Analytical Methods

Simple method to reduce the cyanogen content of gari made from cassava

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The lactic acid content of gari, was determined by pH titration to be about 10 g lactic acid/kg gari. As the size of the gari particles increased from 400 to >1000 μm their total cyanide content increased from 5 to 21 ppm. The acetone cyanohydrin content of gari samples exposed to ambient laboratory conditions of temperature and relative humidity gradually decreased by 58% in 38 weeks whereas linamarin in cassava flour is 100% stable for 6 months. Cyanogens could not be removed from wet gari samples at 30 or 50 °C and were only slowly removed by repeated heating at 100 °C and rewetting. By mixing equal weights of gari (pH 4.1) and low cyanide cassava flour (pH 6.5), wetting and heating at 50 °C for 5 h, the cyanide content was reduced by about one half. Wetting a gari/flour (1:1) sample with 1.5 times its weight of water and standing in a 1 cm thick layer in the sun for 4 h reduced the cyanide content by about one half. This treated mixture may be cooked to prepare stiff porridge with lowered cyanide content, and may help reduce tropical ataxic neuropathy in West Africa.

1. Introduction

Cassava is the staple food of tropical Africa and its production and consumption is increasing rapidly in order to feed a rapidly increasing population (Nhassico, Muquingue, Cliff, Cumbana, & Bradbury, 2008). Cassava is easy to propagate from stem cuttings, yields well in poor soils without fertilizer and is drought resistant, because the plant is sustained during drought by its tuberous roots, which are also a reserve source of food. The roots are very starchy and the leaves are a good source of protein. Cassava produces two cyanogenic glucosides, linamarin and a small amount of lotaustralin (methyl linamarin) that are hydrolysed to give a cyanohydrin and glucose, catalysed by linamarase. The cyanohydrin decomposes spontaneously above pH 5 (White, McMahon, & Sayre, 1994), to liberate poisonous hydrogen cyanide which is a danger to human health.

A new processing method to remove cyanogens from cassava flour involves mixing dry flour with water and leaving the wet flour in a thin layer in the shade for 5 h or for 2 h in the sun to allow the catalysed breakdown of linamarin to hydrogen cyanide. The wet flour is used for cooking the same day. This gives a three- to sixfold reduction in total cyanide content of cassava flour (Bradbury, 2006; Bradbury & Denton, 2010; Cumbana, Mirione, Cliff, & Bradbury, 2007). The method was successfully field-tested in Mozambique and is being introduced also in Tanzania (Nhassico et al., 2008) and Democratic Republic of Congo (Karumba, Denton, & Bradbury, 2009).

In West Africa cassava is processed predominantly by grating the peeled root and allowing fermentation in a sack for about 3 days, followed by expression of the cyanogenic liquid and heating with stirring in a metal pan over a wood fire to give a gritty, roasted product called gari. This removes much more cyanogens than the various methods used to produce cassava flour in Eastern, Southern and Central Africa (Cardoso et al., 2005). Thus, the average total cyanide content of cassava flour in a good season in Mozambique is about 45 ppm (Cardoso, Ernesto, Cliff, Egan, & Bradbury, 1998) and of gari in West Africa is about 20 ppm (Cardoso et al., 2005). The WHO safe level for cyanogens in cassava flour is 10 ppm (FAO/WHO, 1991).

Under drought conditions the total cyanide content of flour is more than doubled and this leads to cyanide poisoning and outbreaks of konzo (Howlett, Brubaker, Mlingi, & Rosling, 1990) in various countries of Eastern and Central Africa. Tropical ataxic neuropathy (TAN) occurs in West Africa (particularly Nigeria), Tanzania, Uganda, Kenya, West Indies and South India (Osuntokun 1981, 1994) and probably results from continuous intake over years of cyanogens from a monotonous diet of cassava. TAN is a progressive neurological disease that causes unsteady walking, loss of sensation in hands, blindness, deafness and weakness. Recent studies have thrown doubt on the linkage of TAN with long term cyanide intake from cassava (Oluwole et al., 2003), but recent work from India (Madhusudanan, Menon, Ummer, & Radhakrishnan, 2008) supports earlier studies by Osuntokun (1994) of a linkage.

The major cyanogen present in gari is acetone cyanohydrin (Bradbury, 2009; Onabolu, Oluwole, Rosling, & Bokanga, 2002), whereas that present in cassava flour is linamarin. Since the total cyanide content of cassava flour is reduced three- to sixfold using...
the wetting method (Bradbury, 2006; Cumbana et al., 2007) it might be expected that a similar reduction would occur with gari. Unfortunately the pH of gari is about 4.1, due to lactic acid fermentation that occurs during processing (Obilie, Tano-Debrah, & Amoateng-Awua, 2004), and acetone cyanohydrin is considered to be stable under these conditions (White et al., 1994). In this paper we have initially studied various properties of gari and different possible methods to remove cyanogens from gari. These initial studies showed the way forward to development of a method by which the pH of the gari is increased by mixing with flour, wetting with water and drying in the sun. This reduces the total cyanide content of the gari/flour mixture by one half. If implemented in West Africa, this would reduce the cyanide intake of people consuming gari and may reduce the incidence of TAN.

2. Materials and methods

2.1. Materials

Five different samples of commercial gari were obtained from local markets in Maputo, Mozambique, and were stored in a deep freeze at −20°C, where they are stable indefinitely. Cassava flour samples were produced from cassava roots grown in a glass house in Canberra, some of which had been used in earlier studies (Bradbury, 2006). Thermochron iButton, model DS1922L-F5 was used as a temperature data logger with a range of temperatures −40 to 85°C, and produced by Embedded Data Systems, Dallas, Texas, USA.

2.2. Methods

2.2.1. Initial measurements of the properties of gari samples and cyanogen loss from gari

2.2.1.1. Measurement of acid content of gari. A sample of gari 2 (10 g) was dispersed in 50 mL water and titrated with 0.100 M NaOH in a 250-ml beaker fitted with a magnetic stirrer using a Beckman pH metre. The pH metre was calibrated with standard buffers at pH 4.00 and 7.00. The pH increased slowly after addition of each 1.00 mL of NaOH from a burette and became constant after <1 min. Titration was continued up to pH 9 and was repeated with gari 5.

2.2.1.2. Total cyanide content of sieved fractions of gari. Samples (10 g) of gari 2, gari 3 and gari 5 were sieved separately by hand using a stack of Endicott sieves of sizes 425, 850 and 1000 μm into fractions <425 μm (very fine), 425–850 μm (fine), 850–1000 μm (medium) and >1000 μm (coarse). The weights of each of the fractions was recorded. In the coarse fraction there were a number of much larger pieces, irregular aggregates of 3–10 mm diameter. These aggregates were picked out using tweezers from the coarse fraction. For total cyanide analysis, duplicate 100 mg samples were placed in plastic vials, a linamarase/buffer paper was added, followed by 1 mL of 1.0 M phosphate buffer at pH 6.8, a sensitive picrate paper and a lid. The vial was maintained at 30°C overnight, the 1 cm² picrate paper was separated from the plastic strip and the paper eluted for 30 min with 0.50 mL of distilled water. The absorbance (A) of the solution was measured using 2 mm cuvettes in a Beckman DU 540 spectrometer in the direct reading mode, against a blank solution prepared from a 1 cm² picrate paper not exposed to HCN and eluted with 0.50 mL water. The total cyanide content in mg HCN equivalents/kg sample = ppm was calculated by the equation, ppm = A × 45.7 (Bradbury, 2009).

2.2.1.3. Loss of cyanide from gari samples under ambient conditions. Samples (about 5 g) of each of the five gari samples were placed in small plastic dishes at ambient conditions of temperature and relative humidity, exposed to the air for 38 weeks and the total cyanide content monitored as a function of time, see Section 2.2.1.2.

2.2.1.4. Attempted loss of cyanide from gari with multiple wetting/ heating at 100°C. Two gari 5 (2 g) samples were taken in weighed 100 mL beakers, 2 mL of water added to 1 and 2 mL of 1 M phosphate buffer (pH 6.9) to the other. After mixing, duplicate samples were withdrawn for total cyanide analysis and the beakers weighed and heated for 30 min at 100°C in an oven. The beakers were removed, weighed, water added with mixing to give the same weight as before the heating, and duplicate 100 mg samples taken for total cyanide analysis. The beakers were again weighed, heated at 100°C for 30 min and the process repeated a second time. This process was repeated five times in all making a total heating time at 100°C of 2.5 h. At the end of the cycle 0.5 g of the wet flour samples were mixed with 2 mL water, the pH of the solutions measured and found to be 4.3 and 6.9, respectively.

2.2.1.5. Loss of total cyanide from wet gari samples with or without pH adjustment. Gari (2 g) was mixed with either 2 or 3 mL of water or with 3 mL of 1.0 M buffer in a weighed 100 mL beaker. Above pH 6, 1.0 M phosphate buffer was used and for pH 4–6, 1.0 M citrate buffer solutions were used. These were prepared from a 1 M citrate buffer by adjustments of pH using conc. HCl or NaOH solutions as required. Duplicate 100 mg samples were taken for total cyanide analysis and a 1.0 g sample of wet gari was added to 2 mL water for pH measurement. The wet gari sample was heated in an oven at 30 or 50°C for 5 h, after which water was added with mixing to give the same weight as at the beginning. Duplicate 100 mg samples were removed for total cyanide analysis (Section 2.2.1.2) and a 1.0 g sample for pH measurement.

2.2.2. Loss of cyanide from wet gari, gari/flour and flour samples

Gari (1 g) was mixed with flour (1 g) and 3 mL water was added with mixing in a weighed 100 mL beaker. Samples of flour (2 g) or gari (2 g) were also mixed with 3 mL water in a weighed 100 mL beaker. Duplicate 100 mg samples were withdrawn for total cyanide analysis and 0.5 g for pH measurement. The mixed wet sample was heated at 30 or 50°C in an oven for 5 h, the beaker weighed and water added to replace that lost by evaporation and mixed well. Duplicate 100 mg samples were taken for total cyanide analysis and 0.5 g for pH measurement (Section 2.2.1.4).

2.2.3. Cyanide loss from wet gari and gari/flour samples treated in the sun

Gari and gari/cassava flour (1:1) samples were placed in small weighed glass Petri dishes, mixed dry and wetted with 1.5 times their weight of water. Sample sizes were either 5 g gari, 5 g flour and 15 g water or 1 g gari, 1 g flour and 3 g water. After mixing, duplicate 100 mg samples were withdrawn for cyanide analysis (see Section 2.2.1.2). The Petri dish was placed on a concrete slab in the sun for 4 h. In some cases an iButton was placed in the dish to monitor the temperature of the wet flour over the course of the experiment. The iButton was either directly exposed to the sun or just covered by about 5 mm of the wet flour. After 4 h the sample was weighed, water added to replace that lost by evaporation with mixing and duplicate 100 mg samples taken for cyanide analysis and 0.5 g for measurement of pH. In several experiments the weighed gari/flour samples were heated in the sun for 2 h followed by addition of water and then heated for another 2 h in the sun followed by addition of water, mixing and cyanide analysis, but this gave similar results to one period of heating for 4 h. Experiments were also made with two UV sources, a Sunlamp and a Spectrolin-
ker Model XL-1500 UV crosslinker (Spectronics Corporation), operating at 254 nm.

3. Results and discussion

Since the aim of this study was to develop a simple method to remove cyanogens from gari, it was necessary to do a number of initial experiments on the properties of gari and on possible methods to remove cyanogens from gari. These initial experiments (see Sections 2.2.1.1 to 2.2.1.5), carried out on one or more of the five gari samples, showed the way forward and allowed development of the final method, which was checked out with all five gari samples and two flour samples of low total cyanide content.

3.1. Acid content of gari

Cassava flour samples dispersed in water have a pH of about 6.5 and are buffered by citrate, malate and succinate anions. The total amount of these anions in fresh cassava roots (which contain ca. 63% moisture) is about 10 g/kg fresh root (Bradbury & Holloway, 1988), and hence the amount present in cassava flour made by sun drying (which contains ca. 9% moisture) would be ca. 24 g/kg fresh flour. In gari the pH has been reduced to 4.1–4.2 by lactic acid fermentation during preparation (Obilie et al., 2004; Sokari & Karibo, 1996). By titration of gari with NaOH up to pH 6.5, one can calculate the amount of lactic acid in the gari sample.

Gari 2 and gari 5 were titrated with 0.1 M NaOH (see Section 2.2.1.1). There was a slow increase of pH from 4.1 up to 5.4 followed by a more rapid increase of pH up to 6.5 and then another slow increase up to pH 7.0. The titration was continued up to pH 9. The 10 g samples of gari 2 and gari 5 in water required 10.0 and 12.6 mL of 0.100 M NaOH, respectively, to raise the pH to 6.5. This corresponds to 9.0 g lactic acid/kg gari 2 and 11 g lactic acid/kg gari 5. On average the lactic acid content of gari is about 10 g/kg gari, which is much less than the total value of the other organic acid anions present in cassava flour and probably also in gari.

3.2. Size distribution and total cyanide content of gari particles

Gari was separated into fractions by sieving and each was analysed for total cyanide content (see Section 2.2.1.2). Similar results were obtained for each gari sample and mean values and standard deviations are recorded in Table 1. The bulk of the gari (62.6%) is in the coarse fraction (>1000 μm) and includes aggregates. Aggregates are particles that were probably fused together during the final roasting of the gari, that involves heating in a metal dish over a wood fire and stirring with a wooden spoon. This roasting volatilises water and hydrogen cyanide (HCN) from the wet solid, partially decomposes acetone cyanohydrin to HCN and also gelatinises some starch granules at 60–70 °C, which also fuses particles together to form aggregates. The roasting process gives gari a gritty, hard surface and it is quite difficult to grind it up into finer particles in a pestle and mortar, in contrast to cassava flour which is soft and fluffy and easy to grind.

It is noteworthy that the total cyanide content of gari particles increases with increasing particle size, such that the cyanide content of aggregates is about 4.5 times as large as that of the very fine particles. This is probably due to residual acetone cyanohydrin and HCN escaping most easily from the very fine particles and not so readily from the gelatinised coarse particles and aggregates. With cassava flour particles there is no difference in cyanide content between coarse and fine flour particles (Bradbury, 2006).

3.3. Loss of cyanide from gari under ambient conditions

The total cyanide content was followed under ambient laboratory conditions of temperature and relative humidity over a period of 38 weeks for the five gari samples. Acetone cyanohydrin (85%) is the major cyanogen present and the remainder is linamarin (Bradbury, 2006). The percentage total cyanide remaining for each gari sample was averaged and the results are shown in Fig. 1. There is a steady slow decline in total cyanide content with time and 42% still remains after 38 weeks. A moisture determination on the five gari samples by oven drying at 80 °C to constant weight, gave 9.5% moisture. This loss of cyanide is very much less than that obtained by Onabolu et al. (2002), who found complete loss of cyanogen in 4 weeks at a slightly higher moisture content, 8.7–13%. Our ambient laboratory temperature averaged about 20 °C, which was lower than the tropical conditions of Onabolu et al. (2002). The slightly higher moisture content and higher temperature and relative humidity used by Onabolu et al. (2002) may explain the difference in results between their work and ours. We conclude that the acetone cyanohydrin remaining in gari at pH 4.1–4.2 is surprisingly stable (42% remaining after 38 weeks), but is much less stable than linamarin in cassava flour, where 100% remained after 6 months (Bradbury, 2006).

3.4. Effect of temperature and pH on cyanide removal from wet gari

The data of White et al. (1994) showed increased instability of acetone cyanohydrin as temperature is increased. Accordingly, two gari samples were wet with water which gave a pH of 4.3 and 1.0 M buffer which gave a pH of 6.9, heated at 100 °C for 30 min and then wet with water and reheated again four times over, see Section 2.2.1.4. The results in Fig. 2 show a slow linear drop in total cyanide content in gari at pH 4.3. This could perhaps be due to evaporation of acetone cyanohydrin (BP 82 °C at 25 mm Hg pressure) from the wet gari, but is more likely due to decomposition of acetone cyanohydrin to acetonitrile and HCN at 100 °C. The very slow loss of acetone cyanohydrin from wet gari at 100 °C at pH 4.3 shows that swelling with water and elevated temperature is not a good method to remove acetone cyanohydrin from wet gari.

However, increasing the pH of wet gari to 6.9 with phosphate buffer at the beginning of the experiment, produced a 75% loss of cyanide in a single 30 min treatment at 100 °C (Fig. 2). These initial experiments show that acetone cyanohydrin in gari is removed at 100 °C if the pH is increased from 4.3 to 6.9, and in the next section we study in detail the effect of pH.

3.5. Effect of pH on cyanide loss from wet gari

Gari samples were mixed either with water to obtain pH 4.2 or with 1.0 M buffers of pH from 4.5–6.8 and heated at 30 or 50 °C for 5 h, with water replacement/mixing at the end of the heating period. The loss of total cyanide was monitored and pH checked (Section 2.2.1.5). Both curves in Fig. 3 show the reduction in % total...
Fig. 1. % Mean total cyanide remaining in five gari samples exposed to ambient laboratory conditions vs time in weeks.

Fig. 2. % Total cyanide remaining in wet gari 5 samples heated at 100 °C for 30 min intervals with water replacement over five cycles at pH 4.3 (△) and pH 6.9 (●).

Fig. 3. % Total cyanide remaining in wet buffered gari 2 samples after heating for 5 h at 30 °C (●) and 50 °C (△) plotted against pH.
cyanide remaining as pH is increased from pH 4.2, where there is no loss. At 30°C only 30% of cyanide is lost at pH 6.5, whereas extensive experiments with cassava flour samples show three- to sixfold losses of cyanide under similar conditions (Bradbury, 2006; Cumbana et al., 2007). This great difference in behaviour between gari and flour is probably due to the hard, gritty nature of roasted gari, and gelatinisation of starch which has fused particles together and restricted the transport of volatile HCN, even when the gari particles are swollen by water. This explanation is consistent with the surprising increased cyanide content of larger gari particles (see Section 3.2).

As shown in Fig. 3 at 50°C there is about twice as much cyanide lost as at 30°C. These initial experiments show that in order to remove acetone cyanohydin from wet gari the temperature of the treatment should be increased above the ambient temperature in the tropics and also the pH of the gari must be increased. There are limited possibilities of methods to increase the pH of gari for use in preparation of thick porridge. One possibility is by mixing gari with low cyanide cassava flour, which has a pH of approximately 6.5 (Bradbury, 2006; Cardoso et al., 2005). This method of increasing the pH by adding cassava flour is explored in Section 3.6.

### 3.6. Cyanide loss from wet gari and wet gari/flour (1:1) mixtures

The five samples of gari were mixed with an equal weight of two different flour samples and excess water, heated at 50°C for 5 h and the loss of total cyanide monitored, see Section 2.2.2. The total cyanide content of the wet gari samples ranged from 1.7 to 8 ppm and of wet MAus7(3) flour was 4.2 ppm and of wet TMS 50395B flour was 10.6 ppm. Mixing gari with an equal weight of flour increased the pH of the mixture to 4.8–4.9, and gari/flour (1:2) gave a pH of 5.1–5.2. The moisture content of the gari/flour (1:1) mixture decreased on heating at 50°C from 60% at zero time to 20, 15 and 8% after 4, 4.5 and 5 h, respectively. The results in Table 2 show as expected, that in the absence of added flour, there is no loss of acetone cyanohydin from wet gari at pH 4.2, but with flour present at pH 4.85, the mean % cyanide remaining is 45–66%. Since the cyanide content of the flour and the gari samples are in the same range (1.7–10.6 ppm, see above) loss of cyanide from both gari and flour has contributed to the result. This is proved by the ca. 66% total cyanide remaining from gari at pH 4.85, see Fig. 3.

### 3.7. Loss of cyanide from wet gari/flour (1:1) in the sun

As shown in Table 3 wet gari did not lose cyanide after 4 h treatment in the sun. There was no breakdown of acetone cyanohydin in wet gari 2 exposed to UV light from a Sunlamp for 2 h. One hour exposure of a wet gari 2 sample to UV light at 254 nm from a Spectrolinker caused heating to 50–60°C with undesirable burning and yellowing of the sample and only a slight reduction in total cyanide content. Clearly, the acetone cyanohydin present in wet gari is not broken down by UV light or by exposure in the sun for 4 h.

### Table 3

<table>
<thead>
<tr>
<th>Experimental conditions</th>
<th>Maximum temperature (°C), using iButton</th>
<th>% Cyanide remaining after 4 h in sun</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wet gari 2, no flour</td>
<td>41</td>
<td>94</td>
</tr>
<tr>
<td>Wet gari 1–5/flour TMS50395B</td>
<td>53–55</td>
<td>58 ± 7</td>
</tr>
<tr>
<td>Wet gari 2/flour MAus7</td>
<td>53</td>
<td>48</td>
</tr>
</tbody>
</table>

* For detailed experimental conditions see Section 2.2.3. pH of all gari/flour (1:1) samples was measured at the end of each experiment and was in the range of 5.0 ± 0.1.
* Mean of 10 experiments.

### 4. Conclusions

It is found that the acetone cyanohydin present in gari can be partially removed by swelling the gari by mixing with water, increasing its pH and raising the temperature of the wet gari. In these experiments the pH was increased by mixing gari with an equal weight of low cyanide cassava flour (but other methods may be discovered which do not destroy the desirable properties of gari as a food) and the temperature raised by exposing the wet mixture in an oven at 50°C for 4 h. The total cyanide content of the mixture was reduced by about one half, which would reduce gari samples that average about 20 ppm total cyanide (Cardoso et al., 2005) to 10 ppm, the WHO safe level (FAO/WHO, 1991). The treated mixture may be cooked in boiling water to produce thick porridge with reduced cyanide content. Adoption...
of this method may reduce the incidence of tropical ataxic neuropathy in Nigeria.

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References


