Influence of Genotype and Harvest Season on the Phytochemical Composition of Araçá (Psidium cattleianum Sabine) Fruit

Andrea Miranda Teixeira, Fabio Clasen Chaves, Rodrigo Cezar Franzon, Cesar Valmor Rombaldi

1Uergs, Unidade Cachoeira do Sul, CEP 96508-010, Cachoeira do Sul, RS, Brazil
2UFpel/FAEM, Department of Agroindustrial Science and Technology, C.P. 354, CEP 90010-900, Pelotas, RS, Brazil
3Embrapa Clima Temperado, BR 392, Km 78, C.P. 403, 96010-971, Pelotas, RS, Brazil

Abstract

L-ascorbic acid, total anthocyanin, total carotenoid, total tannin, total and individual phenolic compounds content, antioxidant potential, and antiproliferative activity against tumoral cells of extracts from six Araçá genotypes (AR9, AR19, AR29, AR27, AR46 and AR72) from six consecutive harvest seasons (2008, 2009, 2010, 2011, 2012, and 2013) were evaluated. Results showed that in general variations in phytochemical content were greater among genotypes than among harvests. Whereas red genotypes (AR9, AR19, and AR29) had higher anthocyanin and total phenolic content, yellow genotypes (AR46, and AR72) had higher L-ascorbic acid content. (-)-Epicatechin was the most abundant phenolic compound found in the fruits, followed by gallic acid, in all genotypes. Only soluble solids, pH, and total tannin content were not affected by harvest year. Regarding the antioxidant capacity against S. cerevisae XV 185-14C cells, all extracts revealed a protective effect from the oxidative stress agent (H2O2). Similarly, all extracts inhibited the proliferation of the human colon adenocarcinoma cells (Caco-2) without cytotoxicity towards fibroblast cells (3T3). Although harvest season influenced the functional potential of Araçá fruit, genotype had the strongest influence.

Keywords: Phytochemicals; Antioxidant capacity; Antiproliferative activity; Genotypes

Introduction

The majority of studies regarding human nutrition and health demonstrate that the occurrence of chronic degenerative diseases is negatively correlated to the dietary intake of grains, fruits, and vegetables[1-6]. Research data are corroborated by adhoc studies, which relate cause and effect in heterologous models (in vitro and in vivo chemical simulations) and by cohort studies[6-11]. On the other hand, although less common, there are studies that show the absence of such a correlation[12].

Health benefits which are attributed to food rich in phenolics compounds and natural antioxidants have increasingly enhanced the search for new species that comprise relevant biological activities, such as protection against diseases. In this regard, Brazil has the advantage of having the planet’s largest biodiversity and variability. However, many fruit species are still either unknown or underutilized in some cases due to difficulties in cultivation. The greatest richness in native Brazilian fruit species may be found in the Amazon region or in the savannahs. However, there are many fruit species in temperate climates with high potentialities[9,13-17]. Araçá (Psidium cattleianum Sabine) fruit, belonging to the Myrtaceae family, produces small sized yellowish or reddish fruits with attractive taste and aroma[9]. Although there are a few studies on the Araçá physicochemical composition, and its antioxidant and antimicrobial properties and on the development of products from its fruit[18-20], the characterization of its phytochemicals in suc-
Materials and Methods

Plant material

Red (AR9, AR19, and AR29) and yellow (AR27, AR46, and AR72) Araçá fruit ( *P. cattleianum* Sabine) were harvested from the Embrapa Clima Temperado germplasm bank, Pelotas, RS, Brazil. One kilogram of fruit from each plant (three plants from each genotype) was collected when ripe (> 50% fruit surface red or yellow in color, accordingly) during the harvest seasons of 2008, 2009, 2010, 2011, 2012, and 2013. Fruit were frozen in liquid nitrogen and stored at -80°C until further analysis. All analyses were performed in triplicate.

Physicochemical composition

Soluble solids content and pH: Soluble solids content (SSC) was determined using a refractometer directly on the juice squeezed from the fruit flesh. Results were expressed in °Brix. pH was measured directly on the fruit juice using a digital pH-meter.

L-ascorbic acid content: L-ascorbic acid content was determined by high performance liquid chromatography (HPLC) using a reverse phase column CLC-ODS (4.6 mm x 150 mm x 5 µm), with a UV detector set at 254 nm[9]. Mean recovery index (RI) was 92.25% and results were expressed in milligrams of L-ascorbic acid per 100 g of fresh fruit.

Total anthocyanin content: Total anthocyanin content was determined spectrophotometrically, following the method described by[23]. One g of ground fruit were extracted with 25 mL acidified ethanol (HCl, pH 1.0) and the solution kept at rest for 1 h, in the dark, and homogenized every 5 min. The mixture was then filtered and the volume completed to 50 mL with acidified ethanol. Absorbance was measured on a spectrophotometer (UV/Vis Ultrospec 2000 Pharmacia Biotech) at 520 nm. Results were expressed in milligrams of cyanidin-3-glucoside per 100 g of fresh fruit.

Total carotenoid content: Total carotenoid content was determined spectrophotometrically, following the method described by[24]. Five grams of fruit were ground with celite, to which were added 20 mL of cold acetone (4°C) for the extraction of pigments and homogenized for approximately 10 min. The sample was vacuum-filtered and the residue was extracted two more times. The filtrate was transferred to a separatory funnel and hexane and distilled water were added. The lower phase was discarded and then washed with distilled water until the total removal of acetone. Absorbance of the hexane extract was measured in a spectrophotometer (UV/Vis Ultrospec 2000 Pharmacia Biotech) at 450 nm. Results were expressed in microgram of β-carotene per gram of fresh fruit.

Total tannin content: Total tannin content was determined spectrophotometrically, according to method described by[25]. Five g of the sample were extracted with methanol (10 mL) under stirring for 1 h at room temperature. The extract was filtered and 100 µL were reacted with the Folin-Denis reagent for 3 minutes when a saturated sodium carbonate solution was added. The solution was kept in the dark at room temperature for 1 h. Absorbance was measured in a spectrophotometer (UV/Vis Ultrospec 2000 Pharmacia Biotech) at 760 nm. Results were expressed in milligrams of tannin per 100 g of fresh fruit.

Total phenolic content: Total phenolic content was determined by spectrophotometry using the Folin-Ciocalteau reagent[9]. Absorbance was measured on a spectrophotometer (UV/Vis Ultrospec 2000 Pharmacia Biotech) at 760 nm. Results were expressed in milligrams of gallic acid equivalent (GAE) per 100 g of fresh fruit.

Individual phenolic compounds contents

Individual phenolic compounds contents were determined by high performance liquid chromatography (HPLC, Shimadzu)[9]. The phenolic compounds were extracted and hydrolyzed with 35 mL acidified methanol (HCl, 15% v/v). The extract was homogenized at 35°C, in the dark, for 24 h, filtered through cotton wool and the supernatant concentrated in a rotary evaporator at 40°C for 20 minutes. The concentrated residue was re-dissolved in methanol (10 mL) and centrifuged at 7000 x g for 10 min. A 30 µL aliquot from the supernatant were extracted in the HPLC-UV with the detector set at 254 nm[9], with a UV detector set at 254 nm[9]. Mean recovery index (RI) was 87.56%, p-coumaric acid (IR 92.34%), ferulic acid (IR 103.87%), quercetin (IR 93.25%) and (-)-epicatechin (IR 90.08%). Results were expressed in milligrams of the respective phenolic compound per 100 grams of fresh fruit.
Antioxidant capacity was determined using the *in vivo* system of eukaryotic *Saccharomyces* XV 185-14C yeasts cell assay [26]. The yeast strain was inoculated in yeast peptone dextrose (YPD) solid medium with 1% (m/v) yeast extract, 2% (m/v) peptone, 2% (m/v) glucose and 2% (m/v) agar. Cells were transferred to a liquid medium (with the same composition of solid media without agar) and placed in an orbital stirrer at 28°C and at 160 rpm. Cell suspensions with 2 x 10⁶ CFU mL⁻¹ were treated with fruit extracts diluted in water at 1:4 (extract:water) and incubated for 1 h at 28°C while stirring in the dark. Cells were centrifuged at 2000 x g, 28°C for 5 min and washed twice with 0.9% (m/v) sodium chloride solution. Cells were then treated with a solution of hydrogen peroxide (50 mM), for 1 h, at 28°C. Samples were diluted in a solution of sodium chloride (0.9% m/v), seeded in a complete YPD medium and incubated at 28°C for 48 h. After incubation, the colonies were counted and survival percentages determined. Test plates were taken as reference, in which the total number of colonies in the control plate (without treatment) was considered as 100% cell survival.

**Antiproliferative activity against tumoral cells**

The antiproliferative effect of fruit extracts was tested in human colon adenocarcinoma cells (Caco-2), whereas a strain of rat’s embryo fibroblasts (3T3) was used as control. RPMI 1640 medium, with bovine fetal serum (20% v/v) and antibiotics (100 UmL⁻¹ penicillin G, 100 µgmL⁻¹ streptomycin) was used for cell culture (5 x 10⁵ mLcells⁻¹). Cells were placed in 96-well micro culture plates, with 100 µL per well, for 24 h, at 37°C in an atmosphere with 95% O₂, 5% CO₂ and 100% relative humidity. After incubation, medium was removed from each well, with cells kept in the lower section. Cells were exposed to a new medium with 80 µgmL⁻¹ extract. After 48-h incubation, cells were fixed by adding trichloroacetic acid (50% m/v) at 4°C for 1h. The colorimetric assay was performed by adding a solution of sulforhodamine B (0.4% m/v) in acidified water (acetic acid 1% v/v) in each well. After 30 min at room temperature, the non-fixed solution was removed by washing with acidified water. Stain fixed to cell proteins was re-solubilized with Tris 10 mM (pH10.0) buffer, on an orbital stirrer at 50 rpm, at room temperature, for 5 min. Optical density was read by ELISA plate spectrophotometric reader at 540 nm. Absorbance data were correlated with a standard curve of viable cells and results were given in cell survival percentage compared to control treatment with cells cultivated in RPMI 1640 medium.

**Experimental design and statistical analysis**

This study was a descriptive analysis of 6 genotypes (3 plants per genotype) during 6 harvest seasons (2008, 2009, 2010, 2011, 2012, and 2013), with three fruit harvests per plant in each year. Analytical replicates were performed in triplicate. Results were evaluated by ANOVA and averages compared by Tukey’s test at 5% probability using Statistica v.7.1 (Statsoft, 2005).

**Results and Discussion**

**Effect of genotype and harvest season on the phytochemical composition of Araca fruit**

The factor genotype affected all the evaluated dependent variables (soluble solids, pH, L-ascorbic acid, anthocyanin, carotenoid, tannin, and phenolic contents) within the set of independent variables (6 genotypes and 6 harvest seasons) under analysis (Table 1). Three of these genotypes produce red fruits (AR9, AR19, and AR29) and 3 produce yellow ones (AR27, AR46, and AR72). In general, both groups differed in pH, anthocyanin, and total phenolic content.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>L-ascorbic acid</th>
<th>Total anthocyanins</th>
<th>Total carotenoids</th>
<th>Total tannins</th>
<th>Total phenolics compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>**</td>
<td>**</td>
<td>**</td>
<td>**</td>
<td>**</td>
</tr>
<tr>
<td>Harvest season</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>ns</td>
<td>**</td>
</tr>
</tbody>
</table>

ns – not significant
* Significantly different only after some years
** Significantly different

Soluble solids content and pH varied only among genotypes and not with harvest season (Table 2). AR72 with 7.17° Brix had the lowest soluble solids content and AR19 with 11.91°Brix the highest. AR29 fruit had the lowest pH (3.08) while AR46 had the highest (3.88). These results characterize Araca as a fruit with relatively low SS and pH.
Table 2: Soluble solids (SS) and pH of red (AR9, AR19, and AR29) and yellow (AR27, AR46, and AR72) Araçá (mean rates correspond to 2008 – 2013 harvest seasons)

<table>
<thead>
<tr>
<th>Genotype</th>
<th>SS* (n = 6)</th>
<th>pH (n = 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR 9</td>
<td>10.04 ± 0.56b</td>
<td>3.19 ± 0.03d</td>
</tr>
<tr>
<td>AR 19</td>
<td>11.91 ± 0.66a</td>
<td>3.34 ± 0.02c</td>
</tr>
<tr>
<td>AR 29</td>
<td>10.67 ± 0.99ab</td>
<td>3.08 ± 0.04e</td>
</tr>
<tr>
<td>AR 27</td>
<td>10.78 ± 1.01ab</td>
<td>3.72 ± 0.02b</td>
</tr>
<tr>
<td>AR 46</td>
<td>11.73 ± 1.16a</td>
<td>3.88 ± 0.07*</td>
</tr>
<tr>
<td>AR 72</td>
<td>7.17 ± 0.75c</td>
<td>3.83 ± 0.05a</td>
</tr>
</tbody>
</table>

Means followed by the same letter in the column are not significantly different by Tukey’s test (p < 0.05). *Soluble solids expressed as °Brix

Genotype and harvest season influenced the L-ascorbic acid content in Araçá fruits (Figure 1A). Genotype AR46 was the one with the highest L-ascorbic acid content in all harvest seasons reaching 11 mg100g^{-1} fruit in 2011 and 2012. L-ascorbic acid content revealed Araçá to be poor in this compound when compared to other fruits such as the acerola (1357 mg 100 g^{-1}) and camu-camu (1882 mg 100 g^{-1}), native fruits to the Amazon biome[27] or to other native species of southern Brazil, such as the guabiroba[28] or butia[29] with approximately 3050 and 9351 mg of L-ascorbic acid per 100 g of fresh fruit, respectively[30] revealing a wide variation for ascorbic acid content among genotypes and species. L-ascorbic acid content may also be affected by climate, such as solar radiation, temperature, and soil conditions[27,28,29,30]. The present study partially corroborated these findings since L-ascorbic acid content varied among genotypes and among harvest seasons.

There was a sharp difference in anthocyanin content between red and yellow fruits (Figure 1B). This fact was expected since these compounds are responsible for the red color of the fruit. Anthocyanin content was significantly higher in red genotypes in all harvests, with maximum of 6.33 mg cyanidin-3-glucoside 100 g^{-1} fruit for the 2012 harvest for genotype AR19, whereas the lowest anthocyanin content was 1.33 mg cyanidin-3-glucoside 100 g^{-1} fruit in yellow genotype AR27. Araçá anthocyanin content was relatively low when compared to fruit known for their high anthocyanin content, such as blackberry (47.7 mg cyanidin-3-glucoside 100 g^{-1} fruit), blueberry (128 mg cyanidin-3-glucoside 100 g^{-1} fruit)[31] and strawberry (23.4 mg cyanidin-3-glucoside100 g^{-1} fruit)[32]. Whereas anthocyanins are present throughout blackberry, blueberry, and strawberry fruit, these pigments occur only in the thin layer of fruit peel in Araçá.

Analysis of total carotenoid content in Araçá fruit showed that it was mainly influenced by genotype (Figure 1C). Only genotype AR46 was affected by the harvest season. Although Araçá genotypes had significantly different total carotenoid contents in each harvest they ranged from 4.33 to 10.0 µg β-carotene.g^{-1} (AR9), without a clear distinction between red and yellow fruits. A previous report also demonstrated carotenoid content to be higher in red fruits when compared to yellow ones[33] higher carotenoid rates in red fruits when compared to those of yellow ones. Among the fruits analyzed by[29], Araçá proved to have the highest carotenoid content, 6.270 µg β-carotene.g^{-1} fruit, significantly higher than the orange Brazilian cherry (4.02 µg β-carotene.g^{-1}) and butia (3.85 µg β-carotene g^{-1}) with orange-colored pulp. Data shown in this report confirm the literature, which reports that carotenoid
content may be influenced by several factors, such as variety, plant tissue, ripening stage, harvest conditions, geographic location and others[34,35].

Total tannin content was influenced only by genotype (Figure 1D). Red-colored fruits AR19 and AR29 had total tannin content significantly higher than those of the other genotypes, in each harvest, with maximum content reaching 398.5 mg 100 g⁻¹ fruit (2009) and 412.5 mg 100 g⁻¹ fruit (2013), respectively. There is no other report in the literature on tannin content of Araçá fruits. This group of phenolic compounds contributes towards the fruit’s astringent characteristic and suggests that the taste of the red-colored fruits is sharper.

Araçá phenolic content varied significantly among genotypes, in the same harvest, with a clear difference between red and yellow fruits ranging from 414.7 to 689.67 mg GAE.100 g⁻¹ (Figure 2A). Each year, red-colored genotypes had higher total phenolic content than the yellow ones, varying between 606.67 (AR9 in 2008) and 689.67 mg GAE.100g⁻¹ fruit (AR29 in 2012). The same genotypes also provided the highest anthocyanin and total tannin content (over 283.5 mg.100g⁻¹ fruit) when compared to yellow fruits. Phenolic content found for red (*Psidium cattleianum*) and yellow (*Psidium cattleianum var. lucidum*) Araçá reported by Biegelmeyer et al. (2011) was in the same range to that found in this study (501.33 mg GAE.100g⁻¹).

Figure 2: Total phenolic(A), (-)-epicatechin (B), gallic acid (C), p-coumaric acid (D), ferulic acid (E), and quercetin (F) content of red (AR9, AR19, and AR29) and yellow (AR27, AR46, and AR72) Araçá genotypes from six consecutive harvest season (2008, 2009, 2010, 2011, 2012, and 2013).

Among the individual phenolic compounds analyzed in the fruit (-)-epicatechin content ranging from 66 to 269 mg.100g⁻¹ and gallic acid from 52 to 147 mg 100g⁻¹ were the predominant phenolics (Figure 2). Variability existed in the composition and in the quantity of phenolic compounds among Araçá genotypes and harvest seasons[28] investigated the profile of phenolic compounds of native fruits and identified molecules derived from gallic acid and quercetin as major components of the Araçá. However, they did not identify the specific phenolic compound measured.

**Effect of genotype and harvest season on the in vivo antioxidant capacity and the anti-proliferative activity of Araçá fruit extracts**

The extract’s *in vivo* antioxidant capacity was evaluated by comparing survival levels of the yeast *S. cerevisiae* against damages caused by the induction of oxidative stress by hydrogen peroxide (negative control) and Araçá extracts and this stress agent. Expectedly, yeast cells revealed sensitiveness to hydrogen peroxide and survival ranged between 42% and 43% in the presence of the oxidative agent. Results corroborated the 46% survival of yeasts treated with H₂O₂ reported by[11]. The protection ability of Araçá fruit extracts was observed when *S. cerevisiae* cells were treated with Araçá fruit extracts and the oxidative stress agent.
Table 3: Antioxidant capacity of red (AR9, AR19, and AR29) and yellow (AR27, AR46, and AR72) araçá extracts towards S. cerevisaeae XV 185-14C

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Survival (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive control (water)</td>
<td>100.00a</td>
</tr>
<tr>
<td>Negative control (H₂O₂ 50 mM)</td>
<td>42.60b</td>
</tr>
<tr>
<td>AR9 + H₂O₂ 50 mM</td>
<td>85.83b</td>
</tr>
<tr>
<td>AR19 + H₂O₂ 50 mM</td>
<td>86.00b</td>
</tr>
<tr>
<td>AR29 + H₂O₂ 50 mM</td>
<td>82.33bc</td>
</tr>
<tr>
<td>AR27 + H₂O₂ 50 mM</td>
<td>79.67c</td>
</tr>
<tr>
<td>AR46 + H₂O₂ 50 mM</td>
<td>80.83bc</td>
</tr>
<tr>
<td>AR72 + H₂O₂ 50 mM</td>
<td>81.50bc</td>
</tr>
</tbody>
</table>

Means followed by the same letter in the column are not significantly different by Tukey’s test (p < 0.05).

The extracts’ antioxidant capacity increased significantly the survival rates of yeast treated with H₂O₂. Red-color genotypes increased survival rate between 40 and 46% when compared to yeast cells exposed to H₂O₂. The same genotypes were found rich in anthocyanin, tannin, and total phenolic contents and thus corroborated reports by authors that assert the capacity of these molecules in protecting cells against damages caused by different oxidative stress agents[11,26,36].

Since low cell death (between 2 and 9%) occurred in control fibroblast cells (3T3), extracts revealed low cytotoxicity. Regardless of the harvest season, the inhibition of cell proliferation in all extracts was reported when the effect of extracts of araçá fruits was evaluated for the proliferation of human colon adenocarcinoma cells (Caco-2). Tumoral Caco-2 cells treated with extracts of yellow-color genotypes had, in general, a higher survival rate (between 45 and 59%) throughout the years (Table 4), that means these extracts have a lower antiproliferative activity when compared to red-color genotypes. The biochemical-molecular mechanism, by which extracts reduced the proliferation of CaCo-2 cells without any cytotoxic activity with cells 3T3, was not studied in current assay.

Table 4: Antiproliferative activity of red (AR9, AR19, and AR29) and yellow (AR27, AR46, and AR72) araçá extracts tested against embryonic fibroblasts (3T3) and human colon adenocarcinoma (Caco-2) cells

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Survival (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3T3</td>
</tr>
<tr>
<td>Control</td>
<td>98.00 a</td>
</tr>
<tr>
<td>AR9</td>
<td>96.50 ab</td>
</tr>
<tr>
<td>AR19</td>
<td>95.67 ab</td>
</tr>
<tr>
<td>AR29</td>
<td>93.50 bc</td>
</tr>
<tr>
<td>AR27</td>
<td>95.00 abc</td>
</tr>
<tr>
<td>AR46</td>
<td>92.33 c</td>
</tr>
<tr>
<td>AR72</td>
<td>94.67 bc</td>
</tr>
</tbody>
</table>

Means followed by the same letter in the column are not significantly different by Tukey’s test (p < 0.05).

Conclusion

Results showed that red and yellow araçá genotypes have great phytochemical variability. araçá fruit phytochemical content also varied with the harvest seasons, probably due to climatic variations. However the influence of genotype was stronger than harvest season on fruit phytochemical content and functional potential. Since fruit extracts increased the survival indexes of S. cerevisaeae cells, they revealed their protection capacity against oxidative stress agents. araçá fruit extracts also inhibited the proliferation of human colon adenocarcinoma cells. Given these results these genotypes show potential to be commercially explored and the consumption of the fruit may be promote in face to the significant functional potential.

Acknowledgments: The authors would like to thank Fapergs and CNPq for providing financial support for research.

References

Harvest Season on the Phytochemical Composition of Araca Fruit


