Grouping of anthurium genotypes based on genetic and morpho-anatomical features

Gabrielen de Maria Gomes Dias1*, Joyce Dória Rodrigues Soares2, Ana Cecília Ribeiro Castro3, Moacir Pasqual2, Ivan Júnior Ascari2 and Adriano Carvalho Costa4

Abstract: The study aimed at investigating the anatomic and DNA content characteristics of 12 Anthurium genotypes. A large genomic amplitude was observed in the genotypes, ranging from 4.95 to 22.74 pg of DNA. Differences in DNA content, assessed by flow cytometry, allowed the differentiation between the genotypes A. andraeanum, A. affine, A. bomplandii, and A. gracile and the hybrid (A. lindmanianum × A. guianesis) and clustered the group A. raimundii, plowmanii and jemanii of IFT 258, in all analyses. Anatomical studies clustered the genotypes A. andraeanum (Astral and Rubi) and distinguished the hybrid and A. bomplandii from the other genotypes, with anatomical features specific to each genotypes. Cluster analysis allowed grouping of the genotypes A. raimundii and A. affine, bomplandii and gracile, A. jemanii and plowmanii, and the cultivars Astral and Rubi, and isolated the hybrid. Demonstrated the importance of grouping in the diagnosis of species belonging to the genus Anthurium.

Key words: Araceae, leaf anatomy, characterization, DNA.

INTRODUCTION

The genus Anthurium is considered taxonomically complex, with many species not very well known, and only a few studies on their geographical distribution to-date. Taxonomic differentiation of the species is difficult, mainly due to their morphological plasticity (Coelho and Catharino 2008). Some are very similar, with subtle morphological and vegetative differences, which leads to taxonomic confusion within the group. Tillich (2003) points out that there are many gaps related to the structure of the seedling in Araceae, and important details about the morphology are still unclear even though 36 genera of the family (including Anthurium) have been studied. Publications on the Araceae family include an extensive work on leaf anatomy, but few anatomical analyses have been performed on the genus Anthurium. The most recent anatomical studies on Anthurium are based on the anatomy of the leaf and spathe (Mantovani and Pereira 2005, Mantovani et al. 2010) and leaf anatomy of young and adult plants in an epiphytic habitat (Lorenzo et al. 2010).

In plant breeding programs, information about the genetic diversity within a population is essential for the effective use of genetic resources. There are several ways of measuring data for the characterization of a germplasm bank. Therefore, data must be collected and interpreted correctly, since, according to Guedes (2002), statistical control of the process is important to avoid multiple-
Flow cytometry has been an attractive alternative to traditional methods of DNA content determination due to its ease, speed and accuracy (Dolezel and Bartos 2005). Flow cytometric analysis of nuclear DNA content is an excellent alternative to traditional methods of chromosome counting (Loureiro and Santos 2004) and can be used to differentiate species not yet identified, as well as for anatomical analysis. Anatomical features of leaves have been studied to resolve taxonomic confusion (Moraes et al. 2011). Therefore, combining leaf anatomy and flow cytometry can provide multiple features that help to clarify relationships within the species *Anthurium*.

Considering the large number of species and their diversity and complexity, it is important to use more advanced technologies, such as anatomy and flow cytometry as previously mentioned, to increase knowledge regarding genus *Anthurium*.

Therefore, the objective of this study was to compare morpho-anatomical characteristics for taxonomic studies of the genus *Anthurium*, by clustering of genotypes through cluster analysis and quantification of DNA content.

**MATERIAL AND METHODS**

The experiment was conducted at the Department of Agriculture of the Federal University of Lavras (UFLA) (lat 21º 14' S, long 45º 00' W and alt 918 m asl), Lavras, south of Minas Gerais.

Twelve genotypes of the genus *Anthurium* from the Embrapa Tropical Agroindustry (CPAT) germplasm bank, collected from different regions of Brazil, were studied: *Anthurium gracile* (IFT 107), *A. bomplandii* (IFT 108), *A. raimundii* (IFT 109), *A. jemanii* (IFT 117), *A. plowmanii* (IFT 118), *Anthurium sp.* (IFT 123), *A. lindmaniarum × A. guianesis* (IFT 128), *A. affine* (IFT 129), *Anthurium sp.* (IFT 255), *Anthurium sp.* (IFT 258), *A. andraeanum* (Astral IAC) and *A. andraeanum* (Rubi IAC).

The genotypes, previously established *in vitro* in the Tissue Culture Laboratory of Embrapa (CPAT), were sent to the UFLA. They were acclimatized in pots and maintained in an acclimatization room at a temperature of ± 28 °C for a photoperiod of 12 hours. After 120 days, three fully expanded leaves from three plants were collected for analysis.

**Flow cytometry**

For the three independent experiments, 30 mg of leaves from each genotypes were used, combined with the same amount of pea leaves (*Pisum sativum*) (DNA content reference standard 9.09 pg) or bean leaves (*Phaseolus vulgaris* L.) (DNA content reference standard 26.90 pg). Leaves were mashed in a petri dish containing 1 mL of nuclei extraction buffer, Tris or Marie (Dolezel et al. 1989), depending on the genotypes. Readings of 5000 nuclei isolated using 1 mL of Tris or Marie buffer and stained with 25 µL propidium iodide were performed. Histograms were acquired from a FacsCalibur® cytometer (Becton Dickinson) using the Cell Quest program (Dickinson 1998). DNA content (pg) of the plants was calculated using the following equation: DNA content (pg) = position of the G1 peak of the sample/position of the standard G1 peak × pg. Statistical analysis was performed using the software WinMDI 2.8 (Trotter 2000).

**Anatomical features**

Leaves were fixed in F.A.A 70% to perform the anatomical analysis (Kraus and Arduim 1997). Paradermic sections of the abaxial and adaxial faces were taken manually and cross-sections were made using a desk microtome (LPC model). For mounting the semi-permanent slides, sections were clarified with sodium hypochlorite 50% (v v⁻¹) and stained with 1.0% safranin and 1.0% safrablau (0.1% safranin and 1.0% Astra blue at a ratio of 7:3), for the paradermic and cross-sections, respectively, following the methodology proposed by Kraus and Arduim (1997). The material was observed under a Olympus CX41 microscope coupled to a Belcam DIV-3000 digital camera and photographed (15 fields), and the measurements were taken using the software ImageTool (UTHSCSA Image Tool for Windows version 3.0, San Antonio, TX, USA).

**Ultrastructural features**

For the ultrastructural analysis, leaves were immersed in fixative Karnovsky (1965) (pH 7.2) for 24 hours under refrigeration, washed in 0.056 M cacodylate buffer (three times, 10 minutes each time) and post-fixed in 1% osmium
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tetroxide in 0.05 M cacodylate buffer for 4 hours at room temperature, following the protocol proposed by Alves (2006). Specimens were observed using a scanning electron microscope (LEO Evo 040), operating between 10 and 20 kV. The entire adaxial and abaxial epidermis was observed for each leaf.

**Experimental design and statistical analysis**

Experiments were conducted using a completely randomized design with 12 treatments (genotypes) and three repetitions (three leaves per genotype). All the data were subjected to Anova using the software R (R Development Core Team 2011) and the means were compared using the Scott-Knott test at 5% probability. Samples were compared based on similarity of anatomical features and DNA content through cluster analysis. After establishing 12 features for the leaf (Density of the stomata (DS), polar diameter (PD) and equatorial diameter (ED) of the stomata and PD/ED ratio, adaxial epidermis (ADE), abaxial epidermis (ABE) and mesophyll (ME) thickness of leaf tissues, presence and absence of drusen, waxes and stomata, DNA content and coefficient of variation (CV) of histograms), cluster analysis was performed using the Euclidean distance and the most suited linkage method (centroid, complete, medium, simple, and ward) was verified and selected for the comparison using the cophenetic correlation coefficient. The software R (R Development Core Team 2010) was used for data analysis.

**RESULTS AND DISCUSSION**

Regarding genomic amplitude, a great amplitude was observed in the genus *Anthurium*, ranging from 4.95 to 22.74 pg (Table 1). There was a clustering of the genotypes, which were divided into seven groups. The first group was only composed of the IFT 128 (hybrid *A. lindmanianum* × *A. guianensis*) with 4.95 pg. The second group was composed of the genotypes IFT 123 (*Anthurium* sp.) and IFT 255 (*Anthurium* sp.), materials not yet identified, with approximately twice the amount of DNA (8.60 pg) when compared with the first group. The third group has the commercial cultivars (Rubi and Astral) belonging to the genotype *A. andraeanum*, with 9.18 and 9.28 pg, respectively. The fourth group had the genotypes IFT 109, IFT 118, IFT 117, and IFT 258, classified as *A. raimundii* (10.21 pg), *A. plowmanii* (10.22 pg), *A. jemanii* (10.64 pg), and *Anthurium* sp. (10.93 pg). The last three groups, with the highest amounts of DNA, were composed of the following isolated genotypes: *A. affine* (IFT 129) with 13.08 pg, *A. gracile* (IFT 107) with 21.55 pg and *A. bomplandii* (IFT 108) with 22.74 pg.

In the experiment, the DNA content of the plants was determined by flow cytometry, which statistically clustered the genotypes *A. raimundii*, *A. jemanii*, *A. plowmanii* and *Anthurium* sp. (IFT 258), and also the group of two genotypes not yet identified: IFT 123 and 255. Furthermore, it confirmed that the cultivars Astral and Rubi (IAC) belong to the same genotypes, which was also observed by Bliss and Suzuki (2012), and isolated *A. bomplandii*, *A. gracile* and *A. affine* from the other genotypes. From a practical point of view, determination of the nuclear DNA content can replace chromosome counting, especially when working with a large number of individuals, as in a germplasm bank (Schifino-Wittmann 2001).

In this study, the mean of the coefficient of variation (CV) for the leaf samples ranged between 0.34 and 0.73. These values demonstrate the quality of the results obtained and the reliability of the estimates of DNA content in the genotypes of *Anthurium* spp. (Table 1). The pea (*Pisum sativum*) and the bean (*Phaseolus vulgaris* L.) were used as internal standards in this study and have a 2 C DNA content of 9.09 and 26.09 pg, respectively (Table 1). These standards were selected because their DNA content lies in the middle of the mean value for most plants, and therefore can be used to assess plants with both small and large genomes. Unlike conventional methods, such as chromosome counting, especially when working with a large number of individuals, as in a germplasm bank (Schifino-Wittmann 2001).

The species of *Anthurium* have a leaf blade with smooth cuticle, polygonal or irregular epidermal cells, with straight or sinuous anticlinal walls, and stomata on both faces or only on the abaxial face (Figure 1). All *Anthurium* genotypes studied in this work showed leaf epidermal cells with straight or wavy walls, corroborating the results obtained by Keating (2002) and Mantovani et al. (2010). The epidermal cells in the leaves are short and have a random distribution. Mantovani et al. (2010) also observed this in a study with nine genotypes of *Anthurium*. There was, however, an exception as *A. bomplandii* showed more elongated cells, an important feature to differentiate it from other genotypes.

The leaf epidermis of the species analyzed had cells with periclinal walls on both faces, ranging from straight to convex, with exception of the hybrid that had sinuous anticlinal and convex periclinal walls, which is a distinctive feature
useful for its identification. In cross-sections, Keating (2002) observed that some genera of Araceae have the outline of the periclinal walls of epidermal cells, as well as the height/width proportion, and considered that these can be used for a taxonomic diagnosis.

Paradermic sections of the abaxial face of leaves are shown in Figure 1, demonstrating that the species have stomata on the abaxial face and are thus classified as hypostomatic leaves. The genotypes A. jemanii, A. affine, A. gracile, A. plowmanii and the hybrid are an exception, having stomata on both faces and at a lower density, and are classified as amphihypostomatic (Figure 1). These genotypes also show stomata in the central nerve or in its proximity. This is in agreement with observations by Keating (2002) (Table 2) showing that the leaf of most genotypes of the Araceae family is hypostomatic and the stomata are mainly concentrated on the abaxial surface.

**Table 1.** DNA quantification for genotypes of *Anthurium* spp. using flow cytometry

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Scientific Name</th>
<th>Buffer</th>
<th>Standard</th>
<th>DNA (pg)*</th>
<th>CV (%)</th>
<th>Histograms</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFT 128</td>
<td>Hybrid</td>
<td>Tris</td>
<td>P. vulgaris</td>
<td>4.95 g**</td>
<td>0.58</td>
<td></td>
</tr>
<tr>
<td>IFT 255</td>
<td><em>Anthurium</em> sp.</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>8.60 f</td>
<td>0.48</td>
<td></td>
</tr>
<tr>
<td>IFT 123</td>
<td><em>Anthurium</em> sp.</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>8.60 f</td>
<td>0.46</td>
<td></td>
</tr>
<tr>
<td>Rubi</td>
<td>A. andraeanum</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>9.18 e</td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td>Astral</td>
<td>A. andraeanum</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>9.28 e</td>
<td>0.40</td>
<td></td>
</tr>
<tr>
<td>IFT 109</td>
<td>A. raimundii</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>10.21 d</td>
<td>0.42</td>
<td></td>
</tr>
<tr>
<td>IFT 118</td>
<td>A. plowmanii</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>10.22 d</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>IFT 117</td>
<td>A. jemanii</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>10.64 d</td>
<td>0.73</td>
<td></td>
</tr>
<tr>
<td>IFT 258</td>
<td><em>Anthurium</em> sp.</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>10.93 d</td>
<td>0.72</td>
<td></td>
</tr>
<tr>
<td>IFT 129</td>
<td>A. affine</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>13.08 c</td>
<td>0.48</td>
<td></td>
</tr>
<tr>
<td>IFT 107</td>
<td>A. gracile</td>
<td>Marie</td>
<td>P. vulgaris</td>
<td>21.55 b</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>IFT 108</td>
<td>A. bomplandii</td>
<td>Tris</td>
<td>P. sativum</td>
<td>22.74 a</td>
<td>0.64</td>
<td></td>
</tr>
</tbody>
</table>

*Means followed by the same lowercase letter within the same column belong to the same group and do not differ significantly according to the Scott-Knott test at 5%.
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*Figure 1.* Photomicrographs of paradermic sections of the abaxial face of leaves from genotypes of *Anthurium* spp. a) IFT 108, b) IFT 117, c) IFT 128, d) IFT 123, e) IFT 129, f) IFT 255, g) IFT 107, h) IFT 258, i) IFT 109, j) IFT 118, l) IAC Astral and m) IAC Rubi. Bar 1 mm.

*Table 2.* Anatomical features of leaves of genotypes of *Anthurium* spp. Density of the stomata (DS), polar diameter (PD) and equatorial diameter (ED) of the stomata and PD/ED ratio, adaxial epidermis (ADE), abaxial epidermis (ABE) and mesophyll (ME) thickness of leaf tissues.

<table>
<thead>
<tr>
<th>Scientific Name</th>
<th>Collection Point</th>
<th>DS (mm²)</th>
<th>PD</th>
<th>ED</th>
<th>PD/ED</th>
<th>ADE (µm)</th>
<th>ABE (µm)</th>
<th>ME (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>A. raimundii</em></td>
<td>Bahia</td>
<td>17.07a</td>
<td>39.96d</td>
<td>33.41a</td>
<td>1.19c</td>
<td>25.5f</td>
<td>21.2d</td>
<td>216.0e</td>
</tr>
<tr>
<td><em>A. jemanii</em></td>
<td>Pará</td>
<td>15.27a</td>
<td>38.15d</td>
<td>33.48a</td>
<td>1.14c</td>
<td>45.7e</td>
<td>50.3c</td>
<td>508.5c</td>
</tr>
<tr>
<td><em>A. sp. (IFT258)</em></td>
<td>Espírito Santo</td>
<td>13.10b</td>
<td>41.78d</td>
<td>30.86b</td>
<td>1.36b</td>
<td>19.2f</td>
<td>16.0d</td>
<td>182.3f</td>
</tr>
<tr>
<td><em>A. plowmanii</em></td>
<td>Pará</td>
<td>12.71b</td>
<td>43.55b</td>
<td>34.20a</td>
<td>1.27c</td>
<td>43.3e</td>
<td>46.0c</td>
<td>523.2c</td>
</tr>
<tr>
<td><em>A. affine</em></td>
<td>Pernambuco</td>
<td>11.60b</td>
<td>38.30d</td>
<td>28.11c</td>
<td>1.36b</td>
<td>109.5c</td>
<td>112.0b</td>
<td>677.6b</td>
</tr>
<tr>
<td>Hybrid</td>
<td>Pará</td>
<td>10.80c</td>
<td>40.14c</td>
<td>29.92c</td>
<td>1.34b</td>
<td>118.6b</td>
<td>117.2a</td>
<td>484.5c</td>
</tr>
<tr>
<td><em>A. sp. (IFT123)</em></td>
<td>Pará</td>
<td>10.00c</td>
<td>33.53e</td>
<td>26.00c</td>
<td>1.28c</td>
<td>115.0c</td>
<td>128.8a</td>
<td>716.6b</td>
</tr>
<tr>
<td><em>A. sp. (IFT255)</em></td>
<td>Rio de Janeiro</td>
<td>8.64d</td>
<td>38.45d</td>
<td>27.43c</td>
<td>1.40b</td>
<td>127.0a</td>
<td>100.0b</td>
<td>916.2a</td>
</tr>
<tr>
<td><em>A. sp. (IFT255)</em></td>
<td>Rio de Janeiro</td>
<td>8.64d</td>
<td>38.45d</td>
<td>27.43c</td>
<td>1.40b</td>
<td>127.0a</td>
<td>100.0b</td>
<td>916.2a</td>
</tr>
<tr>
<td><em>A. andraeanum Astral</em></td>
<td>Cultivated</td>
<td>7.71d</td>
<td>38.17d</td>
<td>27.61c</td>
<td>1.38b</td>
<td>37.0e</td>
<td>36.3c</td>
<td>229.2e</td>
</tr>
<tr>
<td><em>A. andraeanum Rubi</em></td>
<td>Cultivated</td>
<td>7.50d</td>
<td>34.35e</td>
<td>27.76c</td>
<td>1.23c</td>
<td>57.1d</td>
<td>48.6c</td>
<td>303.9d</td>
</tr>
<tr>
<td><em>A. bomplandii</em></td>
<td>Amazonas</td>
<td>7.47d**</td>
<td>43.84b</td>
<td>33.85a</td>
<td>1.29c</td>
<td>118.3b</td>
<td>105.0b</td>
<td>685.1b</td>
</tr>
<tr>
<td><em>A. gracile</em></td>
<td>Paraíba</td>
<td>5.71e</td>
<td>54.58a</td>
<td>34.59a</td>
<td>1.57a</td>
<td>45.1e</td>
<td>40.4c</td>
<td>359.0d</td>
</tr>
</tbody>
</table>

CV % 22.22 9.61 10.68 13.41 16.25 24.47 14.43

Means followed by the same lowercase letter for a given column do not differ significantly from each other by the Scott-Knott test at 5%.
All stomata in this study were identified as paracytic brachyfacial, which are characteristic of genotypes of *Anthurium* (Mantovani and Pereira 2005, Mantovani et al. 2010). In the stomata of the abaxial face of *A. bomplandii*, the guard cells are less prominent than the subsidiary cells, which are more elongated, distinguishing this species from the others that have guard and subsidiary cells at the same level.

All species analyzed in this study have stomata flanked on each side by two subsidiary cells and guard cells with chloroplasts, and are classified as brachyparacytic. Most genotypes have epidermal cells with straight or sinuous walls, with the exception of *A. andraeanum* (Astral and Rubi) that has wavy walls, only on the abaxial face (Figure 1L and M).

Raphides were not found in the species studied although they are frequently found in the Araceae family (Keating 2004). On the other hand, drusen were found in all the species studied, in particular in the genotypes *Anthurium bomplandii*, *A. jemanii*, *A. gracile* and IFT 123, that showed the highest amounts of drusen concentrated in the adaxial face of the leaf blade and main nerve (Figure 2), which can be a distinctive feature for these genotypes.

The presence of small crystals in higher plants is common and is related to physical protection, removal of oxalate from the metabolic system, calcium storage, and regulation of light during photosynthesis in plants that grow in the shade (Franceschi and Nakata 2005). However, the presence, distribution or type of crystal (prismatic, drusen or styloid) can be used as a taxonomic feature. Moraes et al. (2011) suggested that the presence of styloid crystals could be an important feature to identify the genus *Psychotria*. In the present work, drusen crystals were found in all species studied (Figure 2).

According to Keating (2004), the presence of idioblasts with calcium oxalate crystals is a feature of the Araceae family. However, in *Anthurium*, the cells have independent features and the crystals can occur in the leaf blade, petiole, stem, and root while raphides are found in all organs. For the genotypes studied in this work, drusen were found only in the main nerve, leaf blade and below the epidermis, which was also observed by Mantovani et al. (2010).

The main nerve can have similar prominence in both faces for the genotypes *A. bomplandii*, *A. jemanii*, *A. lindmaniarum × A. guianensis* hybrid, *Anthurium* sp. (IFT 123 and 258), *A. raimundii*, *A. plowmanii*, *A. andraeanum* ‘Astral’ and *A. andraeanum* ‘Rubi’ (Figure 2A, B and C) or be more prominent in the abaxial face (*A. gracile*, Figure 2G) or the adaxial face (*A. affine* and *Anthurium* sp. (IFT 255), Figure 2E and F). The genotypes showing the highest prominence of the main nerve was *A. affine* (Figure 2E), while *A. bomplandii*, *Anthurium* sp. (IFT 123) and *A. andraeanum* Astral showed the lowest prominence (Figure 2A, D and L, respectively). These results are in agreement with observations by Keating (2003) that the shape of the main nerve is variable. It can be symmetrical, rounded or with a deltoid shape on both

![Figure 2. Photomicrographs of cross-sections of leaves from genotypes of Anthurium. a) IFT 108, b) IFT 117, c) IFT 128, d) IFT 123, e) IFT 129, f) IFT 255, g) IFT 107, h) IFT 258, i) IFT 109, j) IFT 118, l) IAC Astral and m) IAC Rubi. Bar 5 mm.](image)
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surfaces, flat or largely concave adaxially, and convex to rounded abaxially.

The mesophyll was uniform, with no significant difference between the palisade and spongy parenchyma. The mesophyll cells showed a compact arrangement in *A. lindmaniarum × A. guianesis*, *Anthurium* sp. (IFT 255), *A. plowmanii*, *A. andraeanum* Astral and Rubi, with the exception of *A. bomplandii*, *A. jemanii*, *Anthurium* sp. (IFT 123), *A. affine*, *A. gracile*, *Anthurium* sp. (IFT 258) and *A. raimundii*, in which intercellular spaces were found.

All genotypes showed epicuticular wax deposited on the surface of the cuticle with different shapes. Among others, wax can have the shape of scales (*Anthurium* spp. – IFT 258), a continuous cover (*A. raimundii*) or, as in most cases, granules (*A. jemanii*, hybrid and all others species studied) (Figure 3). The shape of the wax deposited on the surface of the organs is particular to each genotype. The wax also acts as a barrier against fungi, bacteria and insects (Appezzato-da-Glória and Carmello-Guerreiro 2012).

The genotypes analyzed in this study show a smooth cuticle on both faces. Similar results were reported by Mantovani et al. (2010), who observed that the unornamented cuticle is a frequent feature in the Araceae family.

Regarding the density of the stomata, the genotypes that showed a highest number of stomata were *A. jemanii* and *A. raimundii*, while *A. gracile* showed the lowest density of stomata. For the polar and equatorial diameters, *A. gracile* showed higher values for both diameters and also for the ratio between them. The values found for the density of stomata were similar between groups, ranging between 5.71 and 17.07 stomata mm$^2$ (Table 2). In this study, *A. raimundii* and *A. jemanii* had larger stomata while in *A. gracile*, the stomata were smaller. Larger stomata are usually associated with mesophytic environments, allowing a higher consumption of water (Galmes et al. 2007), and smaller stomata would be found in xeric environments, providing a better water control (Aasamaa et al. 2001).

In the adaxial epidermis, *Anthurium* sp. (IFT 255) showed greater epidermal thickness than the other genotypes and also showed a greater thickness of the mesophyll (916.2 µm). However, in the abaxial epidermis, *A. lindmaniarum × A. guianesis* hybrid and *Anthurium* sp. (IFT 123) showed the greatest thickness. *Anthurium raimundii* and IFT 258 had similar values for both faces (adaxial and abaxial), showing the lowest values for the epidermal thickness. The *Anthurium* sp. (IFT 258) showed the smallest thickness of the mesophyll with a value of 182.3 µm (Table 2).

Figure 3. Scanning electron micrographs of the epidermis of the adaxial face of leaves of genotypes of *Anthurium* spp., showing the different types of wax. a) and b) Epicuticular wax in the shape of granules in *A. jemanii* and *A. lindmaniarum × A. guianesis*. Bar 20 µm. c) Epicuticular wax in the shape of scales in IFT 258. d) Epicuticular wax as continuous cover in *A. raimundii*. Bar 30 µm.
According to the dendrogram obtained following cluster analysis (Figure 4), the individuals can be divided into three groups considering the mean Euclidean distance (450), with several subgroups.

Cluster analysis grouped the genotypes A. raimundii and A. affine, A. bomplandii and A. gracile, and A. jemanii and A. plowmanii, and isolated the hybrid, one of the few genotypes that does not have a large amount of wax in the epidermis. The IFT 258, a genotype not yet identified, was clustered with the branch containing the genotypes A. raimundii and A. affine. The other two unidentified genotypes, IFT 123 and 255, were clustered with A. bomplandii and A. gracile. This data confirms that the unidentified genotypes belong to the same section of these.

We also observed that A. andraeanum, represented by the cultivars Astral and Rubi, is closely related to the rest of species examined in the current study. Such results reinforce the potential of anatomical and cytometric analysis for clustering and differentiating genotypes.

Anatomical data and DNA content were helpful for species identification and grouping through cluster analysis. Using cluster analysis of anatomical features of the leaf and spathe, Mantovani et al. (2010) reinforced the importance of clustering in the diagnosis of genotypes belonging to Anthurium.

Thus, knowledge of the genetic makeup of access of several species is important to enrich the knowledge of researchers and producers from around the world (Hornung-Leoni et al. 2013).

CONCLUSIONS

Anatomical studies clustered the genotypes A. andraeanum (Astral and Rubi) and distinguished the hybrid (A. lindmanianum × A. guianesis) and A. bomplandii from the other genotypes, with anatomical features specific to each.

Differences in DNA content, assessed by flow cytometry, allowed the differentiation between the genotypes A. andraeanum, A. affine, A. bomplandii, and A. gracile and the hybrid (A. lindmanianum × A. guianesis) and clustered the group A. raimundii, plowmanii and jemanii of Anthurium sp. (IFT 258), in all analyses.
Grouping of *anthurium* genotypes based on genetic and morpho-anatomical features

Cluster analysis allowed grouping of the genotypes *A. raimundii* and *A. affine*, *A. bomplandii* and *A. gracile*, *A. jemanii* and *A. plowmanii*, and the cultivars Astral and Rubi (*A. andraeanum*), and isolated the hybrid (*A. lindmaniarum × A. guianensis*).

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**REFERENCES**


