Nutritional value and safety of castor bean (Ricinus communis) seeds detoxified in solid-state fermentation by Pleurotus ostreatus

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Abstract. Adebayo OC, Ogidi CO, Akinyele BJ. 2019. Nutritional value and safety of castor bean (Ricinus communis) seed detoxified in solid-state fermentation by Pleurotus ostreatus. Biofarmasi J Nat Prod Biochem 17: 51-60. The nutrient and antinutrient contents of unfermented castor seeds (UCS), castor seeds fermented with Pleurotus ostreatus (CPF) and “Ogiri,” a naturally fermented condiment from castor seeds (CSF) were determined. Proximate analyses of all the samples were carried out using methods of Association of Official Agricultural Chemists. The safety of the samples was carried out using animal model. The raw castor seeds (UCS) had the highest carbohydrates (61.04%), ash (6.02%), fat (6.65%), fiber (6.62%), and calcium (0.30 mg/100g). Castor seeds fermented with Pleurotus ostreatus (CPF) had the highest protein content of 20.47%, magnesium of 7.16 mg/100g, alkaloids (7.40 mg/g) and saponins (6.69 mg/g). CSF had the highest zinc (0.69 mg/100g). CPF had the lowest tamin (0.05 mg/g). All the essential amino acids increased significantly (p<0.05) in the fermented samples. Tryptophan was absent in UCS but present in CSF and CPF with values of 0.78 mg/100g and 1.15 mg/100g, respectively. The hematological analysis of the rats fed CPF had the highest WBC of 5.43x109, which indicated a positive immunomodulatory effect. Hence, this study revealed that Pleurotus ostreatus degraded the toxic compounds in castor seeds to a large extent and enhanced the nutritional contents of the final product.

Keywords: Castor seeds, fermentation, hematological analysis, nutrients, Pleurotus ostreatus

INTRODUCTION

Natural toxins produced by plants are secondary metabolites that protect them from various threats from microorganisms, insects, and predators (Singh et al. 2013). A wide variety of plants such as bitter apricot seeds, bamboo shoots, cassava, flaxseeds, potatoes, wild mushroom, green beans, red and white kidney beans, castor seeds contain natural toxins (RAS 2007). These toxins namely; cyanogenic glycosides, glycoalkaloids, muscarine, lectins may be found throughout the whole plant or in some parts of the plants (seeds, fruits, and leaves). These toxins could be harmful to human health when ingested in a significant amount or when raw materials are not processed properly to final food products.

Castor oil plant (Ricinus communis) belongs to the family of Euphorbiaceae and grows throughout tropical and sub-tropical regions of the world (Ojinnaka et al. 2013; Sousa et al. 2017). Castor seed is a poisonous seed of the castor oil plant (R. communis). The castor seed is inedible because the seed contains a toxic protein, ricin, and other toxic constituents such as ricinidine and ricinoleic acids (Madeira Jr. et al. 2011). The presence of toxic components in castor seeds has remained as a serious impediment in the consumption of castor seeds. Various detoxification methods have been used with varying degrees of success and limitations (Akande et al. 2016). From an economic point of view, these processes are still not practical and not efficient enough to be applied on a large scale.

Fermentation is one of the oldest forms of biotechnology that uses microorganisms to produce secondary metabolites, and recombinant products on an industrial scale (Paulová et al. 2016). The fermentation processes can also be used to enhance food safety by reducing toxic compounds during the production process. Some examples of traditionally fermented condiments produced by solid-substrate fermentation are “Iru” from soya beans (Glycine max), Bambara nut (Vigna subterranea), African locust bean (Parkia biglobosa), “Okpehe” from African Mesquite seeds (Prosopis africana), “Ugba” from oil bean seed (Pentaclethra macrophylla) and “Ogiri” from melon seeds (Cucumropsis manni) or castor seeds (Ricinus communis).

In Nigeria, ‘Ogiri’, a condiment from fermented castor seeds, has a characteristic ammoniacal odor, and its flavor enhances the taste of traditional soups and sauces (Omafuvbe et al. 2004). It serves as a supplement; nutritious non-meat protein substitute and as a functional ingredient in soup. Although, most of the condiments constitute a significant nutritional proportion of diet to many people, there are still a lot of challenges associated with their products such as long hours of boiling, recurrent boiling before the fermentation, having a short shelf life, less acceptable packaging materials, and the typical putrid odour (Ishiwu et al. 2015). Therefore, a new method needs to be introduced by utilizing microorganisms (fungi) to...
make it easier to reduce toxins in the substrate. The use of fungi for eliminating toxins has been adopted in several traditional foods to facilitate the production process.

Some fermented foods have been known in several countries. Aidoo et al. (2006), used Aspergillus oryzae for the fermentation process of rice before the introduction of other yeast to produce Sake (rice wine). ‘Tempeh’, traditional food from Indonesian was produced by fermenting cooked soybeans with a fast-growing fungus: Rhizopus oligosporus (Ilijas et al. 1973). Soy sauce, a condiment made from wheat and soybeans was fermented with Aspergillus spp., yeasts and lactic acid bacteria. ‘Furu’, a condiment side dish consisting of soya bean curd fermented and partly degraded by the mold, Actinomucor elegans (Nout and Aidoo 2011). Fungi such as P. ostreatus possess several lignocellulolytic enzymes (laccase, manganese peroxidase, cellulase, and xylanase), which is selectively used to degrade the toxic compounds in some crops (da Luz et al. 2014). Since, P. ostreatus can be tagged as a Generally recognized as safe (GRAS) fungus because it is a non-pathogenic microorganism and is commonly used to degrade and detoxify food crops before consumption. Therefore, this study was designed to detoxify castor seeds using P. ostreatus in solid-state fermentation and to enhance the nutritional properties of by-product. "

MATERIALS AND METHODS

Collection of castor seeds
Castor seeds were obtained from Akure main market. The raw castor seeds were dehulled, sorted out to remove unwanted materials and dried at 27°C for 7 days. ‘Ogiri’, a locally fermented condiment from castor seeds (CSF) was purchased from the market.

Fungal cultivation
Spawn of Pleurotus ostreatus was obtained from the Federal Institute of Industrial Research, Oshodi (FIIRO), Lagos. The spawn was inoculated into Potato Dextrose Agar (PDA) plate and incubated for seven days at 25°C.

Fermentation of castor seeds with Pleurotus ostreatus
The dehulled castor seeds were divided into two portions. The first portion was boiled for 1 hour, while the second portion was not boiled and left unfermented (UCS). Four agar discs (6 mm) of Pleurotus ostreatus mycelia was inoculated into sterile Potato dextrose broth (20 ml) and left for seven days at 25°C. After that, the broth was aseptically introduced to boiled castor seeds (500 g) and left to ferment for five days using solid-state fermentation. After fermentation, the castor seeds (CPF) were mashed into paste and oven-dried at 28°C for 4 hours.

Proximate analysis of unfermented and fermented castor seeds
Proximate analysis of raw and fermented castor seeds was determined using the method of Association of Official Analytical Chemists (AOAC) (2016). The moisture of the sample was determined by drying 2 g of sample in the oven at 105°C for 3 hours.

% Moisture content = \( \frac{W_2 - W_1}{W_1} \times 100 \)

Where, \( W_1 \) = Weight of crucible; \( W_2 \) = Weight of crucible + sample before drying; \( W_3 \) = Weight of crucible + sample after drying.

The ash content of the sample was determined by incinerating 5.0 g of sample in a muffle furnace at 550°C for 24 h.

% Total ash = \( \frac{W_3 - W_6 \times 100}{W_4} \)

Where, \( W_5 \) = Weight of sample before ashing (g); \( W_6 \) = Weight of dish + sample after ashing; \( W_7 \) = Weight of empty dish.

The fat content was determined by extracting fat from samples using n-hexane in Soxhlet extractor. Briefly, 2 g of sample was wrapped in a filter paper and gradually lowered in the thimble with a fitted flask containing n-hexane. The round-bottomed flask in a Soxhlet extraction unit was slowly heated with thermostatically controlled mantle. The solvent evaporated and passed through the sides tube of the extract to the reflux condenser. The filter paper with the defatted sample was removed from the extractor, and the refluxed solvent distilled out and recovered.

% Crude fat = \( \frac{W_2 - W_1}{S} \times 100 \)

Where, \( W_1 \) = Weight of empty evaporating dish; \( W_2 \) = Weight of evaporating dish + content after drying; \( S \) = Weight of sample after drying.

The protein content was determined using the Kjeldahl nitrogen method. Sample (2.0g) was weighed into the digesting flask. Kjeldahl catalyst and 20 ml of concentrated \( \text{H}_2\text{SO}_4 \) were added to sample and then fixed in the digestion unit (450 °C) of the Kjeldahl apparatus in a fume cupboard for 8 hours. After reaching room temperature, boric acid (20 ml of 4%) was pipetted into a conical flask. A drop of methyl red was added to the flask as an indicator. The sample was diluted with 75 ml of distilled water made alkaline with 20 ml of NaOH (20%) and distilled. The steam exit of the distillatory was closed, and the change in color of boric acid to green was timed. The mixture was distilled for 15 min. The filtrate was then titrated against 0.1N HCl. The protein content (% of Protein) was calculated by the equation:

% Protein content = nitrogen content \times 6.25.

Fiber content was determined using acid and alkaline digestion methods with 20% \( \text{H}_2\text{SO}_4 \) and NaOH solution.

% Fibre content = \( \frac{W_2 - W_3 \times 100}{W_1} \)
Carbohydrate content was determined by the difference method with the equation as follows:

\[
\text{Carbohydrates (\%)} = 100 - (\% \text{ moisture } + \% \text{ protein } + \% \text{ fat } + \% \text{ crude fibre } + \% \text{ ash})
\]

Mineral analysis of unfermented and fermented castor seeds

The method of AOAC (2013) was used to determine the mineral composition of the unfermented and fermented castor seeds. Two g of sample was weighed into a crucible and heated in a muffle furnace at 550°C for 6 hours. The ash was cooled and 6N HCl was added. The mixture was boiled for 10 min, cooled and filtered into 100 ml volumetric flask. The crucible was washed with distilled water, added the filtrate into 100 ml volumetric flask. The volume was made to 100 ml with distilled water. An aliquot of the filtrate was aspirated into the Atomic Absorption Spectrophotometer (Pye Unicam) and the absorbance values corresponding to the different minerals recorded against standard solutions of Ca, Mg, Mn, Fe, Cu, Zn, and Pb.

Phytochemicals analysis of unfermented and fermented castor seeds

The qualitative and quantitative analysis of alkaloids, tannins, saponins, flavonoids were determined by using the methods of Trease and Evans (2005), Sofowora (1993), Harborne and Baxter (1995) and Trease and Evans (2005). Briefly, for alkaloids, sample was stirred with 5 ml of 10% aqueous hydrochloric acid on hot water bath, a few drops of Dragendorf’s reagent was added. The occurrence of orange-red crystalline precipitate indicated the presence of alkaloids. Saponin, 5 g of the sample was grounded and 10 ml of 20% aqueous ethanol was added. The mixture was stirred continuously for 4 hours in hot water bath for 4 hours at about 55°C. The mixture was filtered and the residue was re-extracted using 20% ethanol. The concentrate was transferred into 250 ml separating funnel and 20 ml of diethyl ether was added and shaken together. The aqueous layer was recovered while the ether was later discarded. The purification process was repeated. 60 ml of n-butanol was added and washed twice with 10 ml of 5% aqueous NaCl (Obadoni and Ochuko 2001). Saponin content was calculated as:

\[
\%\text{Saponin} = \frac{\text{Initial weight} - \text{final weight of the sample} \times 100}{\text{Initial weight}}
\]

Flavonoid was quantified by boiling 5 g of each processed sample in 100 ml of 2 M HCl solution for 40 mins. It was allowed to cool to room temperature before being filtered through Whatman filter paper. Flavonoid in the filtrate was precipitated by drop-wise addition of concentrated ethyl acetate until in excess. Following filtration, the flavonoid precipitate recovered was oven-dried and the weight of flavonoid obtained by difference and expressed as a percentage of the sample analyzed (Obadoni and Ochuko 2001).

Tannin was determined by adding 40 ml diethyl ether containing 1% acetic acid (v/v) to 400 mg of each sample. The mixtures were mixed to remove pigment materials. The supernatant was carefully discarded after 5 mins and 20 ml of 70% aqueous acetone added and the flask sealed with cotton plug covered with aluminum foil, then kept in shaker for 2 hours for extraction. The flask content was filtered through Whatman filter paper (No. 2). Aliquot 0.5 ml filtrate was made up to 1.0 ml with distilled water and 0.5 ml Folin Ciocaltiu’s reagent added and then mixed properly before 2.5 ml of 20% sodium carbonate solution was added and further mixed. The mixtures were kept for 40 mins at room temperature, after which absorbance was taken using spectrophotometer and concentration was estimated from the tannic acid standard curve (Sarkiyayi and Agar 2010).

The concentration of phenolics in the sample was determined using the spectrophotometric method at \(\lambda_{max} = 765 \text{ nm} \) (Singleton et al. 1999). The reaction mixture was prepared by mixing 0.5 ml of a methanolic solution of the sample, 2.5 ml of 10% Folin-Ciocaltiu’s reagent dissolved in water and 2.5 ml 7.5% NaHCO_3. Blank was concomitantly prepared, containing 0.5 ml methanol, 2.5 ml 10% Folin-Ciocaltiu’s reagent dissolved in water and 2.5 ml of 7.5% of NaHCO_3. After that, the samples were incubated at 45°C for 45 min. The samples were prepared in triplicate for each analysis, and the mean value of absorbance was obtained. The same procedure was repeated for the standard solution of gallic acid, and the calibration line was constructed. Based on the measured absorbance, the concentration of phenolics was determined (mg/ml) from the calibration curve; then the content of phenolics in extracts was expressed in terms of gallic acid equivalent (mg of GA/g of extract).

Determination of amino acids in unfermented and fermented castor seeds

Initially, 200 mg of each ground seed sample was defatted with a chloroform/methanol mixture in a ratio of 1:1. Then, 30 mg of the defatted sample was put into a glass ampoule, added with 7 ml of 6 M HCl and oxygen expelled by passing nitrogen into the ampoule. The sealed ampoule was put in the oven at 105°C for 22 hours, allowed to cool and filtered. The filtrate was then evaporated to dryness at 40°C under vacuum in a rotary evaporator. The residue was dissolved with 5 ml acetic buffer (pH 2.0) and loaded into the amino acid analyzer where the amino acid compositions of the seed samples were determined by Ion Exchange Chromatographic (IEC) method using the Technicon Sequential Multisample Amino Acid Analyzer (Technicon Instruments Corporation, New York) (Spackman et al. 1958).

Experimental design for in vivo animal model

Twenty albino rats were procured from the Department of Animal Production and Health Science, Federal University of Technology, Akure, Nigeria. The rats were weighed and divided into four groups. They were fed on basal diet and water for seven days before they were fed a different diet. The rats were grouped as follows: (i) BD: Animal fed basal diet only, (ii) BD+UCS: Animal fed basal diet and ground unfermented castor seed, (iii) BD+CSF:...
Animal fed basal diet and commercial fermented castor seeds ("Ogir1"). (iv) BD+CPF: Animal fed basal diet and castor seed fermented with *Pleurotus ostreatus*. The feeding trials were carried out for ten days before the rats were sacrificed.

**Collection of blood sample**

Blood samples were collected by direct cardiac puncture and dispensed into EDTA bottles. The blood samples collected were tested for the red blood cell (RBC) counts, White Blood Cell (WBC) counts, hemoglobin (Hb) concentration, and Packed Cell Volume (PCV) according to the method described by Cheshbrough (2000).

**Histopathological analysis**

Histopathological examination was carried out according to Sarkar et al. (2005). Briefly, liver and kidneys were dissected out and fixed instantaneously in 10% formal saline for 24 hours. The specimens were washed under tap water, dehydrated in ascending grades of ethanol, cleared in xylene, embedded in paraffin wax (melting point of 50-56 °C). Paraffin sections were cut at six µm thicknesses using a rotary microtome; the sections were stained with Harris hematoxylin and eosin. The observation was made using a light microscope and photographs were taken with an automatic photomicrographic system.

**Statistical analysis**

Data obtained from the study were subjected to analysis of variance (ANOVA) using a statistical software SPSS version 22.0 and judged significantly at 95% confidence level (P<0.05).

RESULTS AND DISCUSSION

**Proximate and mineral composition of the unfermented and fermented castor seeds**

Castor seeds fermented with *Pleurotus ostreatus* (CPF) had the highest protein contents of 20.47%, while CSF had the highest moisture contents of 17.29%. UCS had the highest ash (6.02%), fats (6.65%), and fiber (6.62%) (Figure 1). Table 1 shows the mineral composition of raw and fermented castor seeds. CPF had the highest magnesium composition of 7.16 mg/g. CSF had the highest zinc of 0.69 mg/g. UCS had the highest calcium of 0.30 mg/g.

**Phytochemical constituents of the unfermented and fermented castor seeds**

There was no significant difference (P<0.05) in the tannin content of CSF (0.06 mg/g) and CPF (0.05 mg/g). There was a significant increase (P<0.05) in the alkaloid (7.40 mg/g) and saponin contents (6.69 mg/g) of CPF. There was a slightly significant difference in the phenol contents of the unfermented and fermented samples: UCS (1.30 mg/g), CSF and CPF (1.20 mg/g). There were significant differences in flavonoid contents of the samples: UCS (0.65 mg/g), CSF (0.60 mg/g) and CPF (0.48 mg/g).

**Amino acid concentration of the unfermented and fermented castor seeds**

Table 3 revealed the amino acids in the unfermented (UCS) and fermented castor seeds (CSF and CPF). Tryptophan was absent in UCS but had values of 0.78 mg/g and 1.15 mg/g in CSF and CPF respectively. Glutamic acid had the highest concentration in all the samples: UCS had a composition of 10.02 mg/g, CSF had a composition of 10.22 mg/g, and CPF had a composition of 14.59 mg/g. Glutamine and asparagine were absent in all the samples.

**Hematological studies of the albino rats used to test for safety of the unfermented and fermented castor seeds**

Hematological analysis of the rats was tested for safety for unfermented and fermented castor seeds. Rats fed commercial fermented castor seeds (BD+CSF) had the highest Packed Cell Volume (PCV) of 30.67%, while rats fed raw castor seeds (BD+UCS) had the lowest PCV of 16.33%. Rats fed castor seeds fermented with *Pleurotus ostreatus* (BD+CPF) had the highest White Blood Cell (WBC) of 5.43 g/L, while rats fed raw castor seeds (BD+UCS) had the lowest WBC of 3.33 g/L. Rats fed basal diet (BD) had the highest Red Blood Cell (RBC) of 3.42 g/L, while rats fed raw castor seeds (BD+UCS) had the lowest RBC of 1.66 g/L. Rats fed basal diet (BD) had the highest level of hemoglobin of 10.47 g/L, while rats fed raw castor seeds (BD+UCS) had the lowest hemoglobin of 4.47 g/L. Table 5 shows the differential White Blood Cell Count of rats tested for biosafety. BD+ UCS group had the highest lymphocyte count of 59.33%, while BD group had the lowest lymphocyte count, highest neutrophil (46.67%), eosinophils (2.00%), and monocyte (2.00%) count. BD+CPF group had no count for neutrophil, eosinophil, and monocyte. All the groups had no basophil count.

**Histopathological studies of the albino rats used to test for safety of the unfermented and fermented castor seeds**

Plate 1 shows the photomicrograph of the kidneys of rats used to test for biosafety of the raw and fermented castor seed. Kidney of rats fed basal diet (BD) and kidneys of rats fed commercial fermented castor seed (BD+CSF) had the normal histological structure showing the glomerulus, Bowman’s capsule and lymphatic vessels (LV). Kidney of rats fed raw castor seed (BD+UCS) showed enlargement of Bowman’s capsule (EBC), shrunken glomerulus (SG) and distorted proximal and distal convoluted tubules (Z). Kidney of rats fed castor seed fermented with *Pleurotus ostreatus* (BD+CPF) showed disruption of glomerular tuft (D) and hemorrhagic lesions (I). Plate 2 shows the photomicrograph of livers of rats fed raw and fermented castor seeds. Liver of rats fed basal diet and liver of rats fed fermented castor seed showed normal liver structure of sinusoid cells (O) and hepatocytes (N). Liver of rats fed raw castor seed showed dilation of the sinusoid cells (T) and vascular congestion (K). Liver of rats fed castor seeds fermented with *Pleurotus ostreatus* showed dilation of the sinusoid cells (T), vascular congestion (K) and degenerated hepatocytes (DH).
Table 1. Mineral composition (mg/g) of unfermented and fermented castor seeds

<table>
<thead>
<tr>
<th>Minerals</th>
<th>UCS ±SD</th>
<th>CPF ±SD</th>
<th>CSF ±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead</td>
<td>0.03±0.00*</td>
<td>0.02±0.00a</td>
<td>0.02±0.00a</td>
</tr>
<tr>
<td>Zinc</td>
<td>0.13±0.00a</td>
<td>0.38±0.00b</td>
<td>0.69±0.00b</td>
</tr>
<tr>
<td>Iron</td>
<td>0.17±0.00b</td>
<td>0.75±0.00a</td>
<td>0.36±0.00b</td>
</tr>
<tr>
<td>Nickel</td>
<td>0.03±0.00a</td>
<td>0.01±0.00b</td>
<td>0.00±0.00b</td>
</tr>
<tr>
<td>Chromium</td>
<td>0.01±0.00b</td>
<td>0.00±0.00a</td>
<td>0.00±0.00a</td>
</tr>
<tr>
<td>Magnesium</td>
<td>3.79±0.01a</td>
<td>7.16±0.00a</td>
<td>6.86±0.01b</td>
</tr>
<tr>
<td>Manganese</td>
<td>0.04±0.00a</td>
<td>0.21±0.00b</td>
<td>0.21±0.00b</td>
</tr>
<tr>
<td>Copper</td>
<td>0.10±0.00a</td>
<td>0.21±0.00b</td>
<td>0.19±0.00a</td>
</tr>
<tr>
<td>Calcium</td>
<td>0.30±0.00a</td>
<td>0.10±0.00a</td>
<td>0.10±0.00a</td>
</tr>
</tbody>
</table>

Note: Values carrying the same alphabet in the same row are not significantly different (p˃0.05). Values are means of triplicates ±SD. UCS: Unfermented castor seeds, CPF: Castor seeds fermented with Pleurotus ostreatus, CSF: Commercial fermented castor seeds.

Table 2. Quantitative phytochemical compositions (mg/g) of unfermented and fermented castor seeds

<table>
<thead>
<tr>
<th>Parameters</th>
<th>UCS ±SD</th>
<th>CPF ±SD</th>
<th>CSF ±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>*Phenol</td>
<td>1.30±0.01b</td>
<td>1.20±0.00a</td>
<td>1.20±0.01a</td>
</tr>
<tr>
<td>Tannin</td>
<td>0.10±0.01b</td>
<td>0.05±0.00b</td>
<td>0.06±0.01b</td>
</tr>
<tr>
<td>Alkaloids</td>
<td>0.99±0.01a</td>
<td>7.40±0.01c</td>
<td>1.90±0.01a</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>0.65±0.01c</td>
<td>0.48±0.01a</td>
<td>0.60±0.01a</td>
</tr>
<tr>
<td>Saponins</td>
<td>1.95±0.01b</td>
<td>6.69±0.01c</td>
<td>1.40±0.01a</td>
</tr>
</tbody>
</table>

Note: Values carrying the same alphabet in the same row are not significantly different (p˃0.05). Values are means of triplicates ±SD. UCS: Unfermented castor seeds CPF: Castor seeds fermented with Pleurotus ostreatus CSF: Commercial Fermented castor seed, *expressed in terms of Gallic acid equivalent (mg of GA/g of extract) 

Table 3. Amino acid composition (mg/g) of unfermented and fermented castor seeds

<table>
<thead>
<tr>
<th>Amino acids</th>
<th>UCS ±SD</th>
<th>CSF ±SD</th>
<th>CPF ±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leucine</td>
<td>6.55±0.26a</td>
<td>7.58±0.02b</td>
<td>9.92±0.02b</td>
</tr>
<tr>
<td>Lysine</td>
<td>4.49±0.10a</td>
<td>3.71±0.02a</td>
<td>5.62±0.02a</td>
</tr>
<tr>
<td>Phenylnalanine</td>
<td>2.60±0.15b</td>
<td>3.97±0.02b</td>
<td>4.61±0.02b</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>1.64±0.20a</td>
<td>3.81±0.01b</td>
<td>4.30±0.02b</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>0.00±0.00a</td>
<td>0.78±0.03b</td>
<td>1.15±0.01b</td>
</tr>
<tr>
<td>Valine</td>
<td>2.73±0.20b</td>
<td>3.97±0.01b</td>
<td>3.97±0.02b</td>
</tr>
<tr>
<td>Methionine</td>
<td>0.62±0.03b</td>
<td>0.87±0.01b</td>
<td>2.03±0.03a</td>
</tr>
<tr>
<td>Proline</td>
<td>2.96±0.10b</td>
<td>2.35±0.02a</td>
<td>4.06±0.02b</td>
</tr>
<tr>
<td>Arginine</td>
<td>4.67±0.18a</td>
<td>5.31±0.02a</td>
<td>8.52±0.03b</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>1.85±0.05a</td>
<td>2.77±0.02b</td>
<td>4.32±0.02b</td>
</tr>
<tr>
<td>Histidine</td>
<td>1.57±0.11c</td>
<td>2.36±0.02b</td>
<td>2.22±0.02b</td>
</tr>
<tr>
<td>Cystine</td>
<td>0.73±0.08a</td>
<td>0.97±0.02b</td>
<td>1.20±0.01a</td>
</tr>
<tr>
<td>Alanine</td>
<td>2.58±0.35a</td>
<td>3.87±0.02b</td>
<td>4.32±0.02b</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>10.00±0.12a</td>
<td>10.22±0.02a</td>
<td>14.59±0.02a</td>
</tr>
<tr>
<td>Glycine</td>
<td>1.26±0.29a</td>
<td>3.52±0.02b</td>
<td>3.95±0.01b</td>
</tr>
<tr>
<td>Threonine</td>
<td>1.56±0.01a</td>
<td>3.02±0.02b</td>
<td>4.15±0.01b</td>
</tr>
<tr>
<td>Serine</td>
<td>3.32±0.38b</td>
<td>2.32±0.00a</td>
<td>3.62±0.00b</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>7.69±0.15a</td>
<td>7.81±0.03a</td>
<td>12.59±0.01b</td>
</tr>
<tr>
<td>Glutamine</td>
<td>0.00±0.00a</td>
<td>0.00±0.00a</td>
<td>0.00±0.00a</td>
</tr>
<tr>
<td>Asparagin</td>
<td>0.00±0.00a</td>
<td>0.00±0.00a</td>
<td>0.00±0.00a</td>
</tr>
</tbody>
</table>

Note: Values carrying the same superscript in the same row are not significantly different (p˃0.05). Values are means of triplicates ±SD UCS: Unfermented castor seeds CPF: Castor seeds fermented with Pleurotus ostreatus CSF: Commercial Fermented castor seeds.

Table 4. Hematological parameters of rats fed unfermented and fermented castor seeds

<table>
<thead>
<tr>
<th>Groups</th>
<th>PCV (%)</th>
<th>Hb (g/L)</th>
<th>WBC (10^3/g/L)</th>
<th>RBC (10^12/g/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BD+UCS</td>
<td>16.33±1.53a</td>
<td>4.47±0.31a</td>
<td>3.33±0.21a</td>
<td>1.66±0.03a</td>
</tr>
<tr>
<td>BD+CSF</td>
<td>30.67±1.15c</td>
<td>8.43±0.35c</td>
<td>4.30±0.30b</td>
<td>2.23±0.15b</td>
</tr>
<tr>
<td>BD+CPF</td>
<td>22.00±0.00b</td>
<td>7.47±0.15b</td>
<td>5.43±0.31c</td>
<td>2.49±0.10c</td>
</tr>
<tr>
<td>BD</td>
<td>31.67±1.53c</td>
<td>10.47±1.51a</td>
<td>4.17±0.15b</td>
<td>3.42±0.66c</td>
</tr>
</tbody>
</table>

Note: Values carrying the same alphabet in the same column are not significantly different (p˃0.05). Values are means of triplicates ±SD BD: Fed basal diet only, BD+UCS: Fed basal diet mixed with ground unfermented castor seeds, BD+CSF: Fed basal diet mixed with commercial fermented castor seeds, BD+CPF: Fed basal diet mixed with castor seed fermented with Pleurotus ostreatus.

Table 5. Differential white blood cell count of rats fed unfermented and fermented castor seeds

<table>
<thead>
<tr>
<th>Group</th>
<th>Lymphocytes (%)</th>
<th>Neutrophils (%)</th>
<th>Eosinophils (%)</th>
<th>Monocytes (%)</th>
<th>Basophils (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BD+UCS</td>
<td>59.33±1.15c</td>
<td>38.00±2.00b</td>
<td>1.00±0.00b</td>
<td>1.00±0.00b</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>BD+CSF</td>
<td>48.03±1.00b</td>
<td>41.33±1.15c</td>
<td>1.67±0.58c</td>
<td>1.67±0.58c</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>BD+CPF</td>
<td>50.00±2.00b</td>
<td>31.33±1.53a</td>
<td>0.00±0.00b</td>
<td>0.00±0.00b</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>BD</td>
<td>46.00±2.00a</td>
<td>46.67±1.51a</td>
<td>2.00±0.00c</td>
<td>2.00±0.00c</td>
<td>0.00±0.00</td>
</tr>
</tbody>
</table>

Note: Values carrying the same alphabet in the same column are not significantly different (p˃0.05). Values are means of triplicates ±SD. BD: Fed basal diet only, BD+UCS: Fed basal diet mixed with ground unfermented castor seeds, BD+CSF: Fed basal diet mixed with commercial fermented castor seeds, BD+CPF: Fed basal diet mixed with castor seed fermented with Pleurotus ostreatus.
Discussion

The proximate analysis revealed that the fermented castor seeds had higher moisture content than the unfermented castor seeds. The increase in the moisture content could be attributed to the moisture absorbed by the seeds during boiling before fermentation. This result agree with the findings of Ishiwu et al. (2015) that the moisture content of the fermented castor seeds increased after fermentation. However, there was a reduction in the moisture content of castor seeds fermented with *Pleurotus ostreatus* (CPF) when compared to ‘Ogiri’, the naturally fermented condiment from castor seeds (CSF). This result agreed with the findings of Ajala and Akinterinwa (2016) who evaluated the nutrient quality of dried ‘Ogiri’ produced from melon seeds. The researchers observed reduction in the moisture content.

CPF had the highest protein content amongst the three samples (20.47%), while CSF had the lowest protein content (11.14%). This result is in accordance with the findings of Bao et al. (2013) who observed an increase in the protein content (28.17% to 133.42%) of rice fermented with *Pleurotus eryngii*. The increase in the protein content could be due to leaching out of some soluble substances present in the seeds (tannin, trypsin inhibitor) that are protein-bound (Kpanja et al. 2016). The increase in the protein contents could also be due to bioconversion of crude fiber and carbohydrate in the unfermented castor seeds into mycelia protein or single-cell protein (Alemawor et al. 2009). Deepalakshmi and Mirunalini (2014) reported that *Pleurotus ostreatus* are rich in diverse proteins, which possibly enhanced the protein contents of CPF, thereby making its protein content higher than others. Findings of Espinosa-Páez et al. (2017) enhanced the protein digestibility of *Avena sativa* and *Phaseolus vulgaris* by fermentation with *Pleurotus ostreatus*. 
The increase in protein can be attributed to the presence of extracellular enzymes produced by *Pleurotus ostreatus*, which are proteins in nature (Alemawor et al. 2009; Sherief et al. 2010). These enzymes breakdown organic complexes to enhance the release and availability of proteins, which could contribute to the high protein content of the *Pleurotus ostreatus*-fermented castor seeds. The lowest protein content of CSF may be due to leaching during boiling. The lowest protein content may also be caused by the metabolic activities of the proteolytic microorganisms; they used the nitrogen content that contributed to the amount of protein content in the seed, thus, leading to the reduction in the protein contents of the locally fermented castor seeds (Ishiwu et al. 2015).

The fiber contents in the fermented castor seeds were lower compared to the unfermented castor seeds. The lower fiber content could be of medicinal importance. The fiber reduction may be attributed to the ability of the fermenting microfloral to hydrolyze and metabolize castor seed (substrate) as a carbon source to synthesize cell biomass (Madigan et al. 2002). The fat contents in the fermented castor seeds were very low compared to the unfermented castor seeds, which could be attributed to possible degradation of fat by lipase produced by microorganisms during the fermentation (Kpanja et al. 2016). The fat content in castor seeds was deviated from what was reported by Aisha et al. (2013) and Mosquera-Artamonov et al. (2018). This could be as a result of drying castor seed before fermentation and difference in methods adopted.

The ash content of castor seeds fermented with *Pleurotus ostreatus* was the lowest when compared to UCS and CSF. The reduction of ash content in castor seeds fermented with *Pleurotus ostreatus* may be due to leaching as a result of microbial activities that some of the minerals were lost or used (Kpanja et al. 2016). The high ash content of UCS may be due to the high content of mineral present in the seeds, which had been reported by Aisha et al. (2013) when examined wild castor seed from Northern region, Nigeria.

The mineral composition: lead, nickel, chromium, and calcium were reduced in the fermented castor seeds (CSF and CPF) when compared to unfermented (UCS). The finding of Ikanone and Oyekan (2014) revealed that boiling might cause leaching of minerals into water, which may reduce the amount of these minerals in fermented samples. Although, the concentration of zinc, iron, magnesium, manganese, and copper were found to be higher in the fermented castor seeds than in the unfermented castor seeds. This result may be due to the release of these elements from their complex states thereby making it available in the fermented form (Omodara and Olowomofe 2013). Magnesium had the highest concentration in all the samples (UCS, CSF, and CPF), hence fermented castor seed are a source of magnesium. Magnesium is a mineral that is highly essential for human growth; it plays

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**Plate 2.** Photomicrograph of livers of rats fed unfermented and fermented castor seeds. A: fed raw castor seeds (BD+UCS), B: fed commercial fermented castor seeds (BD+CSF), C: fed castor seeds fermented with *Pleurotus ostreatus* (BD+CPF), D: fed basal diet (BD). Dilation of the sinusoid cells (T), degenerated hepatocytes (DN) and vascular congestion (K).
important role in the formation of teeth and bones, in thermoregulation, regulates calcium balance and act as an activator of numerous enzyme systems (Dinu et al. 2014).

The phytochemical study of the unfermented and fermented castor seeds revealed the presence of phenol, saponins, alkaloids, flavonoids, and tannin. The presence of these phytochemicals in foods of plant origin indicates that they can be used as functional foods and for medicinal benefits (Settharaksa et al. 2012). In this study, the concentration of flavonoid was reduced in the fermented castor seeds with CPF having the lowest concentration. Microbial enzymes, such as glucosidase, amylase, cellulase, tannase, esterase, invertase or lipase produced during fermentation are able to hydrolyze glucosides, and break down plant cell walls matrix or starch and consequently facilitating the flavonoids extraction (Hur et al. 2014). During fermentation, the β-glucosidases of microbial origin could also be used to hydrolyze the phenolics and flavonoids (Nazarni et al. 2015) thereby leading to the reduction of flavonoid.

The concentration of tannin was reduced in the fermented castor seeds with CPF having the lowest concentration (0.05 mg/L). Ejikeonye et al. (2018) reported a similar trend when raw watermelon seeds were fermented to a local condiment called ‘Ogiri’. Rodríguez et al. (2009) revealed that during fermentation, tannic acid degrades into smaller components such as glucose and gallic acid. The microorganisms involved during fermentation may also cause a reduction in tannin. According to Fan et al. (2000) and da Luz et al. (2013), *Pleurotus ostreatus* can eliminate antinutrients, mainly tannin by the action of a tannase present in the *Pleurotus ostreatus*, which ultimately destroys the tannins.

Saponin was reduced in ‘Ogiri’, a locally made condiment from castor seeds (CSF), while CPF has the highest concentration. Research carried out by Okwulehie et al. (2018) revealed that *Pleurotus ostreatus* has a higher content of saponin, which could have led to the high saponin content of CPF. The reduction observed in CSF may be as a result of long boiling time and fermentation. Heat processing and fermentation are known to reduce phytochemicals/anti-nutritional concentration in raw food to acceptable nutritional level (Oranusi et al. 2014).

There was no significant difference in the phenolic content of the UCS, CSF, and CPF. This result indicates that castor seeds and its fermented products are a good candidate and a source of phenolic compounds. Phenols are commonly present in foods originated from plant. Phenols protect plants from oxidative damage (Okwu 2005). It has been found highly medicinal for different purposes, specifically block enzymes that cause inflammations, and modify the prostaglandin pathways thereby protecting platelets from clumping (Okwu and Omodamiro 2005). Fermented castor seeds (CPF and CSF) had higher alkaloid content than the raw castor seeds. Research carried out by Okwulehie et al. (2018) revealed that *Pleurotus ostreatus* has high alkaloid content, which may probably be responsible for the increase alkaloid concentration of CPF.

Glutamic acid was the most abundant amino acid followed by aspartic acid, while glutamine and asparagine were absent in the samples (both raw and fermented samples). This trend corresponds with the observations of Khattab et al. (2009) who observed that glutamic and aspartic acid are the most abundant amino acids in some oilseeds and legumes. During the acid hydrolysis, glutamine and asparagine were converted to glutamic and aspartic acids with the liberation of ammonium (NH$_4^+$) ions thereby resulting in the high concentration of glutamic and aspartic acids and also the absence of glutamine and asparagine (Onwuliri and Anekwe 2001). The raw (unfermented) and fermented castor seeds may serve as a good source of monosodium glutamate due to the high concentration of glutamic acid since it is being used as a substrate for a condiment. Monosodium glutamate serves as a major component of food seasonings (Ishiwu et al. 2015). The reduction in protein content may be as a result of boiling time before fermentation, which leads to leaching of the amino acids content. Heat may lead to Maillard reactions and subsequent Amadori rearrangements are making the protein and its amino acids significantly unavailable (Igwe et al. 2012). It was observed that the total content of tryptophan, methionine, and cysteine as essential amino acids was enhanced after fermentation. The increase may be as a result of the action of the microflora and enzymes, which breakdown chemical constituents and thus, enhance the available amino acids (Igwe et al. 2012). Alemawor et al. (2009) reported that *Pleurotus ostreatus* are rich in essential amino acids thereby contributing to an increase of amino acid in CPF. This result agrees with the findings of Bao et al. (2013) who reported the nutritional properties of rice with *Pleurotus eryngii* and found the total content of essential amino acids enhanced after fermentation.

The rats fed UCS and CPF have lower PCV compared to rats fed the basal diet. The decrease in PCV may be as a result of the decreased RBC, which had been earlier reported by Etim et al. (2014). Rats fed CSF had similar PCV to rats fed the basal diet alone. This result suggests that CSF had a positive immunomodulatory effect. The rats fed UCS had the lowest hemoglobin, while the rats fed the basal diet had the highest hemoglobin. The reduced amount of hemoglobin in the groups fed UCS and fermented castor seeds (CPF) may indicate varying degree of unstable alimint. This result agrees with the findings of Ekeh et al. (2010) that observed a reduction in the hemoglobin level of rats drunk with water contaminated with engine oil and suggested that the reduction is due to blood cell deficiency caused by the presence of the toxic substance. The rats fed raw and fermented castor seeds had low RBC compared to rats fed the basal diet. The low values of RBCs may be as a result of their low hemoglobin concentration. RBC is mainly made up of hemoglobin, which gives color to RBCs and helps to transport oxygen throughout the body (Preet and Prakash 2011). The observed reduction in RBC may be attributed to the cytotoxic effects of compounds present in the castor seeds (Eyong et al. 2004). The increase in the lymphocyte count in rat fed castor seeds could have associated with defense mechanism of the body when expose to foreign substance, perhaps the residual compound in castor seeds. Lymphocytes help in the
humoral antibody formation and cellular immunity (Aboderin and Oyetayo 2006). Although, there was relatively increased in White Blood Cells (WBC) and lymphocyte count in rat fed fermented castor seeds, which could also suggest that the fermented castor seeds had an immunostimulatory effect because the rats were stable without any impediments observed. Oluwafemi et al. (2016) reported that Pleurotus ostreatus has high nutritional contents, which boast the immune system of rats since it has been incorporated to the fermented product. The increase in the WBC counts of rat fed CPF suggests that this group would have greater immunity to infections thereby having an immunostimulatory effect. This result agrees with findings of Nfambi et al. (2015) who observed an increase in the WBC count of immunosuppressed rats treated with methanolic leaf extracts of Moringa oleifera. There was a decrease in the neutrophil counts of the rats fed UCS, CSF, and CPF when compared to the control. It may be as a result of the amount of neutrophils used up during sensitization to an antigen (Weber et al. 2015). The significant reduction in the hematological parameters of the rats fed UCS showed the hematotoxicity, a similar effect was reported by Eyyong et al. (2004) who observed a significant reduction in the RBC values of rats and rabbits after ingestion of crude oil, organic molecule containing a toxic compound (aliphatic and aromatic hydrocarbons ), which result in generation of free radical species in various tissues. In red blood cell, free radicals are known to alter erythrocyte membrane as well as other cell membranes as a consequence of oxidative stress (Ita and Udofia 2011). The results of the liver histopathology revealed that the control group and CSF had the normal liver structure of healthy rats, which contains several hepatic lobules. Rats fed UCS and CPF had dilated sinusoid cells and degenerated hepatocytes. The kidney of the rats fed basal diet, and CSF showed normal histological structures of the glomeruli and renal tubules. The kidneys of rat fed the basal diet and CPF had normal kidney structure, while rats fed UCS and CPF showed various defects such as: shrunken glomerulus, hyalinization, enlarged renal tubules, and hemorrhagic lesions. It may be due to the effect of ricin; the toxic compounds present in the seeds. Ricin is quite stable and extremely toxic to the cells of different organs such as the liver, kidney, lung, pancreas, intestine, and thyroid (DaSilva 2003). Ingestion of ricin results in gastrointestinal hemorrhage, necrosis of the liver, spleen, and kidneys; severe localized muscle pain; regional lymph node necrosis, and moderate involvement of visceral organs (Olayiye et al. 2014). This result agreed with the findings of Hassan et al. (2015) who reported the effect of acute phase of toxicity (in different doses) of aqueous extracts of castor seeds on organs of albino rats. They observed that the internal organs showed degenerative changes and proteinaceous material in the spleen and kidney after 24 hours.

In conclusion, this study revealed that Pleurotus ostreatus was able to degrade the toxic compounds present in castor seeds to a greater extent. The results also revealed that Pleurotus ostreatus helped to enhance the nutritional properties of the fermented castor seeds. Pleurotus ostreatus could be used in the detoxification of castor seeds on a larger scale and reduce the boiling time of castor seed before natural fermentation. Pleurotus ostreatus might be used in the detoxification of other poisonous plants.

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