The impact of hydroxypropylated Cassava starch substitution in feed on growth, nutrition, biochemical and toxicological parameters in rat

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Abstract
Experimental diets (0%, 25%, 50% and 100%) using hydroxypropylated cassava starch (HPCS), was evaluated in rat model for one month, growth, antioxidant status, tissue histology and organ toxicity were investigated relative to animals on regular rat chow. Proximate analysis of HPCS compounded-feed showed protein, fibre and ash contents were lower compared to control feed; rats on HPCS had reduced feed-intake, lowered body-weight and organ-weights compared to control. HPCS feeding decreased significantly (P<0.05) urea and protein levels while creatinine and glucose-6-phosphate dehydrogenase (G6DP) increased significantly (P<0.05) in rats on 25% and 50% HPCS feeds compared to both control rats and 0% HPCS. Furthermore, the activity of ALT and ALP increased significantly (P<0.05), while AST significantly (P<0.05) decreased rats on 25%, 50% and 100% HPCS compared to control and 0% HPCS. HPCS feeding in rats elicited increased oxidative stress and lipid peroxidation as liver catalase (CAT), superoxide dismutase (SOD) and malonyl dialdehyde (MDA) were significantly (P<0.05) elevated compared to control and NCS. However, only CAT and SOD levels increased in the kidney of rats on HPCS but not MDA. Only rats on 100% HPCS diet, showed centrilobular fatty degeneration and congested veins in the liver histology while animals on other treatment groups had no lesions in both
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kidney and liver tissues. These results indicate that HPCS may have possible nephrotoxic, hepatotoxic and oxidative effects at different levels of modification and native cassava starch (NCS) or (0%HPCS) should be used with caution for proper growth of rats compared to commercial rat-chow.

Keywords
Feed-intake; Growth; Toxicity; Normal rat chow; NCS; HPCS

Introduction

Starch is one of the most important food ingredients possessing value added attributes for innumerable industrial applications especially when modified as well [1]. Starch is commonly stored in cereals (maize, guinea corn, millet, sorghum, rice) and in roots and tubers (irish/sweet potato, yam, cocoyam and cassava/tapioca). The chemical modification of starch may affect the rate and extent of its digestion in the small intestine [2]. It has been stated that the oxidation or dextrinization of starch, its substitution with hydroxypropyl, acetyl or octenyl succinate groups as well as crosslinking diminish its digestibility [2,3]. Hence, nutritionally starches subjected to chemical modification are acknowledged as resistant starches type 4 (RS 4).

Cassava starch chemical modification is mostly by the reactions of hydroxyl groups on the glucose ring moiety which could be esterification, etherification or oxidation [4]. Cyanogenic glycosides are present in raw unprocessed cassava and it could be contain broken down to produce hydrogen cyanide which can cause both acute and chronic toxicity in humans [5]. Hydrogen cyanide inactivates the enzyme cytochrome oxidase in the mitochondria by binding to the Fe^{3+}/Fe^{2+} and this could cause a decrease in the oxygen utilization of the tissues. Cyanide causes an increase in blood glucose and lactic acid levels but ATP/ADP ratio decreases thus indicating a shift from aerobic to anaerobic metabolism. Processing of cassava such as peeling, slicing, fermentation and cooking helps to reduce both cyanogenic glycosides and hydrogen cyanide significantly and this reduces potential health risk on consumption. Cyanide ion can inhibit several iron, copper or molybdenum metalloenzymes as well as enzymes which contain Schiff base intermediates such as 2-keto-4-hydroxyglutarate aldolase. In female Albino fed raw cassava diet, haemoglobin concentration,
packed cell volume, total serum protein concentration and T4 concentrations decreased while serum thiocyanate levels increased compared to control group [6].

The objective of this study is to evaluate the biochemical and toxicological effects of hydroxypropylated cassava starch (HPCS) and native cassava starch (NCS) which is from a much cheaper carbohydrate source in rat feeding experiment and evaluate its impact on growth and markers of tissue toxicity compared to control rat on commercial chow prepared from maize flour. These would contribute to application and safety issues of both NCS and HPCS in animal feeding and probable the use of either NCS or HPCS as substitute for carbohydrate in feed formulations.

**List of abbreviations**

NCS (native cassava starch), same as 0% hydroxypropylated starch; HPCS (hydroxypropylated cassava starch)

**Material and method**

**Sample collection**

Cassava starch sold for dietary purposes in the Gate market, Ibadan North Central Local Government Area was purchased: samples were air dried and packed into nylon bags until needed for feed formulation. HPCS (7%) was given to us from Industrial Chemistry laboratory, Chemistry Department, Faculty of Science, University of Ibadan, Ibadan already prepared and percentage hydroxypropylation determined.

**Experimental Animals**

Thirty male Wistar rats weighing between 93-110g were purchased from the Animal house of the Faculty of Veterinary Medicine, University of Ibadan, Ibadan, Nigeria. The rats were allowed to acclimatize for 1 week before they were randomly distributed into five groups of six animals. The animals were under standard conditions of temperature and humidity in the Animal house in Biochemistry Department, University of Ibadan. The animals in the control group were fed with normal laboratory rat feed purchased from Ladokun Feeds, Mokola, Ibadan and water was supplied *ad libitum*, while the others had formulated feed using native or hydroxylpropylated (HPCS) starch depending on the group. Protocol for this
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study and animal handling was approved by the Animal Ethics Board, Faculty of Veterinary Medicine, University of Ibadan.

Feed Formulation

The experimental diet was prepared as shown below in Table 1:

<table>
<thead>
<tr>
<th>Components</th>
<th>Concentrations of HPCS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0%</td>
</tr>
<tr>
<td>Soy beans flour</td>
<td>27</td>
</tr>
<tr>
<td>NCS</td>
<td>59</td>
</tr>
<tr>
<td>7% HPCS</td>
<td>-</td>
</tr>
<tr>
<td>Vegetable oil</td>
<td>10</td>
</tr>
<tr>
<td>Vitamin mix</td>
<td>4</td>
</tr>
</tbody>
</table>

The ingredients were thoroughly mixed, pelitized for easy handling by the animals, oven dried at 50°C until properly dried and stored in properly labelled air tight bags to prevent microbial spoilage. Feeds with varying of HPCS were compounded and were fed to rats for four weeks’ duration of the study.

Proximate analysis of diet

The proximate analysis on the NCS and HPCS feed was done using methods described by AOAC (1990) [7].

Sample Collection

The animals were fed for four weeks and sacrificed by cervical dislocation under ether anaesthesia. Blood samples were collected into anticoagulant bottles for plasma and non-anticoagulant tubes. Samples in anticoagulant were centrifuged at 3,500 rpm and plasma was collected into labelled Eppendorf tubes for mineral analysis. Blood samples collected into tubes without anticoagulant were left to stand for 1 hr before centrifugation at 3,500 rpm for 15 minutes to obtain serum. The sera were decanted into tubes and stored at -80°C in the deep freezer. The liver and kidney were washed, dried, weighed and stored in ice-cold 1.15% KCl buffer until needed. These were homogenized in ice-cold 0.1MTris buffer pH 7.4 using Potter-Elvehjem type homogenizer. The homogenates were centrifuged at 10,000 rpm at ≤ 4°C for 15 minutes using Mistral 3000i cold centrifuge and the supernatants containing the liver and
kidney microsomes were collected and stored at -80°C in deep freezer until needed. Small bits of liver and kidney tissue was cut, washed to remove blood stains dried using filter paper, stored in a 40% buffered formaldehyde (formalin) solution before sending it to Veterinary Anatomy Department, University of Ibadan for tissue histology.

**Biochemical assays**

Serum creatinine and urea levels were determined using Randox kit. Glucose-6-phosphate dehydrogenase in liver homogenate was estimated using enzymatic colourimetric method. The activity of catalase in tissue homogenate was determined spectrophotometrically. Briefly, 1ml of sample was mixed with 49ml distilled H₂O to give 1: 50 dilution of the sample. The assay mixture contained 4ml of H₂O₂ solution (800µmoles) and 5ml of phosphate buffer in a 10ml flat bottom flask. 1ml of properly diluted enzyme preparation was rapidly mixed with the reaction mixture by a gentle swirling motion. The reaction was run at room temperature. A 1ml portion of the reaction mixture was withdrawn and blown into 2 ml dichromate/acetic acid reagent at 60 seconds’ intervals. The mononuclear velocity constant, K, for the decomposition of H₂O₂ by catalase was determined using the equation for a first-order reaction: 

\[
K = \frac{1}{t} \log \frac{S_0}{S}
\]

Where S₀ is the initial concentration of H₂O₂ and S is the concentration of the peroxide at t min. The level of SOD activity, measuring thiobarbituric acid reactive substances (TBARS) according whereas earlier described by Nwozo *et al.*, 2012 [8]. Aspartate, alanine aminotransferase and alkaline phosphatase activities were determined using Randox assay kit.

**Statistical analysis**

Results were expressed as the mean ± S.D. (n = 6). A one-way analysis of variance (ANOVA) was used for the data analysis. Significant differences among groups were determined using students t-test at *p* value less than or equal to 0.05.

**Results and discussion**

Starches from different botanical sources such as potatoes, wheat, rice and other foods, vary in colour and physiochemical characteristic. Native, unmodified starches tend to have limited applications in the food industry especially at extreme temperatures and pH ranges. In
general, native starches produce weak-bodied, cohesive, rubbery pastes when heated and undesirable gels when the pastes are cooled. Thus, food manufacturers generally prefer starches with better behavioural characteristic than those provided by native starches which can be achieved by various modifications either by chemical or enzymatic methods. Modified starch is a food additive which is prepared by treating starch or starch granules, causing the starch to be partially degraded [9]. The purposes of this modification are to enhance its properties particularly in specific applications such as to improve the water holding capacity, heat resistant behaviour, reinforce its binding, minimize caking and improve thickening, hence making it possible for it to be useful as fat replacer, thickener etc [10][11].

**Feed composition, intake and growth parameters in rats fed varying percentages of HPCS diet**

Feed composition has previously been shown and proximate analysis of compounded feed using HPCS and NCS are shown on Table 2.

Table 2. Proximate composition of both control diet (normal rat chow) and test diet

<table>
<thead>
<tr>
<th>Components</th>
<th>Commercial Rat Chow (Control feed)</th>
<th>100% HPCS feed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (%)</td>
<td>8.830</td>
<td>8.470</td>
</tr>
<tr>
<td>Fat (%)</td>
<td>4.880</td>
<td>7.650</td>
</tr>
<tr>
<td>Crude fibre (%)</td>
<td>9.220</td>
<td>4.360</td>
</tr>
<tr>
<td>Ash (%)</td>
<td>7.380</td>
<td>6.400</td>
</tr>
<tr>
<td>Protein (%)</td>
<td>18.390</td>
<td>9.630</td>
</tr>
<tr>
<td>Carbohydrate (%)</td>
<td>51.300</td>
<td>63.490</td>
</tr>
</tbody>
</table>

100% HPCS compounded feed had approximately 50% less protein, lower fibre and ash content compared to control diet. Test diet equally had higher carbohydrate and fat contents compared to commercially sold rat chow (Ladokun feed).

Weekly feed intake data are shown on Figure 1 with the control rats having the highest intake especially in the first week and the same group had recorded average intake for four weeks of 94.50±19.41. All test animals had their highest feed intake in the first week except those on 25% HPCS whose highest feed intake was in the second week. Feed intake gradually decreased over time in all feeding groups, average values from control, 0, 25, 50 and 100% HPCS were 94.50±19.41, 68.29±19.32, 66.85±10.89, 79.55±18.77 and 70.89±13.70; with that of 25% HPCS been statistically significantly decreased compared to control.
Changes in total body weights and organs are shown in Figures 2 and 3 respectively. Initial body weight was statistically not different (93-110) g but the final bodyweights after ingesting varying percentages of HPCS for four weeks were significantly decreased compared to control. 100% HPCS was significantly decreased compared to both control and 0% HPCS groups. Control rats had highest weight gain of 81.00±16.35, animals in groups 0, 25 and 50% HPCS had approximately 10g increase in body weight but those on 100% HPCS had weight loss of -18.00±7.53g as shown in Figure 2.
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Rat chow (control feed). This reduced acceptability by the rats is supported by our data on feed intake which had decreased tremendously to about 60% by the fourth week and is further evidenced by decrease in body-weights of the animals to control (Figure 2).

Control rats had biggest liver and kidney weights of 3.24±0.25 and 1.03±0.10 respectively Rats fed varying percentages of HPCS had decreased organ weights compared to the control, only those on 50% HPCS diet had liver weight been significantly lower compared to both control and 0% HPCS groups as shown in Figure 3.

Control rats had biggest liver and kidney weights of 3.24±0.25 and 1.03±0.10 respectively Rats fed varying percentages of HPCS had decreased organ weights compared to the control, only those on 50% HPCS diet had liver weight been significantly lower compared to both control and 0% HPCS groups as shown in Figure 3.

Rats on HPCS and NCS generally lost weight compared to control because of diet given and this lower body weight might have affected the organ weights as well as shown in Figure 3. Organ-body weight ratio may be used as an index of organ swelling, atrophy, or inflammation [12]. The absence of differences in the liver- and kidney-body weight ratio of animals fed HPCS diet is an indication that HPCS feeding did not produce any swelling or inflammation of the organs. By contrast, the increase in the heart-body weight ratio of the rats maintained on the 50% HPCS diet may imply swelling or hypertrophy of the organ and this might imply that the diet elicits toxic effects.

**Effect of feeding different degrees of HPCS on markers of liver toxicity**

Markers of liver tissue toxicity were evaluated using alanine amino transferase ALT, aspartate amino transferase AST and alkaline phosphatases ALP are shown on Figure 4. ALT

![Figure 3. Effect of feeding different percentages of HPCS on organ weights](image-url)
was significantly increased (p<0.05) in the 25, 50 and 100% HPCS fed groups compared to control and NCS fed groups. 0, 50 and 100% HPCS had AST values which were significantly decreased compared to rats fed normal rat chow (control), while 25 and 50% HPCS fed groups had ALP values which were significantly increased relative to normal control rats. Both NCS and HPCS feeding for twenty-eight days caused increase in serum ALT in the liver (Figure 4), elevations in the transaminases levels are considered as the most sensitive markers in the diagnosis of hepatocellular damage and loss of functional integrity of the membrane. Yakubu et al., attributed the increases in serum aminotransferase enzyme activities to their intracellular location in the cytosol, so toxicity affecting the liver with subsequent breakdown in membrane architecture of the cells leads to their spillage into cellular flow where their concentration rises [13]. Leakage of aminotransferases from the liver to the serum thus indicates liver tissue damage. ALP is a “marker” enzyme for the plasma membrane and endoplasmic reticulum, and often it is used to assess the integrity of the plasma membrane [13]. The increase in the serum enzyme levels may be adduced to loss of intact tissue membrane components including ALP from the tissues into the serum [14]. This is an indication that the ordered lipid bilayer of the membrane structure has been damaged or disrupted, leading to the escape of detectable quantity of the enzyme out of the cells of the hepatocytes and nephrons into the extracellular fluid, the serum [14], thus denoting tissue damage.

Figure 4. Effect of feeding different percentages of HPCS on serum ALT, AST and ALP
Effect of feeding different degrees of HPCS markers of kidney tissue toxicity

Results obtained for effect of HPCS feeding on creatinine, urea, serum protein and G6PD are shown on Figures 5 and 6. HPCS feeding caused serum creatinine to be significantly elevated (p<0.05) in both 25 and 100% fed groups compared to rats on control and native starch diets. Urea in rats on HPCS decreased compared to those on normal commercial rat feed. Protein concentration in all treatment groups decreased but only in 50% HPCS treated group was the value significantly decreased compared to control. While G6PD increased in all groups on HPCS but only in 25% HPCS rats were the values increased significantly compared to control.

Figure 5. Effect of feeding different percentages of HPCS on serum creatinine and protein

Figure 6. Effect of feeding different percentages of HPCS on serum urea level and G6PD activity
Creatinine, urea, protein and G6PD levels are known predictors of kidney tissue integrity. Elevations in plasma creatinine levels is indicative of under excretion, hence suggesting kidney impairment [15]. In this study, creatinine significantly increased in rats fed 25% and 100% HPCS starch compared to control, 50% HPCS and NCS, hence test diet did not adversely affect kidney. The decreased levels of serum urea in the rats fed HPCS and NCS diets could be attributed to a decline in liver function as the liver is the primary site for the synthesis of urea. During hepatic injury, blood urea level decreases significantly due to the failure of the liver to convert amino acids and ammonia to urea. G6PD is an enzyme in the pentose phosphate pathway of carbohydrate metabolism and its deficiency results in G6PD haemolytic anaemia. Low levels of G6PD activity, causes a decrease in NADPH and reduced glutathione (GSH) enhance resulting precipitation of deformed hemoglobin, Heinz body formation and red blood cell lysis. In this study NCS and HPCS significantly increased the level of G6PD compared to control. Protein concentrations decreased slightly in rats fed both NCS and HPCS compared to control and altered serum protein is associated with diseased state [16]. Hence, observed results are therefore not indicative of kidney tissue damage in this study.

**Effect of feeding different degrees of HPCS on liver and kidney PMF antioxidant in rats**

On Table 3 is the data obtained on effect of HPCS feeding on lipid peroxidation and enzymatic antioxidants. Liver MDA values increased compared to control and only in groups on 25, 50 and 100% fed rats were the increase statistically significant (p<0.05) but only 100 HPCS was kidney MDA increased compared to control. Liver SOD was increased in the 0, 25 and 50% HPCS treated groups relative to control while kidney SOD was significantly decreased in all treatment groups compared to control except in rats on native cassava starch. Catalase in the liver increased was significantly by HPCS feeding and in kidney only in the 50% HPCS group was the increase significant.

<table>
<thead>
<tr>
<th>Table 3. Effect of feeding of HPCS on lipid peroxidation in liver and kidney of rats</th>
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<tbody>
<tr>
<td>Organs</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>Liver MDA unit/mg protein</td>
</tr>
<tr>
<td>Kidney MDA unit/mg protein</td>
</tr>
<tr>
<td>Liver SOD unit/mg protein</td>
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Lipid peroxidation is an index tissue oxidative damage to membrane lipids [17]. There was a significant increase in MDA levels in the liver of rats fed 25%, 50% and 100% HPCS starch compared NCS and control but not in the kidney, thus implying that there was damage to liver caused by HPCS intake in rats. In order to overcome oxidative stress, cells are equipped with antioxidant enzymes such as superoxide dismutase (SOD) and catalase (CAT). SOD catalysis the reaction superoxide anion radical to hydrogen peroxide and catalase reduces it to water [17]. The increased activities of the antioxidants enzyme could be as a response to mop up accumulated reactive oxygen or increased lipid peroxidation species caused by NCS and HPCS feeding which could have deleterious effect on membrane integrity and function. CAT and SOD activities were increased in rats NCS and HPCS in the liver but not in the kidney, thus the increase implies an activation in the antioxidative enzymes in response increased oxidative stress elicited by dietary intake compared to control.

**Effect of feeding different degrees of HPCS on micronutrient and cyanide levels in rats**

Result obtained on effect of feeding rats with diet compounded with varying percentages of HPCS on plasma levels of zinc, selenium and cyanide are shown on Table 4. HPCS feeding did not affect cyanide levels in rats as there were no significant differences. However, Zn and Se were significantly decreased in rats fed 50% HPCS compared to control.

| Table 4. Effect of feeding HPCS on plasma levels of zinc, selenium and cyanide |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                 | CONTROL         | 0%              | 25%             | 50%             | 100%            |
| Zinc            | 22.624±1.197    | 23.573±3.655    | 23.316±2.824    | 20.046±1.500    | 23.000±0.779    |
| Selenium        | 0.0357±0.0031   | 0.034±0.0072    | 0.0357±0.00569  | 0.0283±0.00153  | 0.0347±0.055    |
| Cyanide         | 0.0123±0.00153  | 0.0107±0.0025   | 0.0147±0.0057   | 0.012±0.0200    | 0.011±0.002     |

Data are presented as mean ± SEM of six rats in each group
* P<0.05 is significant when compared with control
# P<0.05 is significant when compared with rats fed on unmodified cassava starch
Hydroxypropylation did not cause any significant difference in the NCS and HPCS feeding to rats compared to control and this is very important as it indicates that there was no cyanide intoxication, that cassava starch used in this study was safe for animal consumption and had adequate content of essential trace elements such as selenium and zinc.

**Effect of feeding different degrees of HPCS on liver tissue histopathology**

There was no significant lesion in the liver of rats on normal rat chow, native starch as well as 25 and 50% HPCS, however animals on 100% HPCS for four weeks had centrilobular fatty degeneration and congested veins in the tissue histograph (Figure 7a, b, c, d and e). Kidney tissues had normal tissue histology for all treatment groups and as such have not been included for simplicity of data.

![Liver tissue histology](image)

(a) control showing no lesions  (b) native cassava starch showing no lesions  (c) 25% HPCS showing no lesions  
(d) 50% HPCS showing no lesions  (e) 100% HPCS arrow showing centrilobular fatty degeneration and congested veins

**Figure 7. Liver tissue histology**

Histopathology examination showed normal tissue morphology in the kidney, heart and in the liver only the rats on 100% HPCS had centrilobular fatty degeneration and congested veins in the tissue histograph. Histopathology photoptomimrograph for kidney and heart tissue were not included for simplicity of data, however normal kidney morphology obtained was in contrast with serum creatinine result (Figure 7 a-e) which indicated tissue impairment and there is need for caution in interpreting such results. Shahjahan *et al.*, had
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earlier observed that if exposure to a supposed toxicant is low and possibly of a short duration the kidneys could repair the damaged cells and function normally again [18].

Conclusion

Our results indicate that HPCS intake for four weeks, especially when used as total replacement of carbohydrate source in the diet may have possible nephrotoxic and hepatotoxic effects at different levels of modification, it may lead to increase of reactive oxygen species, hence the need for caution.

Also, caution should be exercised in the use of either NCS or HPCS in feed formulation for rats for continual ingestion for four weeks based on both fee intake and bodyweight changes in this study despite the fact that there were no significant differences in cyanide levels.

Observed differences might be due mainly to diminished protein content of cassava starch compared to the grains. Thus, feed formulation from NCS might require fortification for good growth performance.

Acknowledgements

We appreciate Industrial Chemistry Laboratory for the donation of HPCS and Mr Eric Sabo for technical assistance.

References


