained ($P = 0.040$) and final weight attained ($P = 0.028$). However, the growth performance of sheep fed the ensiled or dried cassava peel-based diet did not differ significantly ($P > 0.05$). Dry matter intake of the diets did not differ ($P = 0.436$) between sheep receiving the dried or ensiled supplementary diet even though those in the ensiled group had higher numerical values. Supplementation of sheep with ensiled and dried cassava peels however resulted in higher ($P = 0.031$) average daily weight gain compared to the Control. Ensiling also resulted in higher daily gain than drying even though the difference between the ensiled and the dried were not significant ($P = 0.293$). The growth pattern of the sheep showed that the growth performance of the sheep in the Control group was consistently inferior to those offered the dry and ensiled peels throughout the 70-day feeding period (Fig. 3).

TABLE 4

| Item                         | Dried | Ensiled | Control | SEM | $P$ value |
|------------------------------|-------|---------|---------|-----|-----------|
| Initial weight (kg)          | 12.6  | 11.9    | 11.3    | 1.06| 0.637     |
| Final weight (kg)            | 18.0* | 18.7*   | 16.0*   | 0.55| 0.040     |
| Weight gain (kg)             | 6.1*  | 6.9*    | 4.0*    | 0.53| 0.028     |
| DM intake (g/d)              | 180   | 230     | –       | 0.04| 0.436     |

SEM: Pooled standard error of least square means (n = 6)
Discussion

The reduction in the concentration of NDF during ensilage is suggestive of the solubility of NDF in cassava peels. Hemicellulose and cellulose constitute the digestible fractions of NDF. Hemicellulose contains arabinoxylans and glucuronoarabinoxylans. These can be hydrolysed by the pentose-phosphate pathway of fermentation by heterolactic lactic acid bacteria during ensiling resulting in a reduction in NDF. In previous studies, ensiling cassava peels reduced NDF by 25% compared to sun drying (Olafadehan et al., 2012).

As expected, pH of the ensiled peel was lower than the dried peel and was consistent with the high concentration of lactic acid and the increase in the population of lactic acid-producing bacteria. Lactic acid bacteria are the main micro-organism responsible for the fermentation of sugars to lactic acid. Even though Kung and Shaver (2002) suggested that tropical silages should have a pH of 4.3–4.7 to be graded as good silage, this benchmark cannot be directly applied to tropical crop residues. At harvest, tropical forages especially crop residues have lower water-soluble carbohydrates and high cell wall concentration. These conditions together with higher environmental temperatures in the tropics give the less desirable bacilli a competitive advantage over lactic acid-producing bacteria which are rather more efficient converters of water-soluble carbohydrates to lactic acid (Oude Elferink et al., 2000). A minimum pH of 4.4 has been used as a benchmark for classifying cassava peel silage as being good (Asaolu, 1988) but Meneses et al. (2007) recommended a pH of 3.5–5.5 as an acceptable threshold for good silage made from crop by-products.

A pH of 4.1 observed in this study was associated with good sensory quality characteristics including light-brown colour and a pleasant odour (Table 2), an indication of the less proliferation of saccharolytic spoilage organisms during ensilage. Figure 1 indicates a faster rate of pH decline during fermentation. The rate of pH decline influences the quality of the silage. A slower rate of decline provides more time for the growth of undesirable enterobacteria that grow well at a pH greater than 5 (Schroeder, 2004). The concentration of NH$_3$-N was 13% higher in the ensiled peel compared to the dried peel even though the difference was not significant. Ammonia N is a by-product of proteolytic degradation of protein and is undesirable in the production of good quality silage. The concentration
of NH$_3$-N was higher in the ensiled than dried peel but a reverse trend was the case in the respective diets; the dried peel diet had higher NH$_3$-N concentration than the ensiled peel (0.58 vs. 0.35 mg/kg). Though this may be puzzling, the drastic increase in NH$_3$-N concentration in the dried peel diet could be attributed to the proteolytic activity of yeasts and moulds during storage and feeding. In this study, diets were sampled every 7 d for chemical analysis until the end of the study. Such secondary fermentation in cassava peels has been described as “dry fermentation” and is reported to occur when the peels are improperly stored prior to use (Tewe, 1992). We, therefore, suspect that the dry peels in our study was not properly stored.

The number of moulds was increased by ensiling than drying. The principal factor affecting aerobic deterioration of silage is the population of yeasts that initiate the deterioration. A population threshold of 5 log CFU/g DM has been established for silage to undergo deterioration (Woolford, 1990). Our values were above this threshold meaning our material may have been undergoing deterioration. The ensiled peel generally smelled pleasant, had firm texture with a light brown to brown colour which agrees with a study by Asaolu (1988) who reported that good quality silage is produced when cassava peels silage is light brown in colour, firm in texture with a pleasant odour and a pH of 4.4. The colour of the silage is close to the colour of the original fresh peels hence the silage is considered to be of good quality. Silage should be preserved and be close to the colour of the original fresh forage for it to be considered as good quality silage (Oduguwa et al., 2007; Liu & Guo, 2002).

The population of moulds was increased by ensiling than drying. The increased population of these microbial cells possibly accounted for the higher CP in the ensiled compared to dried peel. The sensory characteristics and rapid decline in pH of the ensiled peel indicated a good trajectory of fermentation. Putrefied silage has a dark colour, recognizable plant parts (bad texture) and a rancid and nauseous smell (Jianxin, 2002).

A further indication that the silage was well-fermented is the fact that ensiling did not depress intake of the supplementary diet. A comparison between silage and hay conserved from forage of the same maturity suggest that voluntary intake of silage is usually about 17% lower than hay (Thiago et al., 1992). This suggests that the end products of the ensiling process may limit intake. The reduction in intake is principally influenced by the end-products of fermentation via their pre-ingestive effects on palatability (Buchanan-Smith, 1990) or post-ingestive effects on ruminal VFA concentrations (Oba & Allen, 2003).

Hydrogen cyanide imposes bitterness on the peel that may reduce palatability. The intake of the diet containing the ensiled peel was 28% greater than that containing the dried peel even though the difference was not significant. The HCN concentration of the dried peel was lower than that of the ensiled peel but ADG of sheep supplemented with the ensiled peel were superior to those supplemented with the dried peel and those on the Control. In general, the depression in growth performance of animals fed cassava-based diets has been attributed to the interference of functions of metal-containing enzyme systems responsible for cell respiration and function by HCN in cassava-based diets. On contrary, however, in pigs, N digestibility and retention have been shown to be greater for pigs fed ensiled cassava leaves diet than for those fed dry leaves diet despite the latter being higher in HCN concentration (Phuc et al., 1996). Tweyongyere & Katongole (2002) examined three methods of HCN
detoxification of cassava peels; sun-drying, fermentation and soaking. They concluded that sun-drying was the most effective method and resulted in the most rapid reduction of HCN; more than 82% reduction in 48 h.

Sun-drying cassava leaves reduced HCN content to 22.5 mg/kg compared to 147 mg/kg for the ensiled material (Phuc et al., 1996). Phuc et al. (1996) therefore concluded that drying may be more effective at removing HCN than ensiling. Other studies however showed that ensiling reduced free HCN of the peel by 36% (Gomez & Valdivieso, 1988) to 98% (Tewe, 1992) compared to a reduction of 82% (Tweyongyere & Katongole, 2002) to 85% (Gomez et al., 1984) by sun-drying. In these latter studies, the greater reduction due to ensiling was generally attributed to the combined effect of lower pH (<4.4) and intense heat generated in the silo during ensiling. Slower rate of sun-drying (1-3 d) in the rainy season removes bound HCN more effectively than other artificial methods of detoxifying cassava peels because apart from the slow generation of heat during sun-drying, “dry fermentation” occurs with cassava peels conserved by sun-drying. This phenomenon further reduces the HCN concentration due to enhanced hydrolysis of cyanogenic glucosides by linamarase caused by slow heating during sun-drying (Famurewa and Emuekele, 2014; Lukuyu et al., 2014).

It has been suggested that improvements in growth performance of sheep fed ensiled compared to dried cassava peels diets could be due to the improvement in CP content and rumen degradability (Asaolu, 1988) rather than reduction in HCN. Indeed, the HCN concentration in this study was not beyond the level of 30 mg/kg recommended to cause depression in growth performance or death (Tweyongyere and Katongole, 2002). The increased populations of bacteria and fungi during fermentation have been reported to increase the microbial protein of the silage (Oboh, 2006). This could account for the higher concentration of CP observed in the silage and silage diets and subsequently, the improved growth performance of sheep fed the ensiled cassava peels diet. The ADG of sheep supplemented with the dried and ensiled cassava peels in the present study compares favourably with those of Asaolu (1988) who fed two groups of sheep with diets containing 80% each of dried and ensiled cassava peels. In that study, ADG for sheep fed 80% ensiled cassava peels was 0.08 kg/d compared to 0.06 kg/d for the dried peel.

The poorest growth performance of sheep in the Control was expected. The quality and quantity of feed resources in the northern parts of Ghana decline greatly during the dry season often necessitating supplementation to avoid weight loss.

**Conclusion**

Conserving cassava peels by sun-drying was more effective at reducing HCN concentration compared to ensiling but the nutritional quality of the peel, in terms of CP content, was improved by ensiling than by sun-drying. This study further indicates that supplementation of sheep with the ensiled peel improves the growth performance of sheep on-farm than supplementation with the dried peel or not providing supplementary feed at all during the dry season. This study therefore recommends conservation of cassava by ensiling in the wet season and sun-drying in the dry season.

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