CHEMICAL AND BIOLOGICAL STUDY OF EUGENIA STIPITATA Mc VAUGH COLLECTED IN THE COLOMBIAN ANDEAN REGION

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INTRODUCTION

The exotic fruit of arazá, Eugenia stipitata (Myrtaceae), is originally from Western Amazon, in the region bordered by Marañon and Ucayali rivers, and the proximities of the Requena Province and Amazon river source [1,2]. Mc Vaugh [3] verified the existence of two different populations of E. stipitata, but with common qualitative features which forced to consider them as subspecies: Stipitata and sororia; this latter is grown in Colombia, Peru, Brazil, Ecuador, Bolivia, and Costa Rica [1,3].

The attractive organoleptic features of the E. stipitata fruit, along with the knowledge of its nutritional properties and production systems, have made the Amazon Institute of Scientific Investigation of Colombia turn it into one of the species prioritized in its strategic line of Development of Productive Alternatives and Green Markets [4]. However, its cultivation and development are not exclusive from the Amazon; there are also monocrops in the departments of Meta, Guaviare, Caquetá, and Putumayo; with lesser frequency, it is located in some departments of the Andean region such as Cundinamarca, Valle del Cauca, Caldas, Antioquia, and Tolima [5,6].

The arazá fruit shows chemical characteristics, intense aroma, and a sweet-sour taste being very appreciated in agroindustrial processes, which has motivated the performance of multiple studies with the purpose of knowing its harvest physiology, ripening process, and post-harvest technologies [6,7]. Nutritional interest studies are also remarkable [4] as well as technological viability of frozen pulp [8], fruit commercialization perspectives [9-11], industrial potentiality [7,12], antioxidant capacity [1,13,14], antimutagenic agent [15], and anti-diabetic [16]; this latter, related to the high content of compound derivatives from quercetin, which are molecules capable of inhibiting the activity of α-glucosidase, an enzyme involved in carbohydrates metabolism.

The attributes mentioned above, as well as the high yield in the pulp of the ripe fruit, would make E. stipitata subs. sororia an important source of basic material for Colombian agroindustry. However, some difficulties in fruit post-harvest handling and the lack of quality technical normativity, among others, have created obstacles in the progress of production line of arazá, in comparison with other fruits related to it, such as guava (Psidium guajava) [1,11,10,17]. Even though most researches have been oriented to the knowledge of the pulp and/or peel [13,14,17-19], leaving aside the biotechnological and/or pharmacological potentialities of the seed, which takes 30% of the fruit volume, it is 22% of fresh weight and 84% of dry matter. The interest of farmers, agricultural technicians, and professionals in regions where there are arazá plantations has made that, until now, studies related to the seeds are fundamentally of agronomic type, although they are mentioned as an important part for the plant growth.

Taking into account the information given above, this study was focused on evaluating the chemical and bioactive properties of the fruit seeds and the organic fractions derived from them. All the data obtained will provide an enhanced knowledge of the promising nutritional/pharmacological value of this product as a social and economic alternative for rural populations in Colombia.
MATERIALS AND METHODS

Chemical reagents and biological material
All chemicals used in analysis as 2,2-diphenyl-1-picrylhydrazyl (DPPH), 3,5-dichloro-1,4-benzoquinone-6-sulfonic acid (ABTS), digitonin, hydrogen peroxide, and 3-(4,5-dimethylthiazol-2-yl)-2,5-tetrazolium diphenyl bromide were purchased from SIGMA-ALDRICH® (St. Louis, MO, USA). The chromatographic solvents were acetone, triluque chromatography-mass spectrometry (LC-MS) grade from J.T. Baker (Deventer, The Netherlands) and water purified in a Milli-Q system from Millipore (Bedford, MA, USA). The other chromatographic chemicals used were high-pressure LC grade supplied also from J.T. Baker. Other chemicals employed were analytical grade (Merck, USA). Free life Panagrellus redivivus nematode strain was provided by the Titaatita research center from Corpoica, Colombia. Gastrointestinal (GI) nematodes were collected from stools taken directly from the rectal ampoule of ruminants at the Northern Regional University Center, University of Tolima, Armero, Tolima (5° 00’ 08.10” N, 74° 54’ 51.33’” O). The permission to use living organisms and/or human samples was authorized by the Ethics Committee of the University of Tolima.

Collecting area
Arazá fruits (E. stipitata Mc Vaugh) were collected in the rural area of the municipality of San Sebastián de Mariquita (05° 12’ 12” N, 74° 53’ 46” O). The region is a bioclimatic transition zone between humid to very humid weather, with an annual precipitation level of 2194 mm, equatorial climate and temperatures between 23°C and 28°C. The land corresponds to the geostucture of the Andean Central Mountain Range and includes the morphostructural sets of the Eastern flank or the Eastern Mountain Range, showing an elevation of 350–1400 m a.s.l. [20]. Mariquita is, nowadays, a developing center being remarkable in the fruit sector of the Tolima Department (Colombian Andean Central Area).

Collection and preparation of plant material
Fruit material, in optimal phytosanitary condition, was collected in February 2016 in small monocrops surrounding Mariquita Municipality. A complete specimen (leaves, flowers, and fruit) was prepared and determined in the National Herbarium of Colombia; a voucher is deposited there. A fraction of the seed, manually separated from the fruit, was submitted to a drying process (45±20°C, 24 h). Then, it was crushed, degreased with n-hexane by Soxhlet extraction method and macerated with ethanol (96%), renewing the solvent after 2 h until depletion of the sample. Raw ethanolic extract (REE) was fractioned liquid-liquid with dichloromethane, obtaining a fraction (DF) and a hydroalcoholic residue (HR). The solvent was vacuum-evaporated (45°C, Rotavapor R-114, Büchi, Flawil, Sweden) and REE, DF, and HR were stored (−85°C, Freezer Kaltis 390°) until further analysis.

Mineral and bromatological analysis
The moisture content, ash, crude fat, crude protein, and crude fiber were determined on a portion of the dry seed. Similarly, major (K, Na, Mg, and Ca) and minor (Fe, Mn, Cu, and Zn) minerals were determined by the atomic absorption spectrophotometric method (ThermoScientific ICE 3000 Series AA), while P, B, and S were quantified by UV-vis spectrophotometry (Evolution 260 BIO). All the determinations were performed under the AOAC standard indications [21]. Total sugar was estimated according to the method described by Nelson [22].

Physicochemical characterization of extract and fractions
REE, DF, and HR were standardized by some physicochemical determinations (color, pH, density, and Brix) and a preliminary phytochemical analysis, attending Harborne’s indications [23]. Besides, in every plant product was measured total phenols and non-tannic phenols content by applying Folin–Ciocalteu method with some modifications reported by Fidrinnay et al. [24]. The tannins content was established as the difference between them [25]. The results were expressed in grams of gallic acid equivalent per every hundred grams of sample (g GAE/100 g).

For REE, the chemical profile was carried out using a high-performance liquid chromatographic analysis on a Waters Alliance 2695 separation module system, coupled to a double canal λ absorbance (280 and 320 nm) detector. Compounds were separated with a Waters Atlantis dC18 column (5 µm; 2.1 mm; 150 mm), using a gradient system with a mobile phase which consisted in H₂O (solvent A, 0.1% v/v formic acid) and methanol 100% (solvent B, 0.1% v/v acetic acid). The flow rate was 0.500 ml/min. A linear gradient elution was performed by varying the proportion of solvent A to solvent B, according to Delpino-Rius et al. [26]. The data were acquired using the chromatographic behavior and UV-vis absorption spectra, together with published data on the main phenolic compounds in the samples. To quantify, calibration curves were built tracing chromatographic peak areas measured in the specific wavelengths of the chemical standards (catechin, epicatechin, procyanidin B1, procyanidin B2, eriodictyol-7-O-rutinoside, quercetin-3-O-galactoside, quercetin-3-O-glucoside, quercetin-3-O-rhamnoside, kaempferol-3-O-rutinoside, isorhamnetin-3-O-rutinoside, arbutin, gallic acid, caffeic acid, p-coumaric acid, ferulic acid, sinapic acid, gallic acid, isoquercitrin, chlorogenic acid, quercitrin, rutin).

Antioxidant and antiradical capacity
Antiradical capacity (the ability of compounds of the tested samples to react with free radicals) was evaluated using DPPH⁺ [27] and ABTS⁺ [28] methods. In terms of the radical-scavenging ability of REE, DF, and HR, the inhibitory concentration 50% (IC₅₀) value (sample concentration that stabilizes in a 50% DPPH⁺ or ABTS⁺ radical, the antioxidant potential (ºP=1/IC₅₀ expressed in percentage)), and Trolox-equivalent antioxidant capacity (TEAC) were established [29]. At that time, the antioxidant capacity (ability to inhibit the process of oxidation) was determined using four in vitro test systems, that is, the oxygen radical absorbance capacity (ORAC), the ferric reducing ability of plasma (FRAP), and OH radical inhibition (30-32), complemented with anti-hemolysis effect on erythrocyte suspension from healthy volunteers [33]. All spectrophotometric measurements were performed with a 96-hole micromplate reader UV-vis (Multiskan® G Thermo scientific).

In vitro anthelmintic assays
The anthelmintic efficacy of REE, DF, and HR against free-living (FL) nematodes (P. redivivus) and GI nematodes was determined. For the culture and maintenance of P. redivivus, the methodology reported by Pica was followed [34]. The GI nematode genres collected were determined according to their main morphological features with foundation on the information stated by Van Wyk and Mayhew [35].

Toxicity over P. redivivus
Toxicity over P. redivivus was estimated by separating the adult organisms from the larvae with a sieve (150 µm). The sample was added (400 µl) to a suspension with 50 larvae (leaving a final volume of 900 µl with isotonic buffer). The sample was replaced with buffer (negative control) and ethanol at 96% (positive control). Treatments were incubated (26°C, 24 h) and the mortality percentage (M%) was established using the following formula:

\[ M\% = \frac{DL}{50} \]

where DL is the number of dead larvae. With this data, IC₅₀ lethal concentration for 50% of larvae was determined [34].

Collection of GI nematodes eggs
Eggs were collected from the feces analyzed using the conventional method of McMaster reported by Paolini et al. [36]. Fecal matter of sheep infected with strains of GI parasites was freshly collected and weighed. First of all, 1 g of feces was analyzed to calculate the number of eggs in the sample (N1). The initial amount of eggs (N) contained in 1 g of feces was estimated by separating the adult larvae from the eggs (N1) by the McMaster method reported by Paolini [24]. The eggs were suspended with 50 ml of isotonic buffer (0.9% NaCl) to a suspension with 50 larvae (leaving a final volume of 50 ml with isotonic buffer). The suspension was then poured through a series of sieves decreasing mesh (200 µm, 363
125 μm, and 40 μm). The residue on the final sievel was collected in a small volume of water and centrifuged. The pellet was added to a saturated sodium chloride (NaCl) solution and then centrifuged again. The supernatant was then filtered and the residue was washed three times following by centrifugation. The resultant pellet was added to a known volume of sterile distilled water.

**GI nematodes egg hatch assay**
Approximately, 100 eggs suspension and 10 μl of the sample were leaving to a final volume of 2 ml with sterile distilled water. Treatments were incubated (30°C, 24 h) and non-hatched number was determined. Benzimidazole solution (400 μg/ml) was used as a positive control and distilled water as negative control. With the obtained information, IC50 (concentration of sample for 50% egg hatching inhibition) was calculated.

**GI nematodes cuticle loss inhibition assay**
The cuticle loss inhibition was evaluated on organisms in the L3 stage [37]. A suspension of approximately 1200 L3 was applied to the sample (30 μl) and the volume was brought to 10 ml with sterile distilled water. Albenazole solution (400 μg/ml) was used as a positive control and sterile distilled water as negative control. Treatments were incubated (22°C, 3 h) and centrifuged (4200 rcf, 2 min); the washing was repeated 3 times with sterile distilled water, discarding 8 ml of the supernatant every time. Each solution treatment (100 L3, 100 μl of the sample, and 250 μl final volume) per well was exposed to sodium hypochlorite (NaClO, 1.5 ml at 0.187%, 10 min) action, a drop of lugol was added and the number of organisms with and free cuticle was counted. Cuticle loss inhibition percentage (CLI%) was calculated using the following equation:

\[ CLI\% = \frac{WC \times 100}{WC + FC} \]

where WC and FC is the number of organisms with cuticle and free cuticle, respectively. With the obtained values, IC50 (50% L3 cuticle loss inhibitory concentration) was calculated.

**GI nematodes motility and mortality inhibition assay**
To verify the effect of the samples phytoconstituents on GI nematodes, motility inhibition and mortality assays were conducted. To eliminate the larval cuticle, 2 ml of L3 suspension was exposed at NaClO (0.187%, 5 min), the treatments were centrifuged (470 rcf, 4°C, 5 min) discarding 1.8 ml of the supernatant; the washing was repeated 3 times with sterile distilled water. The 100 μl of L3 larvae solution were placed in a microplate and equal volume sample was added (250 μl final volume per well). Ivermectin was used as a positive control and sterile distilled water as negative control. The treatments were incubated (26°C, 24 h) and anthelminthic action was determined [38]. The alive (rolled) and dead (immotile) L3 stage larvae were counted in each well. Larval motility inhibition percentage (LM1%) was calculated using the following formula:

\[ LM1\% = \frac{RL \times 100}{TL} \]

where RL is the number of alive rolled larvae in test samples and TL is the total of larvae. With the obtained values, IC50 (motility inhibitory concentration for 50% of L3) was calculated. Mortality percentage (M%) was calculated using the following equation:

\[ M\% = \frac{DL \times 100}{TL} \]

where DL is the number of dead larvae and TL the total of larvae. Using the previous data, (LC50 of L3) was determined.

**Cytotoxicity on human peripheral blood mononuclear cells**
The human blood samples were obtained from healthy volunteers. For leukocyte isolation, the protocol proposed by the Viral and Human Genomics Lab from the Medicine of School from San Luis Potosí Autonomous University [39] was used. Cytotoxic potentials of REE was performed according to the colorimetric model with MTT [40]. The sample (25 μl, 4000 mg/ml), the suspension of leukocytes (25 μl), and MTT tetrazolium salt (50 μl, 0.04 mg/l dissolved in PBS pH 7.4) were added to each well of Microplate reader (Multiskan® GO Thermo scientific), and mixed thoroughly (1 min). After plate incubation (37°C, 2 h), dimethylylphosphate (DMSO, 100 μl) was added to each well and mixed thoroughly in order to dissolve the formed blue crystals of formazan. Dilution solvent was used as sample blank, a digitonin solution (1%) as negative control and cells without extract treatment as a positive control. All plates were then shaken for 10 min and incubated for 2 h. The optical density of wells was measured by microplate reader at wavelength 570 nm [41]. Cell viability (CV%) was calculated as % of the metabolic activity of sample compounds using the following formula:

\[ CV\% = \frac{(AbsT - AbsB) \times 100}{AbsC - AbsB} \]

where AbsT is treatment absorbance; AbsB, sample blank absorbance; AbsC, positive control absorbance.

**Toxicity on human erythrocytes**
The toxicity of plant samples was tested under in vitro conditions on human blood cells [33]. An erythrocyte solution (400 μl at 5% in PBS, pH 7.4) was added to REE, DF, or HR solution (40 μl, 1500–7500 mg/l). Dilution solvent (40 μl) was used as sample blank; negative control was carried out inoculating erythrocytes with PBS (40 μl), whereas digitonin solution (1% w/v) for cytotoxicity positive control was used. Every treatment was adjusted to 1 ml final volume with PBS, incubated (37°C, 20 min), and centrifuged (500 rcf, 5 min). Thereafter, the absorbance of the supernatant (200 μl) was read at 540 nm. Hemolysis percent (H%) was calculated using the following equation:

\[ H\% = \frac{AbsC - (AbsT - AbsB) \times 100}{AbsC} \]

where AbsC is the positive control absorbance; AbsB is sample blank absorbance; AbsT is treatment absorbance.

**Statistical analysis**
The reported values represent the analysis of at least three separate replicates for each sample ± standard deviation. Statistical analysis of the data was done using one-way variance analysis (ANOVA). Values of p<0.05 were considered statistically significant, to which variance normality and homogeneity assumptions were verified (p>0.01). The IC50 values were calculated using linear regression analysis. All the processing was performed with the STATGRAPHICS Centurion XVII Package.

**RESULTS AND DISCUSSION**

**E. stipitata Mc Vaugh fruit** is characterized by being a globose berry (30–800 g) with mild pubescence on the epicarp and because the edible part (bittersweet flavor) holds an approximate number of 20 monoembryonic seeds (0.3–1.5 cm, diameter; 0.3–2.5 cm, length; 0.1–4.3 g, weight; they are 84% of dry matter), distributed in typical form along the internal cavity of the fruit (axial placemant). According to Akamine and Goo [42], fruits of Eugenia genus are not climacteric. However, Galvis and Hernández [43] argue the contrary, even though the total production of ethylene in arazá still is unknown [1].

**Mineral and bromatological analysis**
The results of the nutritional composition of arazá whole seed, in relation to their edible part, are presented in Table 1. A critical look at Table 1 shows that E. stipitata seeds recorded a higher value of nutrient contents than the edible part. Therefore, nutrient profile of
<table>
<thead>
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<th>Parameters</th>
<th>Nutrient content percentage</th>
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<tr>
<td></td>
<td>Seed</td>
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<tr>
<td>Moisture</td>
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<tr>
<td>Dry matter</td>
<td>71.20±1.80</td>
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<tr>
<td>Ashes</td>
<td>1.35±0.05</td>
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<tr>
<td>Crude fat</td>
<td>2.05±0.01</td>
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<tr>
<td>Crude protein</td>
<td>3.06±0.07</td>
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<tr>
<td>Crude fiber</td>
<td>5.9±0.04</td>
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<tr>
<td>Total carbohydrates</td>
<td>31.12±1.49</td>
</tr>
<tr>
<td>Major elements (%)</td>
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<tr>
<td>Calcium</td>
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<tr>
<td>Magnesium</td>
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</tr>
<tr>
<td>Sodium</td>
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</tr>
<tr>
<td>Potassium</td>
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<tr>
<td>Phosphorus</td>
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<tr>
<td>Minor elements (mg/kg)</td>
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<tr>
<td>Zinc</td>
<td>4.90±0.31</td>
</tr>
<tr>
<td>Boron</td>
<td>41.80±0.80</td>
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<tr>
<td>Sulfur</td>
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<tr>
<td>Copper</td>
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<tr>
<td>Iron</td>
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</tr>
<tr>
<td>Manganese</td>
<td>1.40±0.40</td>
</tr>
</tbody>
</table>

*Taken from Barrantes, Yaya, and Arias [45] and Rogers et al. [17].

NR: Non-reported.

arazá seed shall be established as: Carbohydrates (31.12%)>crude fiber (5.98%)>sulfur (100 mg/kg)>boron (41.8 mg/kg)>iron (20.9 mg/kg) and copper (16.2 mg/kg). Such nutritional contributions, added to the succulence, aroma, flavor, texture, and color of the edible portion, make of arazá an appetizing fruit option. The basic chemical composition of the tested seeds was higher to that previously reported by P. guajava fruit, a Myrtaceae of high commercial importance [44].

On the other hand, the moisture content for arazá seed (38.80%) was lower when compared with the edible part (96%). Coriaceous and thin tegument which covers the seed would explain in part the low humidity content of them. Consequently, it must be understood that over 50% are nutrients stored as dry matter in the seed. Moisture content is among the most vital and mostly used measurement in the processing, preservation, and storage of food [46].

It is also of notice the considerable difference in carbohydrate content between the seed and the edible part (16.1); this has a foundation if it is considered that carbohydrates are part of the material that satisfies cell energetic demands for the seedling support. Many primary metabolites act as precursors or are pharmacologically active metabolites in bioactive extracts. Carbohydrates are the main energy reserves in foods of plant origin; they are precursors of certain lipids, proteins, ascorbic acid, and inositol [46], among other functions. In the case of arazá seed, a significant part of these metabolites would be constituted by tegument covering the seed, which possibly helps seed dispersion and fixation on the substrate. Crude fiber in food or plant is an indication of the level of non-digestible carbohydrate and lignin. The value obtained (5.98%) for seeds are much higher than that found for the edible part (0.004%). Crude fiber is the sum total of all those organic compounds of the plant cell membranes and supporting structures which in chemical analysis of plants foodstuff remain after removal of the crude protein, fat, and nitrogen-free extract; they are commonly used as an index of the value in animal feeding [47].

Some researchers state that sulfur, despite being considered a secondary element, is required in plants in levels similar to phosphorus and that also is a nutrient as necessary as nitrogen [47,48]. Boron is an important micronutrient for the formation of cell walls given that it is necessary for pectin synthesis. Likewise, iron is constituent of cytochromes and metalloenzymes; it has roles in photosynthesis, N symbiotic fixation, N metabolism, and redox reactions; meanwhile, copper is cofactor of several enzymes with roles in photosynthesis, respiration, carbohydrate, and protein metabolism, among others. Iron is useful for the prevention of anemia and other related diseases and Zinc plays a role in protein synthesis, normal body development, and recovery from illness [47-49]. The results of the chromatographic composition suggest that the flour from arazá seeds could be utilized as complementary food.

Physicochemical characterization of the extract and fractions

The systematic screening of plant species with the purpose of discovering new bioactive compounds is a routine activity in many laboratories which beginning with the extraction (as the term is pharmaceutically used), using selective solvents of increased polarity and characterization of the extracts obtained [50]. The products from the seeds were relatively complex mixtures of metabolites. The amounts of polar metabolites recovered from arazá seed varied depending on the solvent. The extraction yield for REE was high (84.4%) when compared with 5.54% and 2.9% for HR and DF, respectively. Nevertheless, this value could be considered lower than reported for some Myrtaceae fruits; e.g., in Psidium guajava peel was found 29.53% while the leaves of Psidium guineense revealed a 12.13% [51]. On the other hand, physicochemical features of REE and fractions (DF and HR) showed little variation among them; for example, it was observed in all a brown coloration and densities (1.35–1.64 g/cm³), brix (19.8–20.4), acidity (1.07–1.31% citric acid), and pH (5.5–5.6) values very similar. The basic parameters influencing the yield and quality of an extract are plant portion and solvent used, location, and extraction procedure, among others.

The evaluation of all the drugs is based on physico-chemical and phytochemical approaches which lead to the drug discovery referred as natural product screening. The preliminary qualitative phytochemical screening was carried out on the REE which showed the presence of phenolic compounds, especially tannins. Less abundant were flavonoids, terpenoids, steroids, and saponins. The absence of reducing carbohydrates was noticeable; this is due, at least partially, at the high metabolic rate of the seed at the time of harvest of the fruit in preparation for its germination. The quantitative estimation of phenolic constituents revealed 29.57 g of GAEs per every 100 g of REE (g GAE/100 g). This answer not only confirms what was revealed by the qualitative tests but also shows a higher value than the one found by Neri-Numa et al. [15] in arazá pulp ethanolic extracts (18.40 g GAE/100 g) and to what was reported by Garzón et al. [14] for the peel and the pulp (12.43 and 19.3 mg GAE/100 g, respectively). Castro, Restrepo, and Parada [52] determined in the seeds of P. guajava 176, 72, and 131 mg GAE/100 g applying Soxhlet method, CO2 supercritical, and CO2 supercritical-ethanol, respectively. At the same time, HR and DF showed a value of 17.56 and 13.45 g GAE/100 g, respectively. Matching total phenolic content, the tannic material showed the following order: REE > HR > DF, with values of 29.14, 17.09 and 13.08 g GAE/100 g.

The present investigation shows higher values in the contents of these metabolites when compared to above-mentioned results, suggesting that arazá seeds have phenols of diverse chemical nature. Non-tannic phenols (simple phenols, phenylpropanoids, flavonoids, and phenolic acids among others), followed the same pattern (0.42, 0.23 0.18 g GAE/100 g) and to what was reported by Garzón et al. [14] for the peel and the pulp (12.43 and 19.3 mg GAE/100 g, respectively). It could be thought that in the seed of E. stipitata there is the presence of hydrolyzable and non- hydrolyzable tannins (condensed tannins or proanthocyanidins) which, due to their diverse polarity and/or molecular size, showed an uneven distribution in HR and DF [53]. Tannins are found in foliage tissue, flower buds, stem, roots, and seeds; in this latter, are located mainly on a layer between the external tegument and the aleuronic layer; they have been associated to the maintenance of plant dormancy and have allelopathic and bactericide properties [54].

Phenols are the secondary metabolites of relevant importance because of their contribution to human health, besides their widely known and
recognized as natural antioxidants [55]. In addition, tannins are used in dyeing and industries of wine, beer, tea, and cocoa in pharmacy. They are applied as antidiarrheal [56], antiviral [57], enzymatic inhibitors, and hepatoprotectors [58], angiotensin receptors II AT I [59], lipid peroxidation inhibitors [60], and nitric oxide-dependent vasodilators [61].

To establish the chemical fingerprinting of the extract from seed, 22 authentic samples of phenolic compounds (proanthocyanidins, flavonoids, glycoside flavonoids, hydroxyacinamic acid derivatives, and phenolic acid) were used as reference compounds, which exhibited: Isoquercitrin (0.083%)=>gallic acid (0.055%)=>phloridzin (0.026%)=>sinapic acid (0.023%)=>p-cumaric acid (0.017%) as the main constituents; some of which are recognized as antioxidant and oncoprotective activities [62]. Phloridzin reduces glucose absorption, blocking transporters of this molecule into the small intestine, causing a hypoglycemic effect [63]. Despite the chemical attributes found, arazá in Colombia is still a plant of small gardens with low commercialization. The physicochemical characterization of arazá seeds is not been reported earlier.

Antiradical and antioxidant capacities

Due to the large diversity of antioxidant compounds of REE, DF, and HR, the results of antiradical capacity in this study are given as total antioxidant capacity. The data were uniformly calibrated using the trolox as a reference compound (TEAC). The values of this variable are less influenced by the chemical backbone (main structural elements distinguishing phenolic compound subgroups), than by functional groups attached to the main structure [64]. The comparative assessment showed significant differences between the tested samples as well as between the assays used. There is a great variation in the level of IC50 and AP values (Table 2).

It can be clearly seen that DF and HR displayed no antioxidant capacity significantly higher than REE (p>0.05). Likewise, both fractions showed the lowest IC50, and highest AP values by ABTS assay than the DPPH method (p>0.05). The different AP percentage obtained from the assays may reflect a relative difference in the ability of antioxidant compounds of DF and HR to quench and reduce aqueous radicals. TEAC results (Table 2) confirm the IC50 and AP values. ABTS and DPPH methods have the same mechanism reaction that was electron transfer assays. Phenolic antioxidants (ROH) can donate hydrogen atoms to radicals; then, the presence of total phenolics might contribute to the antioxidant capacity of REE, HR, and DF. In this work, the data illustrated higher total phenolic and tannins values in REE, but its antiradical capacity, by DPPH and ABTS tests, was lower, suggesting an indirect correlation between these variables. ANOVA analysis of one way, applied to the analytic determinations related to antiradical capacity, verified variance normality, and homogeneity assumptions (p>0.01).

Garzón et al. [14] determined in arazá peel 9.0 and 2.0 µmol trolox/g FW by DPPH and ABTS assays, respectively; while Brand-Williams, Culever, and Berset [64] determined inedible part 11.0 and 2.6 µmol trolox/g FW by two radical. Based on the above data could be seen that arazá seeds have higher antiradical capacity than peel and edible part.

Weng and Huang [29] state that there are five factors determining in antioxidant capacity of the phenols: (1) Phenolic compound stability after giving hydrogen atoms; (2) number of hydrogen atoms available in every phenolic compound; (3) easiness of interaction between the free radical and the phenolic; (4) how easily the free radicals from phenolic compound can combine with more active free radicals; and (5) that possibility of a new antioxidant can form after the phenolic compound provides hydrogen atoms. Besides, the chemical nature of the radical used as an experimental model and the unit of expression of the results could be also added. The radical scavenging antioxidants play an essential role in the maintenance of health and prevention of diseases; then, evaluate the antiradical capacity of antioxidants is critically important.

In addition to the previous determinations, antioxidant capacity was measured in the raw extract obtained using ORAC and ferric reducing ability of plasma (FRAP) assays, complemented with •OH radical and hemolysis inhibition tests (Table 2). As a set, the tests made a major approach to bioprospection of the seed in a living organism. The antioxidant capacities obtained in the present study were high compared to other Myrtaceae fruits. Zapata et al. [31] reported the antioxidant capacity of bitter guava (P. aracá), finding 6.6799 mmol Trolox/100 g, 10.3334 mmol Trolox/100 g, 0.6239 g AAE/100 g, and 0.6239 g DMSOE/100 g by ABTS, FRAP, ORAC, and •OH radical (terephthalic acid test), respectively. Thaipong et al. [65] reported the antioxidant activity measured in methanol extracts (AOAM) of guava fruits. The average AOAM values were 31.1, 25.2, 26.1, and 21.3 mM TE/g as determined by the ABTS, DPPH, FRAP, and ORAC assays, respectively. Therefore, comparing the previous results with those obtained in this work it can be affirmed that there are antioxidant compounds in arazá seeds, maybe polyphenols, which may be isolated and analyzed to determine chemical identity. These exhibit biological significant functions such as protection against oxidative stresses and degenerative diseases.

The delicate balance between the advantageous and detrimental effects of free radicals is clearly an important aspect of life. The science of biological redox regulation is a rapidly growing field of research that has an impact on diverse disciplines including physiology, cell biology, and clinical medicine. This seems to be the first report that shows a comparison of four techniques of such activity in arazá seeds.

Antihelminthic activity

The capacity of REE, DF, and HR to inhibit egg hatching, loss of cuticle, motility, and to induce mortality on GI nematodes and P. redivivus mortality induction are given in Table 3. The data revealed that the ethanolic extract showed marked degree antihelminthic activity than DF and HR in all tests on GI nematodes; in addition, it was more effective against free-life organism (P. redivivus) surpassing to DF and HR action, even though all the samples were endowed with the antihelminthic property.

Table 2: Phenolic content, antiradical and antioxidant activity of arazá seeds

<table>
<thead>
<tr>
<th>Assay</th>
<th>REE</th>
<th>DF</th>
<th>HR</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antiradical</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>capacity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DPH•</td>
<td>3.0±0.06</td>
<td>2.65±0.01</td>
<td>2.84±0.17</td>
</tr>
<tr>
<td>IC50</td>
<td>32.70±0.05</td>
<td>37.71±0.08</td>
<td>35.25±0.81</td>
</tr>
<tr>
<td>AP</td>
<td>65.91±15.19</td>
<td>759.76±17.19</td>
<td>710.73±43.84</td>
</tr>
<tr>
<td>TEAC</td>
<td>4.22±0.02</td>
<td>0.83±0.04</td>
<td></td>
</tr>
<tr>
<td>ABTS**</td>
<td>23.69±0.40</td>
<td>100.01±0.11</td>
<td>100.09±0.13</td>
</tr>
<tr>
<td>IC50</td>
<td>44.66±52.82</td>
<td>549.13±86.91</td>
<td>2275.40±110.20</td>
</tr>
<tr>
<td><strong>Antioxidant</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>capacity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HI</td>
<td>24.79±62.40</td>
<td>200.51±98.95</td>
<td>222.09±25.03</td>
</tr>
<tr>
<td>FRAP</td>
<td>4.05±0.36</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>ORAC</td>
<td>27.42±13.76</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>•OH</td>
<td>12.75±1.98</td>
<td>NR</td>
<td>NR</td>
</tr>
</tbody>
</table>

REE: Raw ethanolic extract, DF: Dichloromethane fraction, HR: Hydroalcoholic residue, IC50: 50% inhibitory concentration, mg/L. AP: Antioxidant potential (percent, %). TEAC: Trolox equivalent antioxidant capacity mmol Trolox/100 g. HI: Hemolysis inhibition (IC50). FRAP: Ferric reducing ability of plasma, g AAE/100 g (Ascorbic acid equivalent antioxidant capacity, µmol/100 g). ORAC: Oxygen radical absorbance capacity, TEAC, •OH radical inhibition (DMSOE: Dimethylsulfoxide equivalents, g DMSOE/100 g). NR: Non-reported
Table 3: Anthelmintic activity of extract and fractions

<table>
<thead>
<tr>
<th>Determinations</th>
<th>Samples</th>
<th>REE</th>
<th>DF</th>
<th>HR</th>
</tr>
</thead>
<tbody>
<tr>
<td>GI nematodes</td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>7.18±0.28</td>
<td>11.90±0.10</td>
<td>18.15±0.43</td>
</tr>
<tr>
<td>Cuticle loss inhibition</td>
<td></td>
<td>11.68±0.04</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Mortality</td>
<td>LC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>11.80±0.21</td>
<td>12.05±0.23</td>
<td>17.89±0.52</td>
</tr>
<tr>
<td>P. redivivus</td>
<td></td>
<td>11.26±0.23</td>
<td>ND</td>
<td>11.81±0.06</td>
</tr>
<tr>
<td>Mortality</td>
<td>LC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>2.26±0.31</td>
<td>3.61±0.04</td>
<td>6.46±0.11</td>
</tr>
</tbody>
</table>

REE: Raw ethanolic extract; DF: Dichloromethanic fraction; HR: Hydroalcoholic residue. IC<sub>50</sub>: Inhibitory concentration 50% (mg/ml); LC<sub>50</sub>: Lethal concentration 50% (%/mg/ml); ND: Non-detected under test conditions. P. redivivus: Panagrellus redivivus; GI: Gastrointestinal

Statistical analysis applied to these tests showed differences in the activity of the treatments between egg hatching inhibition, GI nematodes motility inhibition, and mortality. The ANOVA confirmed what had been stated before (p<0.05), was verified variance normality and homogeneity (p>0.01). Ayala et al. [66] reported a lower LC<sub>50</sub> (0.707 mg/ml) in aqueous extracts of leaves of Eucalyptus globus (Myrtaceae). However, these authors allowed a higher exposition time (96 h) to the one used with arazá seeds (24 h).

The anthelmintic activity of REE could be due to the constituents present, mainly the phenols of tannic nature. It is important to mention that condensed tannins have been recognized with the capability to inhibit female fertility and egg hatching in parasitic species T. colubriformis, T. circumcincta, and H. contortus [67,68]. The presence of such constituents, in synergy with other metabolites (hydrolyzable tannins, flavonoids, saponins, terpenes, among others) found in arazá seeds, would explain partially the higher potentiality of REE in antiparasitic tests.

Complementarily, GI nematodes collected from infected sheep in their L<sub>3</sub> stage were identified [35]. The organisms were categorized in three genres according to their main morphological features: Cooperia, as the most abundant (62.50±1.29%); Haemonchus, the second in abundance (24.00±2.30%); and Teladorsagia, the one with the lowest level (12.50±1.29%).

Helminth infections are among the chronic pervasive infection and a foremost degenerative disease distressing a large proportion of the world population [69] and in livestock [70,71]. Evaluation of activities of natural plant products claimed for possessing the anthelmintic property is getting the attention these days. Screening and proper evaluation of the claimed they could offer possible alternatives that may be both sustainable and environmentally acceptable. The results of this study suggested the possible use of E. stipitata seed ethanolic extract in intestinal nematode control.

Cytotoxicity

With the intention of knowing the possibility that derivative products of E. stipitata seeds can be ingested or applied in some way, this work tested cytotoxicity of the raw extract on mononuclear cells. A mononuclear CV of 96.99±4.56% was observed. Rodríguez-Feo et al [72] suggest that extracts that allow cell feasibility below 80% can be considered as toxic. Harmlessness of REE, DF, and HR was confirmed on human erythrocytes, the responses were below 3% which suggests the absence of hemotoxic metabolites in raw extract and fractions obtained from E. stipitata seeds.

CONCLUSIONS

The study revealed that E. stipitata seeds, commonly known as arazá, collected in the Colombia Andean region, can be used as a food supplement. Biological activity tests carried out lead to consider that many of those would be associated with the joint action of several metabolites distributes irregularly in the samples that were studied, mainly to the phenolic constituents. The presence of this kind of compounds, mostly tannic, enables the seed as a raw material in the elaboration of nutraceutical products with antiradical and/or antioxidant action. In any, its low toxicity confers high probabilities for the product to be ingested or applied in some way. This paper seems to be the first work carried out in Colombia interested in knowing physical, chemical, and biological features of arazá seeds; this information can be useful in food engineering, pharmacological industry (human or veterinarian), botanic, and agronomical. Thus, the use of this residue in arazá fruit chain should give added value to the fruit, reducing the possibilities that its bad provision contributes to environmental contamination.

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AUTHORS CONTRIBUTIONS

All the authors made substantial contributions to the conception, design, acquisition, analysis, and interpretation of data.

CONFLICT OF INTEREST

All authors have none to declare.

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