Selected Nigerian Local Diet Effects on Blood Glucose Response of Undergraduates of Imo State University, Owerri

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Abstract: The effects of some local diets on the blood glucose response of Ten (10) selected undergraduates of Imo State University were investigated. The food samples investigated were glucose (control), breadfruit cake (Treculia africana), egusi cake (Agbaratti, Kpurukpuru, Mgbam) (Citrulus lanatus) and groundnut soup (Arachis hypogea). The breadfruit was washed, boiled and cooked with other ingredients such as palm oil, pepper, onions, maggi and salt. The egusi sample was sorted, milled and pepper, onions, red oil and salt added. It was molded into shape, wrapped in a leaf (nturukpa) and boiled. The groundnut was sorted and milled and used to prepare groundnut soup. The result obtained from the study revealed that Sample B (breadfruit cake) has the highest moisture content (55.77%). Sample E (egusi cake) recorded the highest protein (33.39%). Sample G (groundnut soup) has the highest fat (12.24%) and ash (3.21%) content. Crude fiber was highest in Sample G (groundnut soup) (2.00%). The highest carbohydrate was seen in Sample B (breadfruit cake) (20.14%) while (Sample G) groundnut soup has the least (5.30%). The overall acceptability of Sample B (breadfruit cake) is higher (8.1%) compared to the overall rating of other food samples. However, no significant difference (p>0.05) existed in overall rating scores of the food samples. The blood glucose response result shows that Glucose has it’s peak at 30 minutes (118.20mg/dl) followed by Sample B (breadfruit cake) (115.0mg/dl) while Egusi cake has the lowest response (82mg/dl). A close relationship was observed between the responses of glucose and Sample B (breadfruit cake) at 15 minutes (94.1 mg/dl and 96.9mg/dl), 30 minutes (118.2mg/dl and 115 mg/dl) and 45 minutes (111.3mg/dl and 118.3mg/dl) which suggest that Sample B (breadfruit cake) contains more carbohydrate than the other test foods. The glycemic index has been recommended to help guide food choice because low-glycemic index foods have been shown to improve blood glucose control in people with diabetes and to increase insulin sensitivity and β-cell function.

Keywords: Local Diets, Undergraduates of Imo State University, Blood Glucose Response

1. Introduction

1.1. Glycemic Index of Foods

Glycemic index (G.I) is the measure of the rate at which an ingested food causes the level of glucose in the body to rise; this is mainly caused by carbohydrate food [21]. It is a system that ranks food at the rate at which their carbohydrates are converted into glucose in the body. It can also be said to be the measure of the effect of food on the blood sugar levels [21]. The glycemic index compares the blood sugar responses to a particular food to the body’s reaction to pure glucose which is given a value of 100 and also used as the main reference point. For instance if a food
raises the blood sugar levels only half as pure glucose, the food receives a glycemic index of (50) [21]. The glycemic index consist of a scale from 1-100, indicating the rate at which 50 grams of carbohydrate in a particular food is absorbed into the blood stream as blood sugar [19].

1.2. Factors That Influence Glycemic Index

Many factors together including carbohydrate type, fiber, protein, fat, food form and method of preparation, influences the glycemic index of a particular food [4]. Some factors that affect the glycemic index of foods includes; processing, fiber content, fat, method of preparation and the amount eaten. The glycemic index value of foods is determined by measuring out 50g portion of digestible carbohydrate of the test food and administering it to patients/subjects for investigations. Their blood glucose levels is checked every 30 minutes for 2 hours and compared against glucose which is used as a reference point [15]. Foods are classified based on the glycemic index value into high, moderate and low glycemic foods [7]. The contribution of glycemic index to nutrition therapy and medical dietetics cannot be over emphasized and yet very little is known about the glycemic index value of most foods we consume. Less effort has been concentrated on the glycemic index value of most commonly consumed Nigerian foods. The knowledge of the glycemic index value of food is not only limited to diabetics alone, as it can be used to treat obesity which is also very prevalent in our society and insulin insensitivity. The above background motivated the research towards determining the effects of some commonly consumed foods and its corresponding glycemic responses on undergraduates. The result of this study will be of immense significance to dietitians and nutritionists as it will encourage them to use the glycemic index as a tool in the diet therapy.

2. Materials and Methods

2.1. Procurement of Materials

All the materials needed for the experiment (Egusi, Breadfruit, and Groundnut) were purchased randomly from Relief market in Owerri, Imo State.

Glucometer /Blood glucose testing meter, Box of test strips, Sterile Lancet, Cotton wool or gauze swabs, Non-sterile gloves, Sharps container, Methylated spirit, Evans glucose (glucose - D), Weighing scale

2.2. Sample Collection

Evans glucose was purchased from the market. Mature fruit samples of Treculia africana were bought from Relief market. Two bottles of fried groundnut was bought. Required quantity of egusi was bought from the named market.

2.3. Sample Processing

2.3.1. Egusi Cake Preparation

2.5kg of dehulled melon seed known as “egusi” was ground alongside 1.5kg of usu, a thickener with 150g (3 bulbs) of sliced onions until a paste was formed, and 300ml of hot water was added. The mixture was placed in a bowl and 2 cooking spoons of red oil, pepper and salt were added and mixed together. The melon paste was then molded into a desirable size using the hands and wrapped with a leaf (ete). Then it was placed into 3000ml of boiling water in a cooking pot on a cooking gas and was allowed to boil for about 50mins under mild heat until it cakes.

2.3.2. Preparation of Breadfruit Cake

The African breadfruit was thoroughly washed to remove dirt and unwanted materials. They were peeled and washed with clean water. It was placed in a clean pot and water was added just the same level as the breadfruit and allowed to boil. After about 15-20 minutes, trona (kaun) was dissolved in 70 mls of water and strained using a sieve of 0.5mm mesh size. Palm oil, onions, pepper and magi were added. Salt was also added to taste and stirred. The food was covered and allowed to boil for a further 45 minutes to 1 hour till done. When done, a spatula was used to mix the food at 5 minutes interval till it cakes.
2.3.3. Preparation of Groundnut Soup

The soup was prepared by washing 700g of red meat, 400g of dried fish which was placed inside a cooking pot and was allowed to heat for about 10mins on a gas cooker. Then 2 cooking spoons of palm oil was introduced followed by the groundnut paste. After which 50g of crayfish, 10g of fresh pepper, two cubes of maggi and about 500ml of water were all added and allowed to boil for about 30mins, Salt was added to taste and the mixture allowed to heat for 3mins till the soup was ready [11].

2.4. Determination of Glycemic Response of Volunteers

The procedure for determining the glycemic response of volunteers is as described by [20]. Following 12hour fast, 10 volunteers ate 25g available carbohydrate portions of the standard reference food (glucose) dissolved in 200mls of water and other food samples at random on different days. The standard food was repeated three times in each subject and their mean IAUC value was calculated as the IAUC of the glucose.

400mls of table water was given to each of the volunteers so that total meal volume is greater than 300ml to stimulate stomach emptying and reduce the variability of glycemic responses. The foods were consumed within 10-15 minutes and the volunteers were asked to remain seated for the duration of the test. Finger prick capillary blood samples was taken from volunteers using lancets before eating the meals (0 minutes) and at 15, 30, 60, 90 and 120 minutes intervals after consumption of the meals, whole blood glucose was measured by dropping the volunteers blood at each of the intervals in a test strip and inserting at the test spot of a glucometer (a one touch basic system glucometer) and reading taken immediately. Incremental Area Under Curve (IAUC) was measured geometrically using the data obtained from the blood glucose concentration, time graph ignoring area beneath the fasting level.[20]

2.5. Survey Design

A random sampling was used for the study.

2.5.1. Population of Study

Ten undergraduate students from the Department of Nutrition and Dietetics, Faculty of Health Sciences aged 20-24 years old were used for the study.

2.5.2. Sample Selection

A random selection was done after which ten healthy non-diabetic undergraduate students were selected for the study. The goal and objectives of the study including skipping breakfast in order to carry out fasting blood sugar was explained.

2.5.3. Sample Size Determination

The sample size was determined using this formula

$$n = \frac{N}{1 + (e^2)}$$

Where

- $n$ = sample size
- $N$ = Population Size
- $I$ = Constant
- $e$ = Margin of error test of significant

i.e., $9.75 = \frac{10}{1.025} + \frac{10}{1.025} + \frac{10}{1.025}$

Approximately 10

2.5.4. Recruiting of Research Assistance

A 400 level medical laboratory student of Imo State University, Owerri was recruited for blood sample collection.
and determination of blood glucose response. The objectives and techniques of the study were explained to her.

2.5.5. Determination of Glycemic Response of Volunteers

The procedure for determining the glycemic response of volunteers [21]. Following 12hour fast, 10 volunteers ate 25g available carbohydrate portions of the standard reference food (glucose) dissolved in 200mls of water and other food samples at random on different days. The standard food was repeated three times in each subject and their mean IAUC value was calculated as the IAUC of the glucose. 400mls of table water was given to each of the volunteers so that total meal volume is greater than 300ml to stimulate stomach emptying and reduce the variability of glycemic responses. The foods were consumed within 10-15 minutes and the volunteers were asked to remain seated for the duration of the test. Finger prick capillary blood samples was taken from volunteers using lancets before eating the meals (0 minutes) and at 15, 30, 60, 90 and 120 minutes intervals after consumption of the meals, whole blood glucose was measured by dropping the volunteers blood at each of the intervals in a test strip and inserting at the test spot of a glucometer (a one touch basic system glucometer) and reading taken immediately. Incremental Area Under Curve (IAUC) was measured geometrically using the data obtained from the blood glucose concentration, time graph ignoring are beneath the fasting level [23].

2.5.6. Determination of Glycemic Index

The incremental area under the curve (IAUC) was calculated for each meal in every volunteer separately (as the sum of the surface of triangles and trapezoids between the B-glucose curve and horizontal baseline going parallel to x-axis from the beginning of B-glucose curve at time 0 to the point at time 120 min) to reflect the total rise in B-glucose concentration after eating the tested food. The IAUCS for the standard reference food (i.e. 25 g of pure glucose) was obtained similarly to the mean from the first three independent IAUCR, IAUCR2, IAUCR3 in the same volunteer. In each volunteer, the glycemic index (%) was calculated by dividing the IAUC for the tested food by the IAUCS for the standard food and multiplying by 100. The following formula was used:

\[ G.I. = \frac{\text{IAUC of the tested meal}}{\text{Mean IAUCR of reference food}} \times 100 \]

Mean IAUCR of reference food
IAUC – Incremental Area Under the blood glucose response Curve for the tested meal
IAUCR – Incremental Area Under the blood glucose response Curve for the reference meal

The glycemic index for each tested food was calculated as the mean from the respective average glycemic indices of the 10 volunteers.

2.5.7. Data Collection

Each of the samples (3 in number) was collected in a plastic container. The ingredients for the 3 food samples, quantities and methods used in soup preparation were recorded. The collected samples were stored in a deep freezer until used for analysis.

2.5.8. Statistical Analysis

Means and standard deviation were subjected to analysis of Variance (ANOVA) to see if there are significant differences among the five food samples in their proximate composition [18]. Turkey LSD test was used to identify significant difference at 5% level of probability.

2.6. Data Analysis

All determinations were done in triplicates. The methods described below were used for the specific analysis.

2.6.1. Sensory Evaluation of the Food Samples

Organoleptic attributes (color, texture, aroma, mouth feel and general acceptability) of the food samples were evaluated using a nine point hedonic scale [8]. 10 subjects selected among the undergraduates were selected as judges for the study. The judges were asked to taste each of the food samples for color, mouth feel, aroma, texture and general acceptability and indicate their feeling about the product on the sensory evaluation sheet.

2.6.2. Proximate Analysis

A standardized method to determine the values of the macronutrients in the food samples using chemical nutritional analytical processes was employed.

2.6.3. Moisture Determination

This was done by the gravimetric method [2]. A measured weight of the sample (5.0g) was weighed into a previously weighed moisture can. The sample in the can was dried in the oven at 105°C for 3hours. It will be cooled in a desiccator and weighed. It was then returned to the oven for further drying. Drying, Cooling and weighing were done repeatedly at hourly (one hour) interval until there are no further diminutions in the weight (ie constant weight was obtained). The weight of moisture lost was calculated and expressed as a percentage of the weight of sample analyzed. It is given by the expression below:

\[ \%\text{Moisture content}=100 \times \frac{W_1 - W_3}{W_2 - W_1} \]

Where W1 = Weight of empty moisture can
W2 = Weight of empty can + sample before drying
W3 = Weight of can + sample dried to constant weight

2.6.4. Determination of Protein

This was done by the Kjeldahl method described by James [10]. The total Nitrogen was determined and multiplied with factor 6.25 to obtain the protein procedures. Half gram (0.5g) of the sample was mixed with 10mls of Conc. H2SO4 in a digestion flask. A tablet of selenium catalyst was added to it before it was heated under a fume cupboard until a clear solution is obtained (i.e. the digest).

The digest was diluted to 100mls in a volumetric flask and used for the analysis. 10mls of the digest was mixed with equal volume of 45% NaOH solution in a Kjeldahl
portion of hot water, the sample was allowed to drain in dry burnt until only ash is left of it. By difference, the weight of before being transferred quantitatively to a weighed crucible previously weighed porcelain crucible. The sample was burnt to processed sample was boiled in 150mls of 1.25% N\(_2\) solution for 30 minutes under reflux. The boiled sample was collected and titrated against 0.02M EDTA from green to a deep red end point. A reagent blank was also digested, distilled and titrated. The N\(_2\) content and hence the protein content was calculated using the formula below.

\[
1\text{Mole of INH}_2\text{SO}_4 = 14\text{mgN}_2
\]

\[
\%\text{Protein} = \%N_2 \times 6.25
\]

\[
\%N_2 = \left( \frac{10}{W} \times \frac{N \times 14}{100} \times \frac{V_1}{V_2} \right) T
\]

\[W = \text{Weight of sample (0.5g)}\]
\[N = \text{Normality of titrant (0.02N H}_2\text{SO}_4)\]
\[V_t = \text{Total digest volume (100mls)}\]
\[V_a = \text{volume of digest analyzed (10ml)}\]
\[T = \text{Sample titre value}\]
\[B = \text{Blank titre value}\]

### 2.6.5. Determination of Ash

This was done by the furnaces incineration gravimetric method [2] 5.0g of the processed sample was measured into a previously weighed porcelain crucible. The sample was burnt to ashes in a muffle furnace at 550°C. When it became completely ashd, it was cooled in a desiccator and weighed. The weight of ash obtained was calculated by difference and expressed as a percentage of the weight of sample analyzed as shown below.

\[
\%\text{Ash} = \frac{100}{1} \times \frac{W_2 - W_1}{W_{\text{t of Sample}}}
\]

Where \(W_1\) = Weight of empty crucible
\(W_2\) = Weight of crucible + ash.

### 2.6.6. Determination of Crude Fiber

The Wende Method [2] was employed. 5.0g of the processed sample was boiled in 150mls of 1.25% N\(_2\)SO\(_4\) solution for 30 minutes under reflux. The boiled sample was washed in several portions of hot water using a two fold Muslim cloth to trap the particles. It was returned to the flask and boiled again in 150mls of 1.25% NaOH for another 30 minutes under same condition. After washing in several portion of hot water, the sample was allowed to drain dry before being transferred quantitatively to a weighed crucible where it was aired in the oven at 10Soc to a constant weight. It was thereafter taken to a muffle furnace in which it was boiled, vaporized and condensed into the reflux flask. Soon the sample in the thimble was covered with the solvent, which contract the oil (fat). The sample remained in contact with the solvent until the reflux flask was filled up and siphoned over, carrying its oil extract down to the boiling flask. This process was allowed to go on repeatedly for 4 hours before the defatted sample was removed, the solvent recovered and the oil extract was in the flask. The flask (containing the oil extract), was dried in the oven at 60°C for 30 mins (to remove any residual solvent). It was cooled in desiccators and weighed. By difference, the weight of oil (fat) extract was determined and expressed as a percentage of the weight of sample analyzed and given by the expression below:

\[
\%\text{Fat} = \frac{w_2 - w_1}{w_{\text{t of Sample}}} \times \frac{100}{1}
\]

\(w_1\) — Initial weight, \(w_2\) — Final weight.

### 2.6.7. Determination of Fat

The solvent extraction gravimetric method [2] was used, 5.0g of the sample was wrapped in a porous paper (Whiteman filter paper) and put in a thimble. The thimble was put in a Soxhlet reflux flask and mounted into a weigh extraction flask containing 200mls of petroleum ether. The upper end of the reflux flask was connected to a water condenser. The solvent (petroleum ether) was heated; it boiled, vaporized and condensed into the reflux flask. Soon the sample in the thimble was covered with the solvent, which contract the oil (fat). The sample remained in contact with the solvent until the reflux flask was filled up and siphoned over, carrying its oil extract down to the boiling flask. This process was allowed to go on repeatedly for 4 hours before the defatted sample was removed, the solvent recovered and the oil extract was in the flask. The flask (containing the oil extract), was dried in the oven at 60°C for 30 mins (to remove any residual solvent). It was cooled in desiccators and weighed. By difference, the weight of oil (fat) extract was determined and expressed as a percentage of the weight of sample analyzed and given by the expression below:

\[
\%\text{Fat} = \frac{w_2 - w_1}{w_{\text{t of Sample}}} \times \frac{100}{1}
\]

### 2.6.8. Determination of Carbohydrates

It will be calculated using the formula below as described by [9].

\[
\%\text{Carbohydrate} = 100\% \times (\text{protein} + \text{fat} + \text{fiber} + \text{ash} + \text{moisture content}).
\]

### 3. Results

Table 1 Proximate composition of the food samples.

The proximate composition of the food samples are presented in table 1. There was significant difference (p<0.05) in the mean values of proximate composition. Sample B has the highest moisture content (55.77%), followed by (Sample G) groundnut soup (46.12%) with (Sample E) egusi cake having the lowest moisture content (41.56%). Sample E egusi cake recorded the highest protein content (33.39%) while (Sample B) breadfruit cake has the least (14.04%). Sample G (groundnut soup) has the highest fat (12.24%) and ash (3.21%) content while Sample B breadfruit cake recorded the least in the two parameters; fat (7.48%) and ash (1.37%). Crude fiber content of (Sample G) groundnut soup (2.00%) is the highest while (Sample B) breadfruit cake recorded the lowest (1.05%) crude fiber content. The highest carbohydrate content was seen in (Sample B) breadfruit cake (20.14%) while (Sample G) groundnut soup contains the least carbohydrate (5.30%).

Table 2 Sensory evaluation of the food samples

(Sample B) breadfruit cake recorded the highest colour content (8.0%) while both (Sample E) egusi cake and (Sample G) groundnut soup recorded the lowest (7.1%).
However, there is no significant difference between the two food samples. The aroma of the food samples significantly increased (p<0.05) with (Sample E) egusi cake recording the highest aroma (8.0%) while (Sample G) groundnut soup recorded the lowest (7.2%). The taste of the food samples also varied significantly (p<0.05) with (Sample G) groundnut soup having the highest taste (7.8%) and (Sample E) egusi cake the lowest (6.7%). (Sample B) breadfruit cake has the highest texture (7.7%) while (Sample E) egusi cake has the lowest (6.8%). The overall acceptability of (Sample E) breadfruit cake is higher (8.1%) than the rest of the food samples while there is no significant difference between that of (Sample E) egusi cake and (Sample G) groundnut soup, both having 7.4%

**Table 1. Mean values of proximate composition of the food samples.**

| Parameters      | Sample E | Sample B | Sample G | LSD (P<0.05) |
|-----------------|----------|----------|----------|--------------|
| Moisture content % | 41.5650±0.02 | 55.7750±0.02 | 46.1250±0.02 | 0.2121 |
| Protein %       | 33.3950±0.02 | 14.0450±0.02 | 31.1200±0.01 | 0.01915 |
| Fat %           | 12.0350±0.02 | 7.4800±0.03 | 12.2400±0.02 | 0.02380 |
| Ash %           | 2.6200±0.01 | 1.3700±0.01 | 3.2150±0.007 | 0.01225 |
| Carbohydrate %  | 7.9150±0.06 | 20.1450±0.10 | 5.300±0.04 | 0.07550 |

Mean values with different superscripts are significantly different (p<0.05) in the same column, LSD = Least significant difference.

**Key**
- Sample B = breadfruit cake
- Sample E = egusi cake
- Sample G = groundnut soup

**Table 2. Mean scores of sensory attributes of the food samples.**

| Parameters | Sample E | Sample B | Sample G | LSD (P<0.05) |
|------------|----------|----------|----------|--------------|
| Colour     | 7.1000±1.52 | 8.0000±0.94 | 7.1000±0.87 | 0.51496 |
| Aroma      | 8.0000±1.33 | 7.4000±0.69 | 7.2000±0.91 | 0.45542 |
| Taste      | 6.7000±1.63 | 8.3000±0.67 | 7.8000±1.03 | 0.52915 |
| Texture    | 6.8000±1.68 | 7.7000±1.25 | 7.0000±1.41 | 0.65376 |
| Overall acceptability | 7.4000±1.42 | 8.1000±0.31 | 7.4000±0.69 | 0.41099 |

Mean values with different superscripts are significantly different (p<0.05) in the same column, LSD = Least significant difference.

**Key**
- Sample B = breadfruit cake
- Sample E = egusi cake
- Sample G = groundnut soup

Table 3 Glucose Response Level (GRL) of the Foods on the Subjects

Table 3 below shows the mean glucose response level of the food samples on the subjects at 0, 15, 30, 45, 60, 90 and 120 minutes. There was no significant difference on the fasting blood glucose of Sample E, and Sample B respondents. There was a significant difference (p<0.05) in the overall blood glucose response of the respondents. (Sample B) breadfruit cake recorded the highest blood glucose response (96.90mg/dl) at 15 minutes while having it’s peak at 45 minutes (118.20mg/dl). (Sample E) egusi cake has the least response at 15 minutes (78.30mg/dl) with it’s peak at 90 minutes (88.8mg/dl). Glucose has it’s peak at 30 minutes (118.20mg/dl) followed by (Sample B) breadfruit cake (115.0mg/dl) while Egusi cake has the lowest response (82mg/dl). The table also revealed a close relationship between the responses of glucose and (Sample B) breadfruit cake at 15 minutes (94.1 mg/dl and 96.9mg/dl), 30 minutes (118.2mg/dl and 115 mg/dl) and 45 minutes (111.3mg/dl and 118.3mg/dl) which suggests that (Sample B) breadfruit cake contains more carbohydrate than the other test foods. However, the result states that the (Sample G) groundnut soup continued increasing even till 120 minutes (92.3mg/dl)

**Table 3. Mean values of Glucose response level (GRL) of indigenous foods fed respondents.**

| Parameters | Glucose | Sample E | Sample B | Sample G | LSD (P<0.05) |
|------------|---------|----------|----------|----------|--------------|
| GRL at 0 min | 78.50±5.68 | 81.20±6.39 | 81.40±4.81 | 87.0±3.62 | 2.33868 |
| GRL at 15 min | 94.10±7.32 | 78.30±5.98 | 96.90±9.92 | 85.0±3.80 | 3.18146 |
| GRL at 30 min | 118.20±9.22 | 82.0±5.75 | 115.0±9.49 | 86.10±3.10 | 3.30194 |
| GRL at 45 min | 111.30±9.09 | 83.30±4.85 | 118.30±11.60 87.50±4.24 | 3.59846 |
| GRL at 60 min | 97.30±1.26 | 86.70±5.73 | 115.30±12.88 88.0±3.29 | 4.29228 |
| GRL at 90 min | 83.90±7.68 | 88.80±5.30 | 105.90±13.86 91.50±4.88 | 3.89437 |
| GRL at 120 min | 80.50±4.83 | 84.00±3.52 | 93.70±10.01 92.30±4.11 | 2.76556 |

Mean values with different superscripts are significantly different (p<0.05) in the same column, LSD = Least significant difference, GRL= Glucose response level, N= Number of respondents fed per day (10 people).

**Key**
- Glucose = Reference food, Sample E = egusi cake
- Sample B = breadfruit cake
- Sample G = groundnut soup
4. Discussion

4.1. Proximate Composition of the Food Samples

Table 1 shows the proximate composition of the food samples. There is a significant difference (p<0.05) in the moisture content of the samples. Breadfruit cake recorded the highest moisture content 55.77% followed by groundnut soup 46.12% and lastly egusi cake which has 41.56%. The high levels of moisture in all the samples investigated suggests that the indigenous diets would not store for long without spoilage since high water activity could enhance microbial action bringing about food spoilage. The result of protein content of indigenous foods indicated that egusi cake has the highest protein content 33.39% followed by groundnut soup 31.12% while breadfruit cake recorded the least 14.04%. The protein value of breadfruit 14.04% was significantly higher (p<0.05) than that reported by [17] 12.47%. The crude protein values obtained in this study did not differ significantly from the values reported by other authors [10].

However, a study reported by [3] reveals that the protein content of breadfruit 17.18% was significantly (p<0.05) higher than that obtained from the present study. This suggests that the food sample can be used to enhance the protein content of foods.

Groundnut soup and Egusi cake both recorded higher values of fat (12.24% and 12.03% respectively) against breadfruit cake with only 7.48%. The breadfruit cake recorded the highest carbohydrate 20.14% while the least was observed in groundnut soup 5.3%. Breadfruit cake may be ranked as carbohydrate rich due to its high carbohydrate content. Thus, it could serve as a good source of energy. The seeds may also serve as a raw material for production of snacks cookies. The crude fiber of breadfruit cake 1.05% obtained tallied with that reported by [3] 1.04%. The fat content of the breadfruit sample obtained 7.4% was significantly higher (p<0.05) than that reported by [3] 6.4%. This may be due to the quantity of oil added to the breadfruit during preparation which may help increase the energy content of the food and thus provide a suitable source of energy.

The study revealed that the proximate composition of groundnut soup obtained were 46.12% moisture content, 31.12% protein, 12.24% fat, 3.21% ash, 2.00% crude fiber and 5.30% carbohydrate. The values obtained for crude protein, crude fiber and ash were not significantly different (p>0.05) from (38.6%, 2.7% and 4.8%) respectively as reported by [1]. The high protein content of groundnut soup suggests that it can be included in complementary foods and thus improve the nutritional content of food.

The proximate composition of egusi cake obtained were 41.56% moisture, 33.39% protein, 12.03% fat, 2.62% ash, 2.15% crude fiber and 7.91% carbohydrate. The ash and protein contents of the sample were in agreement with (2.22% and 30.16%) reported by [15] and values of 2.15 to 2.53% for ash reported by [13] and 38% protein reported by [14] this result shows that the food sample is more of a body building food and it contains some minerals needed to fuel metabolism. [5] reported a higher moisture content 49.40% as opposed to 41.56% obtained from the study. The crude fiber content 3.43% reported by [5] agrees with 3.57% obtained from the study. There is no significant difference (p<0.05) in carbohydrate content 7.22% reported by [8] with 7.915% obtained from the present study.

4.2. Sensory Properties of the Indigenous Food Samples

There was no significant difference (p>0.05) in the colour of the egusi cake and groundnut soup as both samples recorded the same value 71.1%, while breadfruit cake recorded the highest, 8.0%. The taste of the food samples varied significantly (p<0.05) with breadfruit cake recording the highest taste 8.3% while egusi cake recorded the least 6.7%. There was no significant difference in the texture and overall acceptability of the food samples. However all the samples were statistically similar and were accepted equally on the organoleptic characters tested.

4.3. Glycemic Index of the Food Samples

The glycemic index of breadfruit (61), was significantly higher than those of egusi cake (20), and groundnut soup (15). The glycemic index of the breadfruit (61) obtained from this study however agrees with (60), as reported by [20] for breadfruit eaten in the Carribean. When compared with the glycemic index of raw breadfruit which is 68, it could be deduced from the study that there was a reduction in glycemic index of breadfruit which could be as a result of the oil used in preparing the food and also the effect of heat on the food. Breadfruit cake is therefore a moderate glycemic food. Study reported by [18] showed a glycemic index of (24) for boiled groundnut which varied significantly with that obtained from the present study (15) for groundnut.

5. Conclusion

Consuming too many higher or high-glycemic index starchy staples does not appear to be good enough for people in the general population, particularly, for the obesed, type-2 diabetes and pre-diabetic patients. The selected Nigerian local diets has been shown to all be high in moisture content and so low keeping quality. Breadfruit showed the highest
carbohydrate content and it compared favorably with glucose in terms of release of blood glucose. Interestingly, all selected indigenous diets are good sources of food nutrients and there exist a close relationship between the responses of glucose and their blood low glucose response. Legumes generally have a low glycemic index and are favorably used to monitor the blood glucose level of diabetics. They are useful tools in diet formulation.

6. Recommendation

The glycemic index has been recommended to help guide food choice because low glycemic index foods have been shown to improve blood glucose control in people with diabetes to increase insulin sensitivity and β-cell function (the part of the pancreas responsible for insulin secretion) and to reduce serum triacylglycerol. In addition, a low-glycemic index diet has been associated with reduced risk for developing diabetes and cardiovascular diseases and such local foods should be encouraged in consumption.

It is recommended that future work be done on keeping quality of the selected indigenous diets and their compatibility in diet formulation be increased.

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