Original

Palm tree syrup; nutritional composition of a natural edulcorant

G. Luis1, C. Rubio1, A. J. Gutiérrez1, C. Hernández1, D. González-Weller2, C. Revert1, A. Castilla1, P. Abreu3 and A. Hardisson1


Abstract

Introduction: Palm syrup is a typical product from the Canary Islands, traditionally produced from the sap of the tropical palm tree Phoenix canariensis. Its high caloric content has led to its increasing use as a health food supplement for athletes, children and elderly. Furthermore, demand for this natural syrup is continuously increasing due also to its medicinal uses in homeopathic medicine.

Objective: Palm Tree syrup samples prepared with palm sap from primary producers in La Gomera island (Canary Islands, Spain) were analyzed for their nutritional composition (moisture, ash, sugars, fat, vitamins and minerals).

Methods: 35 syrup samples from five different producing regions in La Gomera island were analyzed. High-performance liquid chromatography (HPLC) was used to determine sugars and vitamins and Flame Atomic Absorption Spectrophotometry (FAAS) was used to analyze the minerals.

Results: Major carbohydrates were sucrose (37.8%), glucose (9.50%) and fructose (4.80%), respectively. The presence of arabinose could not be confirmed. Niacin was the water-soluble vitamin with the highest concentration with an average content of 0.003%. Fat content was found to be under 0.20%. Potassium was the mineral with highest contents (0.45%).

Conclusions: Results suggest that palm tree syrup can play an important role as a sugar and mineral source in human nutrition, suggesting that future applications for this product could be developed.

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Correspondence: Gara Luis.
Department of Toxicology. University of la Laguna.
E-mail: garaluisglez@gmail.com

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Abbreviations

FAAS: Flame Atomic Absorption Spectrometry.
HPLC: High-performance liquid chromatography.
HNO₃: Nitric acid.
H₂SO₄: Sulphuric acid.
b.p.: Bring to boiling.
UV detection: Ultraviolet detection.
NIST: National Institute of Standard and Technology.

Introduction

Palm syrup is a typical product from the Canary Islands, described as having been prepared by the former inhabitants of the islands before the arrival of the Spaniards. Palm syrup has been traditionally produced from the sap of the tropical palm tree Phoenix canariensis. For that reason, the palm tree sap, known as “guarapo”, is collected from the palm trees in the early spring, when temperatures fluctuate between -5 and -10 ºC. Then it must be boiled in a large pot on a wood fired stove until it becomes concentrated. The unique flavour of this natural product develops during this evaporation process (93-110 ºC for 1.5 h). The flow-sheet to obtain the syrup is shown in figure 1, beginning with the extraction of palm tree sap. The resulting syrup is a viscous, sweet liquid, dark with reddish tones. Although this natural product has a similar density to honey, it is less sticky. The composition of the final product depends on the botanical origin, time and temperature of storage. Authors have reported physicochemical changes and degradation of the syrup due to overheating. For that reason, is important to ensure the stability of the syrup to protect against fermentation and crystallization.

Although palm syrup is produced only in limited quantities, this natural product has a high commercial value because its unique flavour has made it popular in the confectionery industry and among consumers. Palm tree syrup is used by the Canarian population to flavour cakes, desserts, food coating or mixed drinks. For that reason, its high caloric content has led to its increasing use as a health food supplement for athletes, children and elderly. Furthermore, the demand for this natural syrup is continuously increasing due also to its medicinal uses in homeopathic medicine.

Palm tree syrup does not contain any kind of additive or artificial colourings and can last for years under the right storage conditions. Very few studies about the chemical and nutrient composition of palm tree syrup have been carried out. Nevertheless, producers and consumers are demanding more and more data about its composition to protect against adulteration via the addition of inexpensive sweeteners, and also for nutritional reasons. As the nutrient composition of Palm Tree syrup varies significantly depending on its geographical origin, it is possible to detect its adulteration and to ensure product purity.

The aim of this study was to determine the nutrient composition of palm tree syrup.

Materials and methods

Samples collection

A total of 35 samples of syrup were studied. Samples were collected during a sixteen month production period, between March 2007 and July 2008, and were analyzed. All samples were provided by producers from five different regions (Alojera, Taguluche, Tazo, Valle Gran Rey, Vallehermoso) of La Gomera island (Canary Islands, Spain). Random samples were collected in a manner proportional to the quantities obtained in each producing area. The sampling carried out in all zones was homogeneous. All samples were stored at -18 ºC before being analyzed. Approximately 100 g of each sample was divided into three sections: (1) 20 g were used to determine sugar and fat contents, (2) 50 g to determine metal content and (3) 30 g to determine vitamin content. All measurements were made in triplicate.

Nutrient analysis

Moisture content was determined by weighing the samples before and after being dried for 48 h at 85 ºC, and ash content was determined by heating the samples for 48 h at 450 ºC, according to the methods of the Association of Official Analytical Chemists.

High-Performance Liquid Chromatography (HPLC) techniques are often employed for the analysis of carbohydrates. Sugars analysis was carried out by means of HPLC employing a Shimadzu chromatograph equipped with a RID-6A detector using an Aminex HPX-87C column (250 x 4 mm). The columns were first conditioned by passing 2 mL of methanol through them and then
washed with 6 mL of watermill-Q. Columns were used with acetonitrile/water mixtures (80:20) as an eluent relying on the hydrophilic interaction of the sugars with a water layer solvating the column. In this case, glucose, fructose, and sucrose concentrations were determined by injecting 20 µL of the standard solutions or sample extracts and eluting them with ultra pure water at flow rate of 2 mL/min. For that reason, refractive index detection has been used with its attendant drawbacks of limited sensitivity and instability with regard to fluctuations in mobile phase composition and eluent temperature.

Carbohydrates were quantified by comparison to appropriate standards.

The extracted fat was determined by a gravimetric procedure. 1 g of palm syrup was accurately weighed in a fat extraction flask. Water was added to bring the volume to 10 mL and mixed. 10 mL of hydrochloric acid (HCl) were added slowly and the content was again well mixed. The fat extraction flask was set in a heated water bath at 70-80 ºC, brought to a boil and heated again at boiling point for 30 min while the flask was shaken carefully every 5 min. The content was cooled to room temperature. 25 mL of ether was added to the flask and then it was shaken vigorously for 1 min. 25 mL of light petroleum ether was added and shaken again. The ether and light petroleum steps were each repeated twice more. Finally, the solvents were evaporated and the extracted fat dried to constant weight in an oven at 103 ± 2 ºC for 1 h, cooled to room temperature in a desiccator and weighed.

Most vitamins were analyzed by HPLC, vitamin B₁ (thiamine) was extracted using 0.1 N sulphuric acid. Quantification was done with reversed-phase HPLC and post-column derivatization with fluorescence detection. Riboflavin (vitamin B₂) and its coenzyme forms were extracted with 0.1 N sulphuric acid in an autoclave at 120 ºC. After neutralization, the enzyme treatment removed the phosphate groups. The riboflavin content in the extract was determined by reversed-phase HPLC and fluorescence detection. Vitamin B₉ (nicotin acid) was determined by reversed-phase HPLC and UV detection after extraction with 0.001 N sulphuric acid.

Vitamin B₉ (pidade) was extracted using sodium acetate buffer and analyzed using reversed-phase HPLC with fluorescence detection. Vitamins B₆ and B₉ were analyzed after aqueous extraction by means of microbiological assays. Vitamin D₃ (cholecalciferol) was extracted with hexane after alkaline saponification. D₃ was used as internal standard. The crude extract was cleaned in a semi-preparative HPLC. The fraction containing D₂ and D₃ was quantified by HPLC with UV detection.

Vitamin E (α-tocopherol) was analyzed after alkaline saponification and extraction with hexane by HPLC and fluorescence detection. Vitamin K was determined following enzymatic digestion, extraction and subsequent HPLC with post-column reduction with zinc and fluorescence detection.¹⁵

Mineral contents were determined using a flame atomic absorption spectrophotometer (FAAS) model Varian Spectr AA-10 Plus equipped with a deuterium lamp for background correction and hollow-cathode lamps for each of the studied elements. Quantification was performed using external standards (Merck IV, multielement standard solution) and all the standard curves were obtained at 5 different concentrations. Table I

<table>
<thead>
<tr>
<th>Trace element</th>
<th>Wavelength (nm)</th>
<th>Spectral band (nm)</th>
<th>Air acetylene flow (mg/L)</th>
<th>Sensibility (mg/L)</th>
<th>Check sensitivity</th>
<th>Linear range (mg/L)</th>
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</thead>
<tbody>
<tr>
<td><strong>Microelements</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Cu</td>
<td>324.8</td>
<td>0.7</td>
<td>8.25</td>
<td>0.077</td>
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<td>5.0</td>
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<tr>
<td>Fe</td>
<td>248.3</td>
<td>0.2</td>
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<td>0.100</td>
<td>5.0</td>
<td>5.0</td>
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<tr>
<td>Mn</td>
<td>279.5</td>
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<td>0.052</td>
<td>2.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Zn</td>
<td>213.9</td>
<td>0.7</td>
<td>8.25</td>
<td>0.018</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Ni</td>
<td>232.0</td>
<td>0.2</td>
<td>8.25</td>
<td>0.140</td>
<td>7.0</td>
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</tr>
<tr>
<td><strong>Macroelements</strong></td>
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<td></td>
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<td></td>
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<tr>
<td>Ca</td>
<td>422.7</td>
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<td>0.092</td>
<td>4.0</td>
<td>5.0</td>
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<td>Mg</td>
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<td>8.25</td>
<td>0.0078</td>
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<tr>
<td>Na</td>
<td>589.0</td>
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<td>8.25</td>
<td>0.012</td>
<td>0.5</td>
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<tr>
<td>K</td>
<td>766.5</td>
<td>1.4</td>
<td>8.25</td>
<td>0.043</td>
<td>2.0</td>
<td>2.0</td>
</tr>
</tbody>
</table>
shows the instrumental parameters for the determination of microelements and macroelements by FAAS.

Quality control

Quality control of the analytical measurements was performed using blank samples and the following reference material: SRM-1515 (Apple leaves) from the National Institute of Standard and Technology (NIST). The recoveries obtained with the reference material were all above 98%.

All glassware was washed with acid and rinsed with purified water.

Results and discussion

Table II shows the results of the moisture, ash, sugars and fat analysis.

Moisture content in this natural product was at 35.3% and the total solid at 64.7%. Water content can vary depending on the environmental conditions of the island.20-21

Ash content in palm syrup was 1.78%. Some studies have found that the ash content for honey in the province of Santa Cruz of Tenerife is 0.31%.22

All palm tree syrup samples contained significant amounts of sugars (66.0%). Sugars analysis indicated that all the palm tree syrup samples were rich in monosaccharides and disaccharides and that the sugars identified in the palm tree syrup samples with semi-quantitative values were sucrose, glucose and fructose. The levels of the three main sugars are quite similar in all samples despite geographical differences.23-24 The major carbohydrate found in the analyzed palm tree syrup samples was sucrose (37.8%) followed by glucose (9.50%) and fructose (4.80%), respectively.

Results published by other authors show that levels of sucrose (in the 60-83% range) are higher that the mean values found in this study and that levels of glucose (in the 3.15-5.04% range) are lower that our mean value (9.50%).1

Table III shows the results of the water-soluble and fat-soluble vitamins in palm tree syrup samples. In general, the total vitamin content was low, below the level of nutritional importance. The water-soluble vitamin content is higher than that of fat-soluble vitamins. Regarding the fat-soluble vitamins, the palm tree syrup samples and the palm tree sap samples showed similar levels. Concentrations are arranged in the following sequence: A > E > D3 > K. Vitamins D (cholecalciferol) and K were rarely detectable, with concentrations less that 5.00 µg/kg and 1.00 µg/kg, respectively.

In all palm tree syrup samples, the water-soluble vitamin niacin showed the highest concentration (31.7 mg/kg) followed by ascorbic acid (5.00 mg/kg), cyanobalamin (5.00 mg/kg), pyridoxine (0.40 mg/kg), thiamine (0.20 mg/kg) riboflavin (0.10 mg/kg) and folic acid (0.10 mg/kg).
Conclusions

Palm tree syrup is a supersaturated solution of sugars and water, suggesting that this natural product could be a suitable raw material in the production of natural sweeteners. The vitamin content of palm tree syrup is higher than that of other syrups of botanical origin. Although the fat, vitamin and mineral contents of palm tree syrup are low, this natural product contains a considerable amount of potassium and sodium. Overall, the results of this study suggest that palm tree syrup can play an important role as a mineral source in human nutrition.

Further studies should investigate new applications for this syrup, like its use as a sweetener.

References