Improving the Nutritional Functionalities of Unimix Porridge by Lactic Fermentation Using Uji Starter Culture

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Abstract

In 1992, UNImix porridge caused an outbreak of anorexia, vomiting and diarrhoea among children in North Eastern Kenya. This was attributed to Trypsin Inhibitors (Ti) and Haemagglutinins (Hgg) in kidney beans and was solved by replacing Kidney beans with soya beans and extrusion cooking. However, the improved UNImix porridge has been associated with cases of flatulence, diarrhoea and poor palatability among children of displaced refugees in Sudan, due to alpha-oligosaccharides in soybeans, which cannot be eliminated by extrusion-cooking but by lactic fermentation, which is also known to eliminate Ti and Hgg and improve other nutritional functionalities in porridges. This study was therefore aimed at replacing extrusion-cooking in UNImix production with lactic fermentation and determination of its effect on Ti and Hgg. A 3×5 factorial experiment was carried out to test effect of lactic fermentation and cooking on UNImix containing soybeans and kidney beans, and compare results with the extruded-cooked UNImix. All unfermented UNImix samples containing kidney beans tested positive for Ti and Hgg even after boiling for 60 minutes; including those processed by fine milling and drum drying in combination with boiling, but negative on autoclaving at 121°C for 15 min, extrusion-cooking at 100 – 140°C and roasting. Lactic Fermentation alone of UNImix with either kidney beans or soybeans failed to inactivate either Ti or Hgg. However, lactic fermentation combined with boiling for 3-5 minutes eliminated Ti in UNImix samples containing soybeans but not kidney beans, and tested positive for Hgg. It is concluded that Ti and Hgg from Kidney beans are more difficult to inactivate than those in Soya beans and were responsible for the diseases’ outbreak in North Eastern of Kenya in 1992. That lactic fermentation in combination with cooking for 3-5 minutes before consumption can replace extrusion-cooking in UNImix containing soybeans on the basis of inactivation of Ti with benefit of eliminating alpha-oligosaccharides the causative agents for flatulence and diarrhoea, and improvement of palatability among other nutritional functionalities.

Key Words: UNImix porridge, Trypsin inhibitors, Haemagglutinins, LAB Uji culture and Lactic fermentation.

Introduction

UNImix porridge is the pioneer for the nutritional therapeutic porridges. It was developed as an emergency ration for UNICEF Somalia in 1992 to cater for nutrition of displaced children and women in North Eastern Kenya, following overthrow of Somalia government in a military coup in 1991. The initial UNImix porridge consisted of a composite flour of maize and kidney beans fortified with vegetable oil and sugar. It was developed by the Nutrition Department of Ministry of Health of Kenya government. Consumption of this porridge by children in the North Eastern Kenya caused a massive outbreak of anorexia, vomiting and diarrhoea. This led to UNICEF Somalia country office based at Gigiiri, Nairobi Kenya at the time, to sanction investigations on causes of the outbreak and to provide solutions through improvement and development by the Department of Food Science, Nutrition and Technology, University of Nairobi [1].

Results from investigations pointed to inadequate inactivation/elimination of antitrypsin and hemagglutinin factors in the UNImix porridge containing kidney beans, and thus established grounds for the development of the present day UNImix porridge which contains soybeans instead of kidney beans and is extruded-cooked to inactivate trypsin inhibitors and haemagglutinin in the soybeans [1].

Since then several therapeutic porridge rations and complementary foods have been developed along the porridge concept for nutritional intervention and rehabilitation such as WFP mix by the World food program and Corn Soy Blend (CSB) from USA by USAID. The use of porridge rations for nutritional intervention and rehabilitation programs across Sub-Saharan region of Africa in refugee camps, famine distress spots and with people living with HIV virus has been a great success in countries such as Somalia, Kenya, Uganda, Democratic Republic of Congo (DRC), Ethiopia, South Sudan, Darfur region of Sudan, Malawi, Mozambique and Angola [2].

However, Nutritionists in Darfur region of Sudan reported cases of flatulence and diarrhea, including reduced UNImix porridge intake by children due to its low palatability. Flatulence and diarrhoea can occur in some individuals caused by alpha-oligosaccharides namely raffinose, verbascose and stachyose found in legumes like soybeans. Unlike Trypsin inhibitors (Ti) and haemagglutinins (Hgg) also found in legumes, alpha-oligosaccharides cannot be eliminated by extrusion-cooking in
production of UNImix porridge [3]. The alpha-oligosaccharides can however be eliminated by lactic fermentation through hydrolysis using appropriate Lactic Acid Bacteria (LAB) strains such as in Uji starter culture reported to also inactivate trypsin inhibitors and hemagglutinins [4]. The Uji culture exhibited \textit{in vitro} antimicrobial properties against some diarrheal causing pathogens but not \textit{in situ} [8]. Several strains of \textit{Lactobacillus plantarum} strains and especially those studied in some fermented milk products and porridges have been shown to have some probiotic properties.

Lactic fermentation can simultaneously impart several other beneficial nutritional functionalities in UNImix porridge such as improved palatability due to developed appealing taste and flavors, cook ability, protein and starch digestibility, minerals’ bioavailability, lowered dietary bulk, and antimicrobial properties [5-10]. Minerals’ bioavailability is made possible by LAB strains with phytase activity, while improved palatability increases porridge intake critical for enhancing convalescence among clinically malnourished children [3, 10 -12].

This study was therefore aimed at substituting extrusion-cooking with lactic fermentation in UNImix porridge production, with special regard to inactivation/elimination of trypsin inhibitors and hemagglutinins using Uji (LAB) starter culture dominated with strains of \textit{Lactobacillus plantarum}, some strains of \textit{Pediococcus acidilactici} and \textit{Pediococcus pentosaceus} known to eliminate by hydrolysis alpha-oligosaccharides and improve palatability of porridges [4, 5].

**Methods**

**Sourcing of Raw Materials**

Maize, Soybean and Kidney bean flours were obtained from Proctor and Allan Co. Ltd Nairobi, while Soybeans and flakes were obtained from Soy Afric Co. Ltd Nairobi, having been imported from Ohio, U.S.A. The two companies were contracted by UNICEF Somalia to manufacture UNImix porridge ration.

**Fermentation of Unimix**

Slurry samples of UNImix porridge were fermented using Uji starter culture prepared according to Mbugua 1992 [13]. Optimal conditions used for fermentation were ambient temperatures ranging from about 22°C -35°C in keeping with traditional fermentation. Uji culture a mesophilic starter culture was developed, from the traditional fermentation process, and reduced traditional fermentation time from about 72 to 24 hrs.

Fermented slurry samples were prepared by mixing flour blends of UNImix with tap water at 40% w/w flour solids, inoculating with Uji starter culture and incubating at 25°C for 24 hrs in a 3 x 5 factorial design, according to Figure 1. Fermented samples: K1 – K5, S1 – S5 and Sf1 – Sf5, were analysed for presence of Trypsin Inhibitors (Ti) and Haemagglutinins (Hgg) without boiling and after boiling for 3, 5, 10, and 15 minutes.

**Boiling of Samples**

The fermented UNImix samples with 40% flour solids were diluted five times to 8% solids in 125 ml volumes in aluminum cups measuring 6.2 and 7.0 cm of diameter and height respectively and with lids. This helped achievement of appropriate consistency and viscosity for consumption. The slurry samples were brought to boil on a hot plate while stirring, covered and allowed to simmer for 3, 5, 10, and 15 minutes and cooled to 25°C before analysis for Ti and Hgg. The boiling temperature for the slurries was 930°C, the boiling and cooking temperature in Nairobi due to the attitude, but higher or lower temperatures due to attitudes would not matter as long as temperatures do not drop below starch gelatinisation of 720°C only possible in inhabitable high mountains.

**Detection of Trypsin Inhibitors (Ti)**

Trypsin inhibitors were detected by a reaction between crude protein extract from slurry samples and gelatin in the presence of trypsin enzyme according to Reddy et al 1986 [3]. The crude protein was extracted from samples by mixing 5g of slurry with 100ml of 0.1N sodium acetate, shaken for 2 hr in a shaker incubator at 30°C, centrifuged and the supernatant used as testing sample. 2mg trypsin enzyme powder was dissolved in 50 ml of sodium phosphate buffer at pH 7.5. The reaction mixture involved 250 mg gelatin, 5 ml trypsin solution and 5 ml crude protein extract solution mixed to dissolve in a vortex mixer, incubated at 37°C for 2 h. and then cooled to 4°C. Samples containing trypsin inhibitors solidified in 15 minutes, while those without remained liquid indefinitely due to hydrolysis of gelatin by the added trypsin enzyme which was not inhibited. Control sample results for protein extracts from raw flour were used to confirm the results.

**Detection of Haemagglutinins (Hgg)**

Haemagglutinins were again detected according to Reddy et al 1986 in the crude protein extracts [3]. The crude protein extract solution in 0.1N sodium acetate was precipitated by mixing in 1:1 ratio with 6 M sodium hydrogen sulphate solution, followed by separation by centrifugation. The precipitate was re-dissolved in 0.1N sodium acetate. 10 ml of this solution were mixed with 1 ml of blood from rabbit in saline solution. Precipitation and haemolysis of the red blood cells were checked as evidence of presence of haemagglutinins. Control sample results for protein extracts from raw bean flour were used to confirm the results.

**Results**

**Trypsin Inhibitors (Ti) and Haemagglutinins (Hgg) in Cooked Unimix Porridge Containing Kidney Bean and Pigeon Pea Flours**

Table 1 shows the effect of cooking unfermented raw UNImix containing kidney beans, kidney beans protein extracts and pigeon peas on Trypsin Inhibitors (Ti) and Haemagglutinins (Hgg). Presence of both trypsin inhibitors (Ti) and haemagglutinins (Hgg) persisted even after boiling for 60 minutes, but were undetectable after autoclaving at 121°C for 15 minutes.
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K1 – K5 = Kidney bean maize (MKB);
S1 – S5 = Soy bean maize (CSB)
Sf1 – Sf5 = Maize Soy flakes (MSF);
K2, S2, Sf2 = Fermented + 3 min boiling;
K3, S3, Sf3 = Fermented + 5 min boiling;
K4, S4, Sf4 = Fermented + 10 min boiling;
K5, S5, Sf5 = Fermented + 15 min boiling.

Figure 1: Research design

Table 1: Heat stability of Trypsin inhibitors (Ti) and Haemagglutinins (Hgg) in Unimix containing Kidney beans

<table>
<thead>
<tr>
<th>Bean Samples</th>
<th>60 min boiling at 93°C</th>
<th>15 min autoclaving at 121°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ti</td>
<td>Hgg</td>
</tr>
<tr>
<td>All bean flours</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Bean protein extract</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Pigeon peas</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

Ti = Trypsin inhibitor; Hgg = Haemagglutinins; + = Presence; - = Absence

Effect of Selected Processing Treatments of Unimix Containing Kidney Beans on Ti and Hgg

Table 2 shows the effect of roasting, drum drying, fine milling and boiling for 15 minutes and extrusion cooking at various temperatures on Ti and Hgg. Roasting inactivated Ti and Hgg completely, but created undesirable off-flavour and discoloration. Extrusion cooking inactivated Ti and Hgg at temperatures above 1000°C, but left traces of Hgg which were eliminated only at 1400°C. Both drum drying and fine milling together with boiling for 15 min were ineffective in elimination of the two nutritional stress factors.
Effect of Lactic Fermentation of Unimix Porridge on Trypsin Inhibitors and Haemagglutinins

The effect of lactic fermentation of UNImix porridge on Trypsin Inhibitors (Ti) and haemagglutinins (Hgg) is shown in Table 3. The UNImix samples had their pH reduced to 3.0-3.2 and titratable acidities increased to between 0.8 – 1.0 % lactic acid equivalent on fermentation. There were no significant differences in these parameters between the samples. Fermentation alone without boiling did not affect either Ti or Hgg, while Hgg persisted in all fermented samples irrespective of cooking time. Trypsin inhibitors in UNImix containing soybeans and soybeans flakes were eliminated by a combination of fermentation and boiling for 3 and 5 minutes respectively. However, presence of trypsin inhibitors in fermented UNImix containing kidney beans persisted even after boiling for up to 15 minutes.

The study established that Ti and Hgg in kidney beans were either completely heat stable or in such quantities unable to be completely inactivated by boiling for up to 60 minutes, but by autoclaving for 15 minutes at 121°C; hence the conclusion that the Ti and Hgg in the UNImix containing Kidney beans and cooked for 15 minutes were responsible for the outbreak of anorexia, vomiting and diarrhoea among the children fed on the porridge in North Eastern Kenya. The ensuing developmental studies with treatments aimed at inactivating Ti and Hgg namely; drum drying and fine milling combined with boiling for 15 minutes were ineffective in eliminating Ti and Hgg, except for roasting and extrusion cooking. Roasting unfortunately caused discoloration and off-flavours in the UNImix porridge due to maillard reaction known to lower protein digestibility in porridges [14]. Extrusion-cooking on the other hand at temperature of 140°C eliminated both Ti and Hgg without creation of discoloration or off-flavours. Accordingly, all factories contracted to manufacture UNImix porridge by UNICEF Somalia namely; Proctor and Allan Ltd Nairobi, Soy Afric Ltd Nairobi and House of Manji Co Ltd Nairobi were required to use soybeans instead of kidney beans for protein fortification, and pre-cook it by extrusion at 140°C to ensure safety from Ti and Hgg [1].

The presence of Hgg in UNImix porridge persisted irrespective of processing treatments with exception of extrusion-cooking and roasting, whether it contained kidney beans or soybeans. In

particular, boiling of UNImix porridge containing kidney beans for up to 2 hours failed to eradicate its presence an indication of unlikelihood of its elimination in recipes containing kidney beans and cooked under normal households’ procedures [1]. This raises questions on harmfulness of Hgg on ingestion, given that no problems have been reported in communities consuming such diets. Reddy et al 1986 reported no problems associated with consumption of kidney beans diets cooked and processed differently but with detectable Hgg [3]. This fact can indirectly be collaborated by reported improvement of rehabilitative value and efficacy of High Energy porridge (HEP) containing kidney beans by lactic fermentation and presumably with residues of Hgg after cooking on clinically malnourished children in Uganda [11].

Current studies on replacement of extrusion-cooking with lactic fermentation using Uji starter culture showed that fermentation alone and even in combination with boiling of UNImix porridge containing kidney beans for up to 15 minutes was unable to eliminate Ti and Hgg, again demonstrating either stability of Ti and Hgg in kidney beans against inactivation under these conditions or their presence in such high quantities in kidney beans requiring more time for inactivation at the given temperature. However, the Ti in the lactic fermented UNImix porridge containing soybeans but not Hgg was eliminated within 3-5 minutes, implying either more heat labile Ti in Soybeans than those in kidney beans or their presence in such lower levels requiring shorter time for inactivation on basis of their inactivation kinetics. This finding however, justifies replacement of extrusion-cooking with lactic fermentation in production of UNImix containing soybeans on the basis of elimination of Ti on minimal boiling for 3-5 minutes, considering that any residue Hgg are apparently harmless. Coincidentally lactic fermentation is reported to improve cookability of porridges by enhancing starch gelatinization cycles during boiling thus lowering energy requirement unlike in extrusion-cooking, and making it appropriate for production of UNImix porridge at house hold level for nutritional intervention in children by mothers in Africa [7, 12-14].The fact that strains of Leuconostoc mesenteroides involved in lactic fermentation of Idi (an Indian legume-based product) have been reported to inactivate Hgg is a clear indication of diversity of LAB strains which can be harnessed and formulated in to desirable starter cultures for production of therapeutic porridge rations like UNImix porridge with desirable nutritional functionalities, which can be produced at household level in Africa for nutritional intervention and rehabilitation on sustainable basis instead of relying on industrially extruded-cooked porridge rations through donations [3, 10, 15-17].

Conclusion

This study has proved that it is possible to replace the extrusion-cooking process with lactic fermentation using appropriate LAB starter cultures in production of UNImix porridge and related therapeutic porridge rations with profound benefits of improvement of nutritional functionalities of the porridge rations, several of which cannot be provided by extrusion-cooking. Additionally, lactic fermentation would make it feasible to produce therapeutic porridge rations on house hold basis and thus empower mothers in Africa technologically to produce such porridge rations using locally available food substrates for nutritional intervention and rehabilitation of their children on a sustainable basis, without relying on donated industrially produced porridge rations.

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References


