In vitro digestibility of globulins from sapucaia (Lecythis pisonis Camb.) nuts by mammalian digestive proteinases

Digestibilidade in vitro de globulinas das amêndoas de sapucaia (Lecythis pisonis Camb.) por proteinases digestivas de mamíferos

Sandra Maria Silveira DENADAI1, Priscila Aiko HIANE2, Sergio MARANGONI1, Paulo Aparecido BALDASSO3, Ana Maria Rauen de Oliveira MIGUEL4, Maria Lígia Rodrigues MACEDO2*

Abstract

Sapucaia (Lecythis pisonis Camb.) raw nuts collected from Brazil were analyzed to determine the proximate composition, amino acid profile of protein fractions, in vitro protein digestibility and antinutritional factors in order to evaluate their potential as a protein alimentary complement. The nuts contained adequate amounts of essential amino acids, fatty acids and minerals. In the present study, no hemagglutinating or inhibitory activities were observed in any of the samples investigated, indicating low or non-detectable levels of proteinase inhibitors or lectins in the samples. In vitro digestibility of in natura and heated nut globulins by mammalian digestive proteinases was carried out using trypsin + chymotrypsin + peptidase, with resulting mean values of approximately 70.30 and 71.35%, respectively. Taken together, the results suggest that sapucaia nuts may provide a new source of protein to use as a potential nutritional agent.

Keywords: protein digestibility; antinutritional factors; globulins; proteinases; Lecythis pisonis.

1 Introduction

Plants offer an enormous variety of macro and micronutrients for human consumption. The value of plant proteins in supplying the protein needs in developing countries has been recognized in recent years. Furthermore, fruit, seeds, nuts and almonds from regional native plants have been used to complement the diet of indigenous populations and animal feeding23.33 as they are as rich in proteins, carbohydrates, lipids, vitamins and minerals as legume grains37. Brazilian flora has many native fruit-bearing forest species18,19. Among the edible species consumed in some regions of Brazilian flora, the sapucaia is found in the Atlantic forest and as ‘cumbuca-de-macaco’, among other names. Native to Brazilian rainforests, the sapucaia is found in the Atlantic forest and in the Amazon region34. Its aromatic, sweet-tasting, oleaginous nuts can be consumed raw, boiled or roasted.

The search for alternative, nutritionally suitable and affordable food sources is thus highly desirable. The chemical composition and nutritional value are yet to be studied in many of the native species in Brazilian regions, although regional fruit, which was recently investigated, has been shown to be a good source of nutrients such as amino acids, sugars, fats, vitamins and fibers16,19. Among the edible species consumed in some regions of Brazil is the sapucaia (Lecythis pisonis Camb.), locally known as ‘cumbuca-de-macaco’, among other names. Native to Brazilian rainforests, the sapucaia is found in the Atlantic forest and in the Amazon region34. Its aromatic, sweet-tasting, oleaginous nuts can be consumed raw, boiled or roasted.

Before products of plant origin can be indicated to use as food complements, mainly as proteins sources, investigations should be conducted to determine the amino acid composition of their proteins and protein digestibility, as well as the presence of antinutritional factors. While the amino acid proportionality pattern of a protein is probably the most important determinant of protein quality, digestibility of protein and bioavailability of its constituent amino acids are the next most important factors28. Differences in protein digestibility may arise from inherent differences in the nature of food protein, from the presence of non-protein constituents, which may modify digestion, from the presence of antiphysiological factors or from processing conditions that alter the release of amino acids from proteins by enzymatic processes. However, digestibility still provides a satisfactory rate of protein use4.
In addition, high levels of insoluble fiber and high concentrations of antinutritional factors in diets are also responsible for poor digestibility of proteins\(^1\). Food and feed products may contain a number of antinutritional factors that may adversely affect protein digestibility and amino acid availability\(^2\). Inhibitory proteases, abundant in the plant kingdom, are proteins that can inhibit trypsin, chymotrypsin, amylase, and carboxypeptidase activities\(^1,3,26\). Chronic ingestion of residual levels of antinutritional factors is unlikely to pose risks to human health.

Most animal proteins are well digested, resulting in efficient absorption of amino acids. In contrast, plant proteins are not usually well digested, and are thus nutritionally inferior. The value of plant proteins in supplying the protein needs in developing countries is well acknowledged. If some of the peptide bonds fail to be hydrolyzed in the digestive process, part of the protein content is excreted in the feces or altered into metabolic products by intestinal microorganisms in the large intestine\(^33\). Good plant protein sources are essentially plant foodstuffs whose proteins are well digested.

Most studies designed to provide answers to questions related to the nutritional quality of plant proteins have focused on proteins in the globulin fraction. Globulins are globular proteins that are widely distributed throughout the plant and animal kingdoms. They are soluble in water or in dilute salt solutions. Globulins are involved in transporting a variety of substances, including lipids, hormones, and inorganic ions, in addition to playing a role in the immune system. They are present in seeds in high amounts as storage proteins, and are also found in fractions of antinutritional factors\(^7\). They have structural and enzymatic functions and are important in the germination process\(^33\).

Because the proteins of sapucaia nuts have not been previously characterized, the objective of this work was to study in vitro the action of mammalian proteases—trypsin, chymotrypsin, and pepsin—on globulins and to determine the nutritional value of dry mature sapucaia nuts.

### 2 Materials and methods

#### 2.1 Material

Nuts were obtained from dry mature sapucaia fruit collected from native trees at Estação Experimental de Santa Rita do Passa Quatro, SP at the Instituto de Pesquisa e Estudos Florestais do Estado de São Paulo, Brazil.

#### 2.2 Preparation of defatted meal

The nuts were ground in a Delta Ultrassônico grinder (Delta, São Paulo, SP Brazil) and pulverized to a homogenous powder, which was named whole meal. The whole meal was defatted with petroleum ether PA (40-60 °C) in a Soxhlet extractor (Sebelin TE-188, TECNAL, São Paulo, SP Brazil) for 24 hours. After extraction, the ether was evaporated at 105 °C for 4 hours. The meal was again triturated and pulverized, resulting in a very fine powder that was called defatted meal, which was used as the source of proteins in all the experiments.

#### 2.3 Fatty acid composition

Fatty acid composition of the lipid fraction was obtained after methyl etherification by a procedure described by HARTMAN and LAGO\(^16\). Identification and quantification were carried out using gas-liquid chromatography and flame ionization detection, according to the procedure outlined by FIRESTONE\(^9\) and HORWITZ\(^22\).

#### 2.4 Mineral content

Micro and macromineral contents were determined in the Animal Nutrition Laboratory of EMBRAPA, Campo Grande, MS, Brazil, and the analysis was conducted according to the methodology described by SALINAS and GARCIA\(^32\), involving organic digestion with acids. Manganese, zinc, copper, magnesium, and iron contents were determined using an atomic absorption spectrophotometer. Sodium and potassium were determined by flame photometry and phosphorus and calcium by visible-light spectrophotometry.

#### 2.5 Proximate composition

##### Moisture

The moisture content of the whole meal was determined by stov drying at 105 °C for approximately 4 hours according to methods described in the analytical norms of INSTITUTO ADOLFO LUTZ\(^24\).

##### Total sugar

The total sugar was determined by the reduction method using Fehling’s reagent, according to the procedure described in the norms of INSTITUTO ADOLFO LUTZ\(^24\).

##### Ash

Ash (fixed mineral residue) was determined according to AOAC\(^3\). The total fiber was estimated by differences.

##### Protein

The protein content was measured with the procedure developed by BRADFORD\(^4\) with bovine serum albumin (BSA) as the protein standard and by total nitrogen content (%) according to the Kjeldahl method described in AOAC\(^3\) and multiplied by a factor of 6.25.

#### 2.6 Fractionation of meal proteins

The nut protein fractions used in the present study namely, albumins, globulins, prolamins, glutelins and residue were prepared according to an extraction procedure with NaCl, ethanol, and NaOH\(^27\). Fifteen-gram portions of nut flour were extracted with 150 mL of 4% NaCl for 1 hour. The slurry was centrifuged at 17 000 x g for 30 minutes at 4 °C and the supernatant was then dialyzed against distilled water for albumin and globulin separation. The residue of the salt extraction was suspended in 70% ethanol for 1 hour and again centrifuged as described above to obtain prolamins. The alcohol-insoluble pellet was
suspended in 100 mM NaOH and extracted for 1 hour. Glutelins were then obtained by centrifugation as described above. All the fractions plus the final insoluble residue were recovered by dialysis and freeze-drying.

2.7 SDS-PAGE-polyacrylamide gel electrophoresis

This method was carried out using a LAEMMLI system (1970). The proteins used as molecular mass standards were: phosphorylase b (94 kDa), bovine serum albumin (66 kDa), ovalbumin (45 kDa), carbonic anhydrase (30 kDa), soybean trypsin inhibitor (20 kDa), and α-lactoglobulin (14.2 kDa).

2.8 Hemagglutination assay

Hemagglutination assays were done in microtiter U-plates using serial dilutions with 50 μL volumes of 105 mM NaCl. A 50 μL volume of a 2% suspension of type A human erythrocytes was added and, after 1 hour at room temperature, the results were read. The hemagglutination titer corresponding to the reciprocal of the highest dilution showing hemagglutination was defined as one hemagglutination unit11.

2.9 Inhibitory activity assay

Bovine pancreatic trypsin and bovine pancreatic chymotrypsin were used for the enzymatic assays. Trypsin-like activities were assayed using the N-α-benzoyl-DL-arginine p-nitroanilide (BAPNA) as a substrate. Chymotrypsin-like activities were assayed using N-benzoyl-L-tyrosine p-nitroanilide (BTpNA) as substrate27. In a standard assay, a reaction mixture contained 50 μL of each enzyme extract, reaction buffer (100 mM Tris-HCl buffer, pH 8.0), and 50 μL of 1 mM substrate to a final volume of 500 μL. The reaction was stopped by adding 200 μL of 30% acetic acid. The release of p-nitroaniline groups was measured spectrophotometrically at 410 nm. The proteinase inhibitor was assayed by preincubating 50 μL of fraction at concentrations ranging from 25 to 200 μg with 50 μL of proteinase and 350 μL of reaction buffer at 37 °C for 15 minutes. The reaction was started by adding the substrate and was performed as described above. The remaining activity was expressed as the percentage of enzymatic activity in the absence of an inhibitor.

2.10 Amino acid composition

Amino acid analysis was performed on a PicoTag amino acid analyzer (Waters) as described by HENRIKSON and MEREDITH27. One nanomole of protein fraction was hydrolyzed in 6 M HCl/1% phenol at 106 °C for 24 hours. The hydrolyzed was reacted with 20 μL of fresh derivatization solution (methanol : triethylamine : water : phenylisothiocyanate, 7:1:1:1, v.v –1) for 1 hour at room temperature. After pre-column derivatization, phenylisothiocyanate (PTC) amino acids were identified on a reverse-phase HPLC column by comparing their retention times to those of standard PTC amino acids (Pierce). Cysteine residues were quantified as cystic acid.

2.11 Purification of globulins

Globulins were prepared from sapucaia nuts by the procedure described by MACEDO27. Ground meals, extracted with 50 mM borate buffer at pH 8.0 for 30 min at room temperature, were centrifuged (30 minutes at 8000 x g at 5 °C) and the supernatant proteins were fractioned by ammonium sulfate precipitation. The 70-90% saturation fraction was dialyzed against water, freeze-dried, and applied to a Sephacryl S-200 column (3 cm x 50 cm) equilibrated and eluted with the same buffer used for extraction. The globulin-rich fractions were recovered using an ion-exchange chromatography on a DEAE-Sepharose column (2 cm x 20 cm), equilibrated with 50 mM Tris-HCl at pH 8.0 and eluted with a NaCl gradient (100 mM) in the same buffer. Globulins were dialyzed in water and freeze-dried.

2.12 In vitro digestibility of globulins

Globulins were dissolved in a 100 mM phosphate buffer at pH 6.0 at a 0.5 mg mL–1 concentration. Globulins (250 mL aliquots) were separately assayed for digestion by 10 mL of pepsin (25 mg mL–1 in 50 mM HCl), trypsin (25 mg mL–1 in 100 mM phosphate buffer, pH 7.0), or chymotrypsin (25 mg mL–1 in 100 mM phosphate buffer, pH 7.0), at 37 °C for periods of 15 minutes, 30 minutes, 1, 2, and 4 hours. The substrate-to-proteinase ratio was 20:1. Adding a 10% SDS solution stopped the digestion. The enzymes used for the globulin digestibility assay were purchased from the Sigma Chemical Co. (St. Louis, MO, USA).

2.13 Multienzymatic assays

Globulin digestibility was assayed by the in vitro method described by HSU et al.29. Calculated control (casein) and samples were weighed, dissolved in 10 mL of distilled water and refrigerated at 5 °C for 1 hour. The globulin-containing samples and enzymes were all adjusted to pH 8.0 at 37 °C. Globulin digestibility was determined by digesting the protein-containing sample with a multienzyme mixture—trypsin (porcine pancreatic trypsin—Type IX) with 14 190 BAEE units/mg protein; α-chymotrypsin (bovine pancreatic chymotrypsin—Type II), 60 units/mg powder; and peptidase (porcine intestinal peptidase—Grade III), 40 units/g powder—at 37 °C. A pH drop from 8.0 in the samples was recorded after 20 min of incubation. Globulin digestibility was calculated according to the regression equation (Y = 234.84 – 22.56X, where Y = % digestibility, X = pH drop) described by HSU29. The assays were performed using native and heated globulins.

2.14 Statistical tests

The results were expressed as mean ± S.D., the level of significance was 5% (p < 0.05), when applicable. The data were analyzed using an analysis of variance (ANOVA) (general linear models or GLM procedure)30.

3 Results and discussion

3.1 Proximate composition

Table 1 shows the proximate composition and total calorie content (kcal.100 g–1) of the whole meal. The lipid and protein contents are in accordance with values found in the...
literature (60.61 and 20.47%, respectively) with L. pisonis seeds collected in Brazil, revealing a high energy content (645.05 kcal 100 g⁻¹)¹⁰,³⁸,³⁹.

According to MELLO et al.²⁹, the cashew nut (Anacardium occidentale L.) contains 46.3% of total lipids and 24% of total carbohydrates; the carbohydrate content is significantly higher than that found in sapucaia of this work and of that reported by other authors²⁸,³⁹. However the amount of proteins found for cashew nuts (22%) is similar to the value of the studied sapucaia.

Lipid contents of the Pará nuts (Bertholletia excelsa) (69.3%) are higher than those of the sapucaia (60.61%). However, the protein (16.4%), carbohydrate (3.2%) and fiber (4.6%) contents are lower in the Pará nuts than in the sapucaia nuts, when compared to the values contained in Table 1.

### Table 1. Proximate composition of sapucaia (Lecythis pisonis Camb.) nuts, expressed as g 100 g⁻¹ of raw matter.

<table>
<thead>
<tr>
<th>Component</th>
<th>Results*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>5.04 ± 0.03</td>
</tr>
<tr>
<td>Ash</td>
<td>3.80 ± 0.01</td>
</tr>
<tr>
<td>Crude lipid (ether extract)</td>
<td>60.61 ± 0.33</td>
</tr>
<tr>
<td>Total sugars</td>
<td>4.42 ± 0.23</td>
</tr>
<tr>
<td>Protein (N x 6.25)</td>
<td>20.47 ± 0.38</td>
</tr>
<tr>
<td>Total dietary fiber (by difference)</td>
<td>5.67</td>
</tr>
<tr>
<td>Total calorie content (kcal 100 g⁻¹)**</td>
<td>645.05 ± 2.07</td>
</tr>
</tbody>
</table>

*Mean values ± standard deviation of triplicate determinations; and **Total calorie content was calculated with these factors: 4 for protein and sugars and 9 for lipids according to FAO/WHO.

### 3.2 Fatty acid composition

The fatty acid profile of the oils analyzed (Table 2) indicates a high content of unsaturated acids (monounsaturated, 34.22%; polyunsaturated, 42.73%; omega 3, 0.19%) and a predominance of linoleic (42.54%) and oleic acids (33.94%). The concentration of linoleic acid is in accordance with the levels recommended in BRAZIL⁸ and by the AOCS³ for peanut oil (Brazil, 13.0-45.0%; AOCS, 14.0-43.0%).

Sapucaia nuts were found to be an excellent source of linoleic acid, an essential fatty acid. Moreover, their high lipid content and high level of oil unsaturation indicate their potential use for human consumption, in addition to being a good source of calories in nutritional diets; the data obtained are similar to the values found in the literature²⁸,³⁹. Regarding the quality of oils, the acid contents were within the international standards for the processing of crude vegetable oils for human consumption²⁶.

Comparing the sapucaia nuts to fruit nuts from the Palmae family, the bocaiúva seeds (Acrocomia aculeata (Jacq.) Lodd.)⁹ showed monounsaturated fatty acids (42.5%) and saturated fatty acid (49.7%) values higher than those found for sapucaia nuts, in this study.

Linoleic acid contents in the sapucaia nuts (42.54%) are high when compared to values found for cashew nut (19.6%)²⁹.

### Table 3. Lipid contents and main fatty acid composition of sapucaia (Lecythis pisonis, Camb.) nuts, expressed as g 100 g⁻¹.

<table>
<thead>
<tr>
<th>Fatty acid Results**</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lauric (C12:0)</td>
<td>0.10</td>
</tr>
<tr>
<td>Myristic (C14:0)</td>
<td>0.10</td>
</tr>
<tr>
<td>Palmitic (C16:0)</td>
<td>12.14</td>
</tr>
<tr>
<td>Palmitoleic (C16:1ω7)</td>
<td>0.19</td>
</tr>
<tr>
<td>Stearic (C18:0)</td>
<td>6.31</td>
</tr>
<tr>
<td>Oleic (C18:1ω9)</td>
<td>33.94</td>
</tr>
<tr>
<td>Linoleic (C18:2ω6)</td>
<td>42.54</td>
</tr>
<tr>
<td>cis-11-eicosanoic (C20:1ω11)</td>
<td>0.10</td>
</tr>
<tr>
<td>Alpha linolenic (C18:3ω3x)</td>
<td>0.19</td>
</tr>
</tbody>
</table>

**Saturated | 18.64 **
| Monounsaturated | 34.22 **
| Polyunsaturated | 42.73 **
| Omega 3 | 0.19 **
| Total trans-isomers | ND**

*Mean values ± standard deviation of triplicate determinations; and **Total calorie content was calculated with these factors: 4 for protein and sugars and 9 for lipids according to FAO/WHO.

### 3.3 Mineral composition

The mineral composition (macro and microminerals) of the nuts is shown in Table 3. Compared with the recommended dietary allowances for adults³⁵,³⁶, sapucaia nuts presented high values of nutritionally important minerals required in the human diet.

High levels of Cu, Mn, P, Mg, Fe and Zn were found in sapucaia nut samples, and 100 g of this nut may provide more than 30% of each mineral dietary recommendation intake for adults. Thus, it was classified here as a food rich in these minerals; the levels of Cu and Mn are 3.5 times higher than the dietary reference intake; and regarding the K content, the studied sapucaia nut can be classified as a source of this mineral, for male adults³⁵,³⁶.

Comparing the data obtained in this study to the data from papers with L. pisonis seeds (sapucaia) collected in Brazil³⁸,³⁹, high levels of Cu and significant quantities of Zn and Mn were also observed by other authors.

### Table 3. Macro- and microminerals* of sapucaia (Lecythis pisonis, Camb.) nuts.

<table>
<thead>
<tr>
<th>Element</th>
<th>Sapucaia</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>5.28 ± 0.00</td>
<td>1.200</td>
</tr>
<tr>
<td>Fe</td>
<td>32.65 ± 1.36</td>
<td>8</td>
</tr>
<tr>
<td>Mn</td>
<td>80.69 ± 2.20</td>
<td>2.3</td>
</tr>
<tr>
<td>Zn</td>
<td>40.37 ± 0.38</td>
<td>11</td>
</tr>
<tr>
<td>Cu</td>
<td>32.76 ± 1.14</td>
<td>0.9</td>
</tr>
<tr>
<td>Ca</td>
<td>1.72 ± 0.02</td>
<td>1.200</td>
</tr>
<tr>
<td>Mg</td>
<td>2.79 ± 0.10</td>
<td>420</td>
</tr>
<tr>
<td>P</td>
<td>8.75 ± 0.51</td>
<td>700</td>
</tr>
<tr>
<td>K</td>
<td>8.90 ± 0.04</td>
<td>4.700</td>
</tr>
</tbody>
</table>

*Mean values ± standard deviation of triplicate determinations; and *The mineral dietary reference intake for male adults as expressed in mg d⁻¹.³⁵,³⁶
3.4 Protein fractionation

As shown in Figure 1, albumins are composed of many different polypeptides covering a wide range of molecular masses (18-94 kDa), whereas globulins (major fraction) are essentially represented by four major polypeptides (18, 34, 40, and 50 kDa). Prolamins show fractions ranging from 38 to 50 kDa and the polypeptide composition of glutelins has fractions from 18 to 50 kDa. Several other protein bands were found in each of these fractions.

![Image of molecular mass marker and protein bands](image)

Figure 1. Polypeptide patterns of molecular mass marker (lane 1), crude extract (lane 2), globulins (lane 3), albumins (lane 4), glutelins (lane 5) and prolamins (lane 6) of sapucaia (Lecythis pisonis, Camb.) nuts.

The protein content of sapucaia defatted nut meal determined by BRADFORD was 66%. In the present investigation, globulins made up the major protein fraction component of sapucaia nuts (58.7%), whereas glutelins, albumins and prolamins accounted for 20.2, 20.1, and 1.0%, respectively. Table 4. The globulin fraction showed a notably high protein concentration (84%) when compared with the other fractions (data not shown), and was thus chosen for the purification and assays conducted in the present work.

Table 4. Protein fractions of sapucaia (Lecythis pisonis, Camb.) nuts, by solubility.

<table>
<thead>
<tr>
<th>Protein fractions</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Globulins</td>
<td>58.7</td>
</tr>
<tr>
<td>Glutelins</td>
<td>20.2</td>
</tr>
<tr>
<td>Albumins</td>
<td>20.1</td>
</tr>
<tr>
<td>Prolamins</td>
<td>1.0</td>
</tr>
</tbody>
</table>

3.5 Amino acid analysis

The quality of seed proteins as sources of amino acids can usually be evaluated by comparing the FAO/WHO recommended standards for essential amino acids. As shown in Table 5, the proteins from sapucaia nuts contained adequate levels of phenylalanine, lysine, leucine, methionine, valine, and arginine, and the other amino acids were found in high or moderate amounts, based on the FAO/WHO standards for children. The proteins also contained adequate amounts of essential amino acids for pre-school children and all the essential amino acids for adults. The concentration of tryptophan was not determined. When the amino acid content of sapucaia nuts is compared with that of animal proteins from eggs, cow's milk or beef, sapucaia nuts are found to be an excellent amino acids source.

3.6 Antinutritional factors

Protein quality is affected by antinutritional factors that interact with cells of the intestinal tract, such as proteinase inhibitors, lectins, and tannins, which reduce protein digestibility and amino acid absorption. Unless destroyed or inactivated by heat or by some other suitable treatment, these substances can exert adverse physiological effects when ingested by man and animals. In the present study, no hemagglutination or inhibition were observed in any of the samples assayed (data not shown), revealing low or non-detectable levels of lectin and proteinase inhibitors and demonstrating that the nuts analyzed were free of these major antinutritional factors.

This is a relevant finding, because feeding raw soybean and many other legume products, which contain high levels of proteinase inhibitors, to experimental animals such as rats, mice and chickens leads to growth depression, pancreatic hypertrophy, and/or hyperplasia and a potentiation of pancreatic carcinogenesis. Most of these compounds inhibit the digestive enzymes or react with essential amino acids, limiting the use of whole seeds in food products. Lectins bind to the intestinal mucosa, impairing digestion and absorption of nutrients and reducing protein digestibility by inhibiting digestive enzymes.

3.7 In vitro digestibility of globulins

The in vitro digestibility of sapucaia nut globulins by mammalian digestive proteinases was carried out using trypsin, chymotrypsin and pepsin, separately. Incubation of purified globulins showed that trypsin digested the 18 and 66 kDa fractions, but the globulins were resistant to hydrolysis by chymotrypsin or pepsin.

After heat treatment, however, the 50 and 66 kDa fractions were digested by chymotrypsin and the 18, 50, and 66 kDa fractions were hydrolyzed by trypsin, although no hydrolysis by pepsin was observed on SDS-PAGE. Trypsin showed hydrolytic activity on both native and heated globulins. These results were compatible with previous findings that globulins are resistant to hydrolysis by pepsin.

3.8 In vitro digestibility by multienzymatic assays

Figure 4 shows the SDS-PAGE patterns of native and heated globulins digested by multienzymes. The electrophoretic pattern of their in vitro digestibility is shown in Figure 5.
In vitro digestibility of sapucaia globulins

**Figure 2.** SDS-PAGE patterns of digestion of native globulins of sapucaia (*Lecythis pisonis*, Camb.) nuts by trypsin, chymotrypsin and pepsin, separately. a) Digestion by trypsin; b) Digestion by chymotrypsin; and c) Digestion by pepsin. Vertical numbers indicate molecular weight marker in kDa. Horizontal numbers refer to times of digestion.

**Figure 3.** SDS-PAGE pattern digestion of heated globulins of sapucaia (*Lecythis pisonis*, Camb.) nuts by trypsin, chymotrypsin and pepsin, separately. a) Digestion by trypsin; b) Digestion by chymotrypsin; and c) Digestion by pepsin. Vertical numbers indicate molecular weight marker in kDa. Horizontal numbers refer to times of digestion.
Table 5. Amino acid composition of proteins from sapucaia (Lecythis pisonis, Camb.) nuts (mg·g⁻¹ protein).

<table>
<thead>
<tr>
<th></th>
<th>Albumins</th>
<th>Glutelins</th>
<th>Globulins</th>
<th>Prolamins</th>
<th>Total proteins</th>
<th>Requirement standard*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>10.0</td>
<td>32.2</td>
<td>44.2</td>
<td>1.4</td>
<td>87.8</td>
<td>-</td>
</tr>
<tr>
<td>Arginine</td>
<td>2.4</td>
<td>57.6</td>
<td>108.6</td>
<td>0.5</td>
<td>169.1</td>
<td>-</td>
</tr>
<tr>
<td>Aspartate</td>
<td>11.7</td>
<td>38.7</td>
<td>56.1</td>
<td>1.5</td>
<td>110.0</td>
<td>-</td>
</tr>
<tr>
<td>Cystine</td>
<td>-</td>
<td>3.7</td>
<td>11.7</td>
<td>-</td>
<td>15.4</td>
<td>25**</td>
</tr>
<tr>
<td>Glutamate</td>
<td>36.4</td>
<td>83.8</td>
<td>153.3</td>
<td>1.8</td>
<td>275.3</td>
<td>-</td>
</tr>
<tr>
<td>Glucose</td>
<td>39.0</td>
<td>43.2</td>
<td>73.2</td>
<td>1.7</td>
<td>157.1</td>
<td>-</td>
</tr>
<tr>
<td>Histidine</td>
<td>-</td>
<td>7.4</td>
<td>11.8</td>
<td>0.1</td>
<td>19.3</td>
<td>19</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>2.7</td>
<td>10.6</td>
<td>15.8</td>
<td>0.4</td>
<td>29.5</td>
<td>28</td>
</tr>
<tr>
<td>Leucine</td>
<td>1.0</td>
<td>42.8</td>
<td>69.0</td>
<td>1.0</td>
<td>113.8</td>
<td>66</td>
</tr>
<tr>
<td>Lysine</td>
<td>28.4</td>
<td>20.7</td>
<td>31.9</td>
<td>1.1</td>
<td>82.1</td>
<td>58</td>
</tr>
<tr>
<td>Methionine</td>
<td>-</td>
<td>30.1</td>
<td>59.2</td>
<td>0.3</td>
<td>89.6</td>
<td>-</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>77.2</td>
<td>12.5</td>
<td>16.6</td>
<td>0.3</td>
<td>106.6</td>
<td>63***</td>
</tr>
<tr>
<td>Proline</td>
<td>117.5</td>
<td>39.3</td>
<td>70.0</td>
<td>2.2</td>
<td>229.0</td>
<td>-</td>
</tr>
<tr>
<td>Seryne</td>
<td>7.2</td>
<td>31.5</td>
<td>46.9</td>
<td>0.9</td>
<td>86.5</td>
<td>-</td>
</tr>
<tr>
<td>Threonine</td>
<td>3.2</td>
<td>13.2</td>
<td>17.3</td>
<td>0.7</td>
<td>34.4</td>
<td>34</td>
</tr>
<tr>
<td>Triptophan</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>11</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>1.7</td>
<td>12.0</td>
<td>18.3</td>
<td>0.5</td>
<td>32.5</td>
<td>-</td>
</tr>
<tr>
<td>Valine</td>
<td>4.7</td>
<td>22.4</td>
<td>34.1</td>
<td>0.8</td>
<td>62.0</td>
<td>35</td>
</tr>
</tbody>
</table>

*FAO/WHO; **Cystine + methionine; ***Phenylalanine + tyrosine; Tryptofan was not determined – ND. Essential amino acids are in bold letters.

Figure 4. In vitro digestibility of sapucaia (Lecythis pisonis, Camb.) nut globulins by multienzymes (trypsin, chymotrypsin, and peptidase) in comparison with casein (mean ± SD, n = 3). Black columns: native protein; white columns: heated samples. Experimental error is indicated by standard deviation bars. The effect of heating on protein digestibility was statistically insignificant, according to ANOVA (p < 0.05).

Figures 4 and 5 reveal that the multienzymatic complex was efficient in digesting globulins. Digestibility was as high as 70.30% (Figure 4). Heating of globulins for 10 minutes led to an insignificant increase in digestibility, to 71.35%. The increased digestibility of sapucaia globulins by multienzymes suggests that digestive enzymes may have a joint action, making all the bonds more accessible to proteases. The low increase in digestibility after heating suggests that sapucaia globulins can be ingested as fresh protein as the nuts do not contain antinutritional factors such as lectins and protease inhibitors.

Figure 5. Electrophoresis showing in vitro digestibility of native and heated globulins of sapucaia (Lecythis pisonis, Camb.) nuts in a multienzymatic assay (trypsin, chymotrypsin, and peptidase); molecular mass marker (Lane 1), casein (lane 2), casein + multienzymes (lane 3), globulins (lane 4), and globulins + multienzymes (lane 5).

4 Conclusions
This study revealed that sapucaia nuts are a valuable source of proteins, with higher levels of essential amino acids, fatty acids and minerals than the recommended ones. No hemagglutinating or inhibitory activities were observed in any
of the samples investigated, indicating low or non-detectable levels of lectin and proteinase inhibitors, thus demonstrating that sapucaia nuts are free of these antinutritional factors. In addition, the in vitro digestibility of globulins by a multienzymatic complex was significant. These observations suggest that sapucaia nuts may be a new source of proteins for human consumption, a potential functional and nutritional agent, and an economically important oil source.

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References


