Physical and Chemical Characteristics of Depigmented Oven Dried Dehulled Millet Flours

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ABSTRACT

The physical and chemical characteristics oven dried millet flour from dehulled millet soaked in different media was studied. Depigmented oven dried dehulled millet flour was produced by soaking dehulled millet in different solutions; water, 1% NaCl, 1% Na2CO3, and 1% citric acid. All samples were soaked in their respective solution for 12h, dried in the oven at 60°C for 6h then milled into flour and sieved through 710µm mesh size. The physical properties of oven dried dehulled millet; colour had L* (whiteness value), a* (redness values), and b* (yellowness values) that differed significantly ranging from 66.74 to 84.21, 0.85 to 1.43, and 6.89 to 12.69 respectively. The minimum, mean and maximum particle size distributions of samples ranged from 9.53 to 23.41µm, 29.09 to 50.15µm and 59.46 to 176.01 µm respectively. Starch granules in micrographs of oven dried depigmented millet flours were irregular, compact and polygonally shaped. Gelatinization properties of oven dried millet flour; the onset temperature (To), peak temperature (Tp), end set temperature (Te) and enthalpy varied significantly ranging from 70.15 to 97.65 °C, 79.48 to 102.31 °C, 83.30 to 104.96 °C, and 6.70 to 28.41 (J/g) respectively. Chemical properties comprising of moisture, fats, ash, protein, crude fiber, pH, total titratable acidity, pH of soak solution, phytates and tannins varied significantly ranging from 10.19 to 10.88%, 4.27 to 4.61%, 0.71 to 1.19%, 7.25 to 8.67%, 1.00 to 1.08%, 4.35 to 8.45, 0.001 to 0.084%, 3.32 to 9.93, 0.60 to 0.62, 1.84 to 6.45mg/g and 3.06 to 6.68 mg/g respectively. Depigmenting dehulled millet by soaking in 1% Na2CO3 impaired the colour of sample ODMF2 while depigmenting dehulled millet grains by soaking in 1% citric acid solution for 12 h improved colour of sample ODMF1.

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Introduction

Millet is regarded as an under-utilized cereal, because of its minimal inclusion in commercial food systems, lack of research and novel product development processes even in agro-ecological systems where grown (Shahidi and Chandrasekar, 2013). Hence, millet is often referred to as coarse grain cereal and poor man’s crop due to its use by economically disadvantaged population in Asian and African countries. Millet is a drought, disease and pest resistant crop (Saleh et al., 2013), thus it can be considered as a suitable adaptable crop to mitigate the effect climate change especially in areas where conditions have become desert-like, thereby conferring it a leading role in food security. Millet has a good nutritional profile of either superiority or comparison to rice, maize and sorghum in terms of energy density, protein, fat, minerals and vitamins, yet it has remained a food for low socio-economic groups due to the presence of pigments in the pericarp and endosperm layers, giving the millet products an undesirable color and taste. The seed coat normally imparts dark colour, chewy texture and characteristic musty odour to the food products and these largely hinder their acceptability by the non-traditional millet consumers (Shobana and Malleshi, 2007). The increased use of millet is dependent on the removal of these pigments. Nigeria produces about 5.0 million tonnes of millet annually and had evolved a traditional method of acidifying millet with lime or tamarind juice. Acidification of millet with lime or tamarind juice improved the colour and taste of millet based foods (Nkama and Malleshi, 1998). The coloration of pearl millet is highly correlated with the phenolics inherent in the grain, however, acidification with HCl showed effective reduction of the phenolic content (Reichert, 1979). Anti-nutritional factors of millet, namely phytates, tannins and polyphenols are mainly concentrated in the seed coat and aleurone layers. Different procedures have been used to reduce anti-nutritional factors of grains, namely; soaking, dehulling and application of heat which
in turn effectively improves the nutritional value of grains. The cooking quality and protein digestibility of horse gram (*Dolichos biflorus*) were shown to improve by pre-soaking treatment with salt solution, 1.5% NaHCO$_3$ and 0.75% citric acid for 12h (Kadam and Salunkhe, 1989). In view of the foregoing, there is paucity of information on the use of water and different solutions (1%) of citric acid, Na$_2$CO$_3$ and NaCl as alternatives for the depigmentation of millet. The specific objective of the study was to produce flours from depigmented oven dried millet and evaluate the physical and chemical properties of the flour. It is envisaged that, increased utilization will lead to improving and standardizing the processing of millet not only at household levels but provide an opportunity at industrial level.

**Materials and Methods**

**Materials**

Millet (*Pennisetum glaucum*) was purchased from Alamis market, Lafia in Nasarawa State - Nigeria.

**Sample preparation:** Dehulled millet was processed using the modified traditional method as shown in Fig.1. Dehulled millet was depigmented by soaking in solutions (1%) of citric acid, Na$_2$CO$_3$, NaCl and water. Oven dried millet flours were produced from dehulled depigmented dried grain.

**Preparation of depigmented oven dried millet flour:** Depigmented oven dried millet flour was produced from dehulled millet grains. Dehulled millet grains were cleaned, divided into eight (5) portions of 200g each and soaked for 12h in distilled water and 1% solutions of citric acid, Na$_2$CO$_3$ and NaCl, respectively to effect depigmentation. The 5th portion served as control (no soaking). Production of depigmented oven dried millet flour is as shown in Fig. 2.

**Physical Analyses**

*Colour:* The colour of depigmented oven dried millet flours was measured using HunterLab Colorimeter (UltraScan PRO, USA). Samples were put into an optically transparent glass cell, CIELAB value readings were taken. The ‘L*’ is a measure of whiteness (lightness) ranging from 0 (black) to (100) white, the ‘a*’ value ranges from maximum (redness) to minimum (greenness) and the ‘b*’ value ranges from maximum (yellowness) to minimum (blueness).

*Water activity:* Water activity of oven dried millet flours was measured using a water activity analyser, (Aqualab Model Series 3TE 08038569B Decagon Devices Inc. Pullman, Washington). The water activity was effective in the drying of flour. The moisture content of oven-dried millet was determined using the modified traditional method as shown in Fig.1. The moisture content was determined using a weight loss to equilibrium (AOAC, 2012).

*Particle size:* Particle size of oven dried flour was determined using a laser particle size analyzer (MasterSizer 2000, Malvern Instrument LTD, Malvern UK). Samples were fed into the mastersizer at a vibration feed rate of 50% and a dispersive air pressure of 1.475bar. Particle sizes were classified into average minimum, mean and maximum particle sizes (µm) by an in-built software, Scirocco 2000.

*Scanning Electron Microscopy (SEM):* Depigmented oven dried millet flours were analysed by scanning electron microscopy for microstructure evaluation. Samples were fixed on aluminium stubs using double-sided tape and coated with a layer of gold using the sputter coater to improve conductivity. The coated samples were viewed under a scanning electron microscope (JOEL, made in Japan) using an electron voltage of 15-22v during the scan (Pellisari et al., 2012).

**Differential Scanning Calorimetry:** The gelatinization properties of ODMF were studied using a differential scanning calorimeter (DSC), (Mettler Toledo, Switzerland, UK) equipped with an in-built thermal analysis software, STAR$^\circledR$. Sample (3mg) was loaded into a 40µl capacity aluminium pan with lid (Mettler, ME-26763) and distilled water was added by Hamilton microsyringe, to achieve a flour water suspension containing 70% water. Samples were hermetically sealed and allowed to stand for 1h at room temp before heating in the DSC. The DSC analyser was calibrated using indium and an empty aluminium pan was used as a reference. Sample pans were heated at a rate of 10°C/min from 30 to 120°C. The gelatinization properties of millet flour were defined as To (Onset gelatinization temperature), Tp (Peak gelatinization temperature), Te (end set gelatinization temperature), and AH (enthalpy of gelatinization). These were calculated automatically with the in-built software, STAR$^\circledR$.

**Chemical Analyses**

*Proximate composition:* Ash, crude protein, ether extract, crude fiber and moisture content were determined by the methods of AOAC, (2012).

*Tannin:* Tannin was determined by the method described by Pearson, (1976). One gram of sample was dispersed in 50ml of 70% acetone, placed in an ice bag and allowed to stand for 30min at room temperature, and centrifuged to obtain the tannin extract. Folin- Ciocalteu reagent (1ml) was measured into each flask containing 10ml of extract, followed by 2.5ml of 20% Na$_2$CO$_3$ solution, the mixture was made up to mark in a 50ml flask and incubated for 90min at room temperature. The absorbance was measured at 720nm in a spectrophotometer (Genway model 6000). Tannin solution was prepared and the absorbance read at 720nm. The values were used to prepare the tannin standard curve a series of standard solutions (2-10µg/ml). The concentration of tannin was calculated from standard curve. Readings were taken with the reagent blank at zero. Tannin content was calculated with the expression;

\[
\%\text{Tannins} = \frac{\text{An}}{\text{As}} \times C \times \frac{100}{w} \times \frac{V_F}{V_a}
\]

Where:
- An = Absorbance of test material
- As = Absorbance of standard solution
- C = Concentration of standard solution
- W = Weight of sample
- Vf = Total volume of extract
- Va = Volume of extract analysed

*Phytates:* Phytate was determined by the method of Latta and Eskin, (1980) as modified by Vaintraub and Latta (1980). One gram of sample was extracted with 50ml of 2.4% HCl for 1h at ambient temperature and centrifuged at 3000 rpm for 30min. The clear supernatant was used for the phytate estimation, 1ml of Wade reagent (0.03% FeCl$_3$.6H$_2$O and 0.3% Sulphosalicylic acid) was added and the mixture was made up to mark and incubated for 30min. The absorbance was measured at 267nm using a spectrophotometer (UV-2000 Shimadzu). A series of standard solutions (0.03-0.1µl) was prepared and the absorbance read at 267nm. The values were used to prepare the phytate standard curve. Readings were taken with the reagent blank at zero. Phytate content was calculated with the expression;

\[
\%\text{Phytates} = \frac{\text{An}}{\text{As}} \times C \times \frac{100}{w} \times \frac{V_F}{V_a}
\]

Where:
- An = Absorbance of test material
- As = Absorbance of standard solution
- C = Concentration of standard solution
- W = Weight of sample
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**Results and Discussion**

The gelatinization properties of ODMF were studied using a differential scanning calorimeter (DSC), (Mettler Toledo, Switzerland, UK) equipped with an in-built thermal analysis software, STAR$^\circledR$. Sample (3mg) was loaded into a 40µl capacity aluminium pan with lid (Mettler, ME-26763) and distilled water was added by Hamilton microsyringe, to achieve a flour water suspension containing 70% water. Samples were hermetically sealed and allowed to stand for 1h at room temp before heating in the DSC. The DSC analyser was calibrated using indium and an empty aluminium pan was used as a reference. Sample pans were heated at a rate of 10°C/min from 30 to 120°C. The gelatinization properties of millet flour were defined as To (Onset gelatinization temperature), Tp (Peak gelatinization temperature), Te (end set gelatinization temperature), and AH (enthalpy of gelatinization). These were calculated automatically with the in-built software, STAR$^\circledR$.
added to 3ml of sample solution and centrifuged. The absorbance of the supernatant was measured at 500nm using a spectrophotometer. The standard curve was prepared using absorbance values of working standard solutions of 5-40mg/ml concentrations. The absorbance of Wade reagent not bound by phytic acid was read as the blank. The amount of phytic acid extracted from sample was calculated by subtracting from the absorbance of the blank. The difference was compared with the absorbance readings of the standard solutions, which corresponds with the various concentrations of phytic acid and expressed as phytic acid in mg /100g.

**pH:** pH of a 10% (w/v) dispersion of the samples in distilled water was read with a hand held pH meter (E. Merck model). Each dispersion was mixed thoroughly before the pH was read.

**Total titratable acidity:** Total titratable acidity (TTA) was determined by the method of Nielsen (2002). Ten grammes of sample was added to 100 ml of distilled water, stirred and allowed to stand for 10min. Phenolphthalein indicator (3 drops) was added to 10ml suspension of the mixture and swirled gently. This was titrated against 0.1M NaOH until there was a colour change from colourless to pale pink.

TTA was calculated and expressed as % lactic acid with the expression,

\[
\text{%Lactic acid} = \frac{\text{Volume of NaOH (ml)}}{\text{Volume of Sample (ml)}} \times 0.9 \quad (2)
\]

**Iron, Iodine and Zinc:** Iron, iodine, and zinc were determined by the method documented by Onwuka, (2005). Five grams of each sample was placed in a crucible for ashing in a muffle furnace at 550°C for 6h, after which it was allowed to cool for 1h in the furnace before being transferred to the desiccator. Two grams of ash sample was weighed in a conical flask and 20ml of acid mixture (65% ml Conc. HNO₃, 80ml Perchloric acid and 20ml Conc. H₂SO₄) was added. The conical flask was heated until a clear digest was obtained. The digest was diluted with distilled water to 500ml mark. Ten millilitres of the diluted digest was injected into the atomic absorption spectrophotometer and the absorbance was read at the wavelength (λ_max) of absorption of the respective element. Standard curves were plotted, from which the concentration of each mineral (Iodine, Iron and Zinc) was extrapolated.

**Preparation of standard curves for minerals:** The working standard solutions (0.2 – 2.0µg/ml) for each mineral were prepared from the respective stock solutions. Absorbance of each standard solution was read at the maximum wavelength (Zn at 213.9nm, Fe at 248.3nm and I at 539.5nm) of absorptions of each mineral used to plot the standard curve. The respective mineral concentration was obtained by extrapolation from the standard curve.

**Experimental Design**

The experimental design was Completely Randomized Design (CRD). The data generated from the study was subjected to analysis of variance (ANOVA) using the statistical software IBM SPSS version 20. Means were separated using Duncan Multiple Range Test (DMRT) and significance was accepted at P<0.05.
Significantly from that of depigmented samples
n aggregated mass comprising of several
es of the starch
2°
ern are not
et soaked in 1% citric
pplications.
ODMF2 to 50.15µm
- products undesirable colour and taste,
samples. The morphological featur
drying treatment did not affect the starch granules in the
compact. The photomicrographs show that the integrity of
starch is a homogenous mass of starch granules. Apparently
reported that the endosperm of decorticated finger millet
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(2005) that starch granule population of finger millet
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drying of samples at 60°C were not strong enough
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granules remained intact with no obvious morphological
changes. Okafor et al. (2009) reported that transversely
cut cowpea blanched in 0.5% citric acid solution viewed

Results and Discussion

Physical Properties of Depigmented Oven Dried Millet Flour

Physical properties of depigmented oven dried millet flour is shown in Table 1. The L* (66.74), a* (0.69) and
b* (6.89) values of the depigmented sample (control) differed significantly from that of depigmented samples
which ranged from 70.86 (ODMF2) to 84.21 (ODMF1), 0.85 (ODMF4) to 1.43 (ODMF2) and 8.91 (ODMF3) to
12.69 (ODMF2), respectively.

Higher L* (whiteness) value was observed in oven
dried millet flour from depigmented millet soaked in 1% citric
acid solution followed by oven dried millet flour from
dehulled millet soaked in 1% NaCl and water. A similar
trend was observed in redness value (a*), however for
yellowness value (b*), samples from depigmented millet
soaked in 1% citric acid and 1% Na2CO3 solutions had
higher values. The lower L* value (whiteness) observed in
millet soaked in 1% Na2CO3 solution was attributed to
the polymerization of carbohydrates and proteins which
led to the formation of compounds that were readily
oxidizable at high pH to form dark coloured compounds.
The observed higher whiteness value in sample ODMF1
may be associated with the anti-oxidative effect of citric
acid, which may have caused the whitening effect on the
millet samples. Similar observations were made on

Table 1 Effect of depigmentation treatments on selected physical properties of oven dried millet flour

<table>
<thead>
<tr>
<th>Samples</th>
<th>Colour</th>
<th>Particle size distribution (µm)</th>
<th>Water activity (aw)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L*</td>
<td>a*</td>
<td>b*</td>
</tr>
<tr>
<td>ODMF1</td>
<td>84.21± ±0.21</td>
<td>1.38a± ±0.00</td>
<td>11.37b± ±0.01</td>
</tr>
<tr>
<td>ODMF2</td>
<td>70.86± ±0.17</td>
<td>1.43b± ±0.05</td>
<td>12.69a± ±0.29</td>
</tr>
<tr>
<td>ODMF3</td>
<td>75.72± ±0.24</td>
<td>0.97c± ±0.04</td>
<td>8.91d± ±0.04</td>
</tr>
<tr>
<td>ODMF4</td>
<td>76.45± ±0.02</td>
<td>0.85c± ±0.01</td>
<td>9.63b± ±0.03</td>
</tr>
<tr>
<td>ODMF5</td>
<td>66.74± ±0.08</td>
<td>0.69a± ±0.03</td>
<td>6.89c± ±0.09</td>
</tr>
</tbody>
</table>

Values are means of triplicate determination ± standard error mean (SEM). Means bearing the same superscript in the same column are not significantly different (P>0.05)

Morphology of Starch Granules of Oven Dried Millet Flour from Different Depigmentation Treatments

Scanning electron photomicrographs of starch
granules of differently treated millet flour samples is
shown in Plates 1 to 5. The codes X, L, E, F, J and R in
plates 1, 2, 3, 4 and 5, respectively show ungelatinized
starch granules that are distinctive, interlocking and
compact. The photomicrographs show that the integrity of
the starch granules remained intact suggesting that oven
drying treatment did not affect the starch granules in the
samples. The morphological features of the starch
granules in all the treatment samples were found to be

irregular, compact and polygonally-shaped. The observed
features confirm the earlier report of Adebowale et al.
(2005) that starch granule population of finger millet
cluster to form an aggregated mass comprising of several
small granules. Shobana and Mallesh, (2007) also
reported that the endosperm of decorticated finger millet
is a homogenous mass of starch granules. Apparently
drying of samples at 60°C were not strong enough
treatments to cause gelatinization hence the starch
granules remained intact with no obvious morphological
changes. Okafor et al. (2009) reported that transversely
cut cowpea blanched in 0.5% citric acid solution viewed
through scanning electron microscope showed a closely
knit mat-like arrangement of cells, which is indicative of
structural rearrangement and modification of cowpea cells
by citric acid. This is in agreement with the granule
arrangement of depigmented dehulled millet soaked in
1% citric acid solution (Plate 1) that showed a closely knit
mat-like arrangement with slightly disintegrated cells.

Plate 1 Oven dried millet flour from dehulled millet
soaked in 1% citric acid ×1000

Plate 2 Oven dried millet flour from dehulled millet
soaked in 1% Na$_2$CO$_3$ ×1000

Plate 3 Oven dried millet flour from dehulled millet
soaked in 1% NaCl ×1000

Gelatinization Properties of Oven Dried Millet Flour
from Dehulled Millet Depigmented in Different Solutions

Table 2 shows the gelatinization properties of oven
dried millet flour from dehulled millet depigmented in
different solutions as monitored by Differential Scanning
Calorimetry [DSC]

The onset (To), peak (Tp), end set temperatures of
gelatinization for oven dried millet flour from
undepigmented dehulled millet differed significantly
($P<0.05$) from those of the depigmented samples. The
onset temperature of gelatinization for control sample
(ODMF5) was 97.65°C, while from those of the
depigmented samples which ranged from 70.15°C for
sample ODMF4 to 85.65°C for sample ODMF3. The
higher To temperature observed in sample ODMF5 could
be associated with inadequate uptake of water required for
gelatinization to occur. This invariably suggests that the
sample was not sufficiently hydrated thus the
gelatinization temperature was affected.

The peak temperature of gelatinization (Tp) for
control sample (ODMF5) was 102.31°C, while from those
of the depigmented samples ranged from 79.63°C
(ODMF2) to 96.35°C (ODMF3). Na$_2$CO$_3$ affects the
texture of foods by causing hardening, this effect may
have enhanced the weakening of granules and caused the
sudden collapse of starch structure thus resulting in the
noticeable low peak temperature of gelatinization.

The highest end set temperature (104.96°C) of
gelatinization (Te) was observed in the control sample
(ODMF5) while lowest Te value was observed in sample
ODMF2. Samples ODMF2 (83.30°C) and ODMF4
(84.23°C) had comparable Te values.

Oven dried millet flour from
undepigmented dehulled
millet (control) had an enthalpy ($\Delta H$) value of 6.701J/g
which differed significantly ($P<0.05$) from that of the
depigmented samples which ranged from 7.44 J/g
(ODMF3) to 18.22 J/g (ODMF1). Starch gelatinization is
an order-disorder phase transition of starch granules in the
presence of water and heat which results in loss of the
crystalline order, swelling of granules and solubilization
of starch molecules. This transition is endothermic and
the heat change is called gelatinization enthalpy
(Nwokocha and Williams, 2011).

Higher To, Tp, Te values were observed in the control
sample (ODMF5) than in samples ODMF4, ODMF3 and
ODMF2 probably due to the fact that there was no
hydration so the starch granules did not imbibe enough
water required for gelatinization to occur. High To, Tp,
Te values were observed in the sample soaked in 1%
NaCl (ODMF3). According to Ratnayake et al. (2009)
NaCl exert a protective or restrictive effect on starch
gelatinization which could reflect as high To, Tp, Te
values. Chiotelli et al. (2002) and Day et al. (2013)
reported that NaCl concentrations below 7% increases the
temperature at which starch gelatinization occurs or
begins, whereas high concentrations exert an opposite
effect. Ratnayake et al. (2009) reported that both sodium
chloride and sucrose increase starch gelatinization
temperature and decrease the degree of gelatinization.
According to Bonnet (1993), the amount of amylase in
solution increase with increased concentrations of citric
acid thus hastening granule disruption during
gelatinization while with increased concentrations of
sugar, amyllose in solution decreases thus delaying granule disruption. Acid hydrolysis has been reported to increase the crystallinity of starch granules, this is attributed to the preferential breakdown of amorphous region of the granules (Wang et al., 2006). Starch gelatinization temperature range is a measure of the cooking quality of starch and it is an important parameter in food processing. Starches with low gelatinization temperature ranges are known to have good cooking quality (Waters et al., 2005). High gelatinization temperatures indicate higher stability of starch crystallites in starch molecules.

The pH of the control sample was 6.28 while that of the depigmented samples varied significantly (P<0.05) from that of the depigmented samples, which ranged from 4.35 (ODMF1) to 8.45 in oven dried millet flour from dehulled millet grain soaked in 1% NaCl solution and leaching of protein into soaking media. The percentage crude protein, fat, crude fiber and ash contents of pearl millet reported by various workers (Seghal and Kawatra, 2008) ranged from 7.02 to 13.67, 4.02 to 7.80, 0.54 to 3.00 and 0.25 to 2.54, respectively. The variation in protein content values could be attributed to the effect of the soaking media (1% citric acid, 1% NaCO₃ and water) which may have enhanced leaching of soluble protein. Rath et al. (2003) reported that low values of protein (6.73%) in biscuits prepared from pearl millet depigmented with 0.2N HCl was partly due to hydrolysis and leaching of protein into soaking medium during depigmentation.

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The total ash content of flour from dehulled undepigmented millet (control) was 1.00% while that of the depigmented samples ranged from 0.71 (ODMF1) to 1.19% (ODMF2). The observed high ash content of sample ODMF2 may depict minimal leaching losses when soaked in 1% Na₂CO₃ compared to other soak solutions. The low total ash content recorded in this study compare with the values (0.25 to 2.54%) reported by Seghal and Kawatra, (2008). Karle and Beleia, (2010) reported that the removal of the outer layer of pearl millet accounted for mineral losses.

The undepigmented dehulled sample (control) showed a crude protein content of 8.55% while depigmented samples had crude protein content that ranged from 7.25% in sample ODMF1 to 8.67% in sample ODMF3. Though treatments caused variation in the crude protein content, samples ODMF3 and ODMF5 showed comparable crude protein content which differed significantly (P<0.05) from other samples. The high protein content (8.67%) observed in sample ODMF3 could suggest that NaCl solution was stable, that is had equal number of H⁺ and OH⁻ ion, thus there was no leaching of protein into the soak solution. The percentage crude protein, fat, crude fiber and ash contents of pearl millet reported by various workers (Seghal and Kawatra, 2008) ranged from 7.02 to 13.67, 4.02 to 7.80, 0.54 to 3.00 and 0.25 to 2.54, respectively. The variation in protein content values could be attributed to the effect of the soaking media (1% citric acid, 1% NaCO₃ and water) which may have enhanced leaching of soluble protein. Rath et al. (2003) reported that low values of protein (6.73%) in biscuits prepared from pearl millet depigmented with 0.2N HCl was partly due to hydrolysis and leaching of protein into soaking medium during depigmentation.

The pH of the control sample was 6.28 while that of the depigmented samples varied significantly (P<0.05) from that of the depigmented samples, which ranged from 4.35 (ODMF1) to 8.45 in oven dried millet flour from dehulled millet grain soaked in 1% NaCl solution and leaching of protein into soaking media. The percentage crude protein, fat, crude fiber and ash contents of pearl millet reported by various workers (Seghal and Kawatra, 2008) ranged from 7.02 to 13.67, 4.02 to 7.80, 0.54 to 3.00 and 0.25 to 2.54, respectively. The variation in protein content values could be attributed to the effect of the soaking media (1% citric acid, 1% NaCO₃ and water) which may have enhanced leaching of soluble protein. Rath et al. (2003) reported that low values of protein (6.73%) in biscuits prepared from pearl millet depigmented with 0.2N HCl was partly due to hydrolysis and leaching of protein into soaking medium during depigmentation.
The phytate and tannin contents of control sample (ODMF5) was 6.45mg/100g and 6.68mg/100g, respectively while that of depigmented samples ranged from 1.84mg/100g (ODMF2) to 5.13 mg/100g (ODMF4) and 3.06 (ODMF2) to 5.80 mg/100g, respectively. There were significant differences (P<0.05) between phytate and tannin contents of the control and the depigmented samples. It was observed in this study that the phytate and tannin contents of samples were generally low when compared to reports of previous studies (El-Maki et al., 2007; Mohammed et al., 2011), this could be attributed to the effect of initial dehulling prior to soaking. Also, samples soaked in 1% citric acid solution showed significantly (P<0.05) reduced levels of phytates and tannins than those soaked in water and the undepigmented dehulled sample (ODMF5). Similarly, soaking in 1% Na₂CO₃ significantly (P<0.05) reduced the levels of phytates and tannins, a similar trend was reported by Towo et al. (2003), who observed that soaking in alkali (NaHCO₃, pH 8.4) and acidic media (lactic acid, pH 3) were effective in reducing total amount of extractable phenolics in sorghum and millet. Reichert, (1979) also reported that depigmentation of pearl millet with acid medium was most effective.

It was observed that sample (ODMF1) depigmented in 1% citric acid solution had lower mineral contents (Na, K, P, Mg, Ca, Zn and I) than the control sample (ODMF5) and other samples (ODMF2, ODMF3 and ODMF4) depigmented in other solution. This could be due to the increased concentration of H⁺ ions in the citric acid solution that may have affected the leaching of minerals as the pH of sample was 4.35. Saleh et al. (2013) reported that mineral contents especially P, Ca and Fe were reduced with increased period of soaking pearl millet in acid and attributed it to leaching out of the minerals into the soaking solution.

**Conclusion**

The study has been able to establish that oven dried depigmented millet flours were produced by soaking dehulled millet in different solutions. Depigmenting dehulled millet by soaking in 1% Na₂CO₃ impaired the colour of sample ODMF2 while depigmenting dehulled millet grains by soaking in 1% citric acid solution for 12 h improved colour of sample ODMF1. Higher To, Tp and Te (onset, peak and end gelatinization temperatures) values were observed for undepigmented oven dried millet flour (ODMF5) compared to depigmented samples (ODMF1, ODMF2, ODMF3 and ODMF4). The pH of soaking solutions influenced the chemical composition of oven dried millet flours. Phytate content of sample ODMF2 decreased when dehulled millet was soaked in 1% Na₂CO₃ for 12h. The method most preferred for ODMF involves depigmenting dehulled millet grains by soaking in 1% citric acid solution.

Oven dried millet flours can be used in producing thick porridge such as ‘tuwo’ in accompaniment with soup or thin porridges (gruels) are usually taken for breakfast.

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**Table 2** Gelatinization temperatures of oven dried millet flour from dehulled millet depigmented in different solutions

<table>
<thead>
<tr>
<th>Samples</th>
<th>To (°C)</th>
<th>Tp (°C)</th>
<th>Te (°C)</th>
<th>ΔH (J/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ODMF1</td>
<td>78.78±0.56</td>
<td>90.95±0.11</td>
<td>96.52±0.06</td>
<td>18.22±1.20</td>
</tr>
<tr>
<td>ODMF2</td>
<td>77.13±0.05</td>
<td>79.48±0.00</td>
<td>83.30±0.59</td>
<td>15.17±1.34</td>
</tr>
<tr>
<td>ODMF3</td>
<td>85.65±0.44</td>
<td>96.35±0.00</td>
<td>101.76±0.01</td>
<td>7.44±0.89</td>
</tr>
<tr>
<td>ODMF4</td>
<td>70.15±0.36</td>
<td>79.63±0.50</td>
<td>84.23±0.05</td>
<td>10.81±0.49</td>
</tr>
<tr>
<td>ODMF5</td>
<td>97.65±0.39</td>
<td>102.31±0.09</td>
<td>104.96±0.00</td>
<td>6.70±1.45</td>
</tr>
</tbody>
</table>

Values are means of triplicate determination ± standard error mean (SEM). Means bearing the same superscript in the same column are not significantly different (P>0.05).

**Table 3** Effects of depigmentation treatments on the Chemical composition of oven dried millet flour

<table>
<thead>
<tr>
<th>Parameters</th>
<th>ODMF1</th>
<th>ODMF2</th>
<th>ODMF3</th>
<th>ODMF4</th>
<th>ODMF5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (%)</td>
<td>10.53±0.21</td>
<td>10.70±0.17</td>
<td>10.43±0.24</td>
<td>10.88±0.02</td>
<td>10.19±0.08</td>
</tr>
<tr>
<td>Ether Extract (%)</td>
<td>4.45±0.01</td>
<td>4.27±0.02</td>
<td>4.43±0.11</td>
<td>4.61±0.14</td>
<td>4.49±0.05</td>
</tr>
<tr>
<td>Total Ash (%)</td>
<td>0.71±0.01</td>
<td>1.19±0.03</td>
<td>0.86±0.05</td>
<td>0.77±0.12</td>
<td>1.00±0.08</td>
</tr>
<tr>
<td>Crude Protein (%)</td>
<td>7.25±0.12</td>
<td>7.53±0.25</td>
<td>8.67±0.04</td>
<td>7.77±0.14</td>
<td>8.55±0.05</td>
</tr>
<tr>
<td>Crude fiber (%)</td>
<td>1.08±0.03</td>
<td>1.00±0.01</td>
<td>1.00±0.06</td>
<td>1.07±0.05</td>
<td>1.04±0.08</td>
</tr>
<tr>
<td>pH</td>
<td>4.35±0.01</td>
<td>8.45±0.01</td>
<td>6.08±0.01</td>
<td>6.12±0.01</td>
<td>6.28±0.00</td>
</tr>
<tr>
<td>Total titratable acidity (TTA)</td>
<td>0.084±0.01</td>
<td>0.001±0.00</td>
<td>0.036±0.00</td>
<td>0.027±0.00</td>
<td>0.039±0.00</td>
</tr>
<tr>
<td>pH of soak solution</td>
<td>3.32±0.00</td>
<td>9.93±0.05</td>
<td>5.45±0.00</td>
<td>5.30±0.00</td>
<td>NA</td>
</tr>
<tr>
<td>Phytates (mg/100g)</td>
<td>2.56±0.01</td>
<td>1.84±0.01</td>
<td>3.31±0.02</td>
<td>5.13±0.03</td>
<td>6.45±0.04</td>
</tr>
<tr>
<td>Tannins(mg/100g)</td>
<td>3.77±0.07</td>
<td>3.06±0.05</td>
<td>3.43±0.04</td>
<td>5.48±0.04</td>
<td>6.68±0.01</td>
</tr>
<tr>
<td>Na (mg/100g)</td>
<td>40.50±0.15</td>
<td>72.00±0.10</td>
<td>76.50±0.15</td>
<td>83.50±0.15</td>
<td>65.50±0.15</td>
</tr>
<tr>
<td>K (mg/100g)</td>
<td>127.50±0.15</td>
<td>168.00±0.10</td>
<td>199.50±0.10</td>
<td>193.50±0.15</td>
<td>160.50±0.15</td>
</tr>
<tr>
<td>P (mg/100g)</td>
<td>188.00±0.10</td>
<td>275.00±0.10</td>
<td>284.00±0.10</td>
<td>305.00±0.10</td>
<td>262.50±0.15</td>
</tr>
<tr>
<td>Mg (mg/100g)</td>
<td>286.00±0.01</td>
<td>465.00±0.01</td>
<td>474.00±0.01</td>
<td>533.00±0.01</td>
<td>364.00±0.10</td>
</tr>
<tr>
<td>Ca (mg/100g)</td>
<td>109.50±0.15</td>
<td>268.00±0.10</td>
<td>290.00±0.10</td>
<td>316.50±0.15</td>
<td>205.05±0.10</td>
</tr>
<tr>
<td>Fe (mg/100g)</td>
<td>1.48±0.03</td>
<td>1.46±0.01</td>
<td>1.54±0.00</td>
<td>1.75±0.00</td>
<td>1.32±0.00</td>
</tr>
<tr>
<td>Zn (mg/100g)</td>
<td>2.19±0.15</td>
<td>2.77±0.10</td>
<td>2.94±0.00</td>
<td>3.14±0.00</td>
<td>2.65±0.00</td>
</tr>
<tr>
<td>I (mg/100g)</td>
<td>0.12±0.15</td>
<td>0.27±0.00</td>
<td>0.29±0.00</td>
<td>0.31±0.00</td>
<td>0.20±0.00</td>
</tr>
</tbody>
</table>

Values are means of triplicate determination ± standard error mean (SEM). Means bearing the same superscript on the same row are not significantly different (P>0.05)
References


