Genetic diversity of root anatomy in wild and cultivated *Manihot* species

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ABSTRACT. An anatomical study of roots was conducted on two wild *Manihot* species, namely *M. glaziovii* and *M. fortalezensis*, and two cassava varieties, *M. esculenta* Crantz variety UnB 201 and *M. esculenta* variety UnB 122, to identify taxonomic differences in primary growth. Anatomical characters of cassava roots have been rarely investigated. Their study may help cassava breeders to identify varieties with economically important characters, such as tolerance to drought. We investigated tap and lateral adventitious roots of two specimens of each clone or species. Free-hand cross-sections of roots were drawn; these had been clarified with 20% sodium hypochlorite solution, stained with 1% safranin-alcian blue ethanolic solution, dehydrated in ethanol series and butyl acetate and mounted in synthetic resin. Anatomical differences among *Manihot* species and varieties were found in the epidermal and exodermal cell shape and wall thickness, content of cortical parenchyma, and number of xylem poles. Wall thickness of the epidermis and exodermis of tap root were similar in all species, while in the lateral root there were differences in cell shape and wall thickness. Epidermal cells with thick walls were found in the tap root of all species and in lateral
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roots of cassava varieties. This character is apparently associated
with tolerance to drought and disease. The variation in the number
of xylem poles of cassava varieties was larger (4-8) than in wild
species (4-6), and appears to support the hybrid origin of cassava.

**Key word:** Primary growth; *Manihot glaziovii*; *Manihot esculenta*
*Manihot fortalezensis*; Xylem poles

**INTRODUCTION**

Wild *Manihot* species are important for improving cassava, *Manihot esculenta* Crantz. The root is of interest to research on this biological group, because it is a consumed part of the cultivated plant. Acquiring information on this part may help improve the crop for different affinities. There are scarce reports in the literature on the subject. Our study is probably the first one on the subject.

Economic characters such as tolerance to drought and insects are found in wild *Manihot* species (Nichols, 1947; Nassar, 1986). Knowing more about root anatomy would help in understanding the mechanism and nature of these characters.

Hybridization between *Manihot* species occurs naturally (Nassar, 2001; Chacón et al., 2008) and could lead to variation and diversity in different characters, either of the stem, flower or fruit. This presumed diversity may have resulted as a result of domestication action on all material. Also, knowledge of the anatomy of the cultivated plant compared to the wild species may contribute to information on the domestication.

In this article, we present the findings on the anatomy of 2 wild *Manihot* species and 2 varieties of cassava and their economic significance.

**MATERIAL AND METHODS**

Two wild *Manihot* species, namely *Manihot glaziovii* Muller and *M. fortalezensis* Nassar et al., and two varieties, *M. esculenta* Crantz variety UnB 201 and *M. esculenta* variety UnB 122, were our study material.

*Manihot glaziovii* and *M. fortalezensis* were collected from Ceará State, northeast region of Brazil, and grown at the experimental station of Universidade de Brasília. *M. esculenta* variety UnB 201 was collected from Amapá State, northern region of Brazil, while the variety UnB 122 was selected from the progeny of the *M. esculenta* and *M. anomala* Pohl hybrid. All specimens were kept with the live collection at the Universidade de Brasília, Brasília, Brazil.

Adventitious roots in primary growth were collected at 1-3 months after planting. The roots collected were fixed in 70% FAA (Johansen, 1940) for a period of 24 h and preserved in 70% ethanol. Free-hand cross-sections were made of primary roots 10 cm in length, using a microtome, and clarified in 20% sodium hypochlorite (Kraus and Arduin, 1997). Next, the sections were stained with 1% safranin and 1% aqueous alcian blue (Luque et al., 1996), dehydrated in an ethanol series, cleared in butyl acetate, and mounted in synthetic resin (Paiva et al., 2006). All sections were made from the tap root and from lateral ones.
RESULTS AND DISCUSSION

The four specimens studied showed anatomical differences in the epidermis, exodermis, cortical parenchyma, and number of vascular poles. The differences are described in Table 1.

<table>
<thead>
<tr>
<th>Species</th>
<th>Tap root Epidermis</th>
<th>Lateral root Epidermis</th>
<th>Exodermis</th>
<th>Cortical parenchyma (at 7 mm to root tip)</th>
<th>Number of vascular poles</th>
</tr>
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<tbody>
<tr>
<td><em>M. fortalezensis</em></td>
<td>Circular outer cells with wall thickened</td>
<td>Circular to narrow cells with walls slightly thickened</td>
<td>Circular to polygonal (4-7 sides). Cells longer than in varieties UnB 201 and UnB 122.</td>
<td>Up to 11 layers. Abundant druses</td>
<td>5, 6, 4-6</td>
</tr>
<tr>
<td><em>M. glaziovii</em></td>
<td>Circular to tabular cell with wall thickened</td>
<td>Circular to polygonal (4-7 sides). Cells longer than in varieties <em>M. esculenta</em> and <em>M. fortalezensis</em></td>
<td>Up to 10 layers. Rare druses</td>
<td>4, 5</td>
<td></td>
</tr>
<tr>
<td><em>M. esculenta</em> Crantz (variety UnB 201)</td>
<td>Circular cells with thickened wall</td>
<td>Circular to polygonal (4-7 sides). Cells longer than in varieties <em>M. glaziovii</em> and <em>M. fortalezensis</em></td>
<td>Up to 14 layers. Rare druses</td>
<td>6, 7, 5</td>
<td></td>
</tr>
<tr>
<td><em>M. esculenta</em> (variety UnB 122)</td>
<td>Circular to irregular cells with thickened</td>
<td>Up to 11 layers. Frequent druses</td>
<td>5, 6, 8, 4, 5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The tap root in primary growth showed a cuticle and unistratified epidermis, which was formed by common epidermal cells with outer circular shape and thick wall (Figure 1A). These cells showed little variation in size. In *M. esculenta* variety UnB 122, the cells have an acute end and irregular shape (Figure 1B). Trichomes were short and broad, with rounded tips, and they were observed in both varieties (Figure 1C).

In all species studied, the exodermis is a uniseriate layer, which differs in shape. Its cells vary from circular to polygonal (4-7 sides) in shape, may be homogeneous and have outer periclinal walls, thick and lignified (Figure 1D), with 2-3 times the length of epidermal cells. Cultivated species have smaller-sized cells, with twice the length of epidermal cells.

The cortex was found to be composed of isodiametric cells of varying size and thin wall. Cells in the middle of the cortex are twice the size of other cortical cells. The parenchyma differed between species as to the content and number of layers, as described in Table 1. Calcium oxalate druses were also noted in the cortex (Figure 1E).

Endoderm was found to be formed by small, circular to square cells of uniform size, and the Casparian bands were present in the anticlinal walls (Figure 1F). In the variety UnB 122, it was also possible to observe Casparian bands in the periclinal walls (Figure 2A). The central cylinder was composed of uniseriate pericycle with circular to square, small cells involving other tissues. The primary phloem in all species studied consisted of 1-4 layers of polygonal to round cells with walls slightly thicker than the walls of the parenchyma (Figure 1E).

Primary xylem vessel elements with lignified walls and vessel gaps were observed. The number of vascular poles differed between species. In *M. glaziovii*, 4 and 5 poles were observed (Figure 2B), while 5 and 6 poles in *M. fortalezensis* (Figure 1F). In the variety UnB 201, 6 (Figure 2C) and sometimes 7 (Figure 2D) poles were observed. The variety UnB 122 showed 4, 5, 6 (Figure 2E) and sometimes 8 (Figure 2F) poles. Lateral roots were seen in front of vascular poles (Figure 3A). Surrounding xylem vessels were parenchyma cells of variable size and shape. The larger cells were present at the center to form pith, and cells varied in shape from circular to polygonal.
Figure 1. A. Epidermal cells with thick walls in *Manihot esculenta* variety UnB 201. B. Epidermal cells with acute end and irregular shape in *M. esculenta* variety UnB 122. C. Trichomes short and broad, with rounded tip in *M. esculenta* variety UnB 122. D. Exodermal cells with thick and lignified outer periclinal wall in *M. fortalezensis*. E. Cortical parenchyma with calcium oxalate druses (arrows) in *M. fortalezensis*. F. Casparian bands on anticlinal walls of exodermal cells in *M. fortalezensis* with 7 mm to root tip (tap root). Bar: 100 µm.
Figure 2. A. Casparian bands on anticlinal and periclinal walls of endodermal cells in *Manihot esculenta* variety UnB 122 (lateral root). B. Cross-section of *M. glaziovii* with 5 vascular poles (tap root). C. Six vascular poles in *M. esculenta* variety UnB 201 (lateral root). D. Seven vascular poles and cortical parenchyma *M. esculenta* variety UnB 201 (tap root). E. Six vascular poles in *M. esculenta* variety UnB 122, 7 mm to root tip (tap root). F. Eight vascular poles and Casparian bands on anticlinal walls of endodermis in *M. esculenta* variety UnB 122 (tap root). Bar: 100 µm.
Figure 3. A. Lateral root emission in *Manihot glaziovii*. B. Tabular epidermal cells with thick wall in *M. glaziovii* (lateral root). C. Narrow epidermal cells with thick wall in *M. fortalezensis* (lateral root). D. Four vascular poles in lateral root of *M. glaziovii* in 7 mm to root tip. E. Four vascular poles in lateral root of *M. glaziovii*, 7 mm to root tip. Bar: 100 µm.
The lateral roots showed a structure similar to that of the tap root, except for the shape of common epidermal cells in *M. glaziovii* and *M. fortalezensis*. Circular to tabular epidermal cells were observed in *M. glaziovii* (Figure 3B), while in *M. fortalezensis*, cell shape was circular to narrow (Figure 3C). The number of poles of the vascular system also varied between the lateral roots and tap roots. *M. glaziovii* had lateral roots with 4 poles and the center occupied by the vascular vessel (Figure 3D). *M. fortalezensis* exhibited 4 poles (Figure 3E), sometimes 5 and 6 poles in the lateral roots. The variety UnB 201 showed 5 poles, while in the variety UnB 122 there were 4 and 5 poles.

The thickness of epidermal and exodermal cell walls may have significance in relation to drought and tolerance to disease, because Casparian bands and suberin lamellae may restrict water loss (Peterson et al., 1993) and pathogen colonization (Schreiber et al., 1994). There are reports on the increase of cell wall thickness in plants under drought conditions (Cruz et al., 1992; Mostajeran and Rahimi-Eichi, 2008). All *Manihot* species studied here have thick cell walls of epidermis and the outer periclinal walls of the exodermis being thick. This finding may help understand the rustic characteristic in the *Manihot* species reported by Nassar et al. (2008, 2010), Graciano-Ribeiro (2008), and Graciano-Ribeiro et al. (2009).

Within the Euphorbiaceae, some genera such as *Euphorbia* have their roots with 3 or 4 poles (Gales and Toma, 2006), while *Jatropha cordata*, a species belonging to the same subfamily of *Manihot*, has 2 vascular poles (Popham, 1947). *Manihot pilosa* and *M. esculenta* have 4 or 5 poles (Indira and Kurian, 1977; Vanucci, 1982), and 6 poles (Moraes-Dalaqua and Coral, 2002), but this is the first report of more than 6 poles in *Manihot* species and the plants studied. Variation in the same species is frequently reported (Esau, 1965). Although *Manihot* species are dicots, they showed more than 4 poles and lateral root position in front of poles. These characteristics are commonly reported in monocotyledons (Rudall, 2007).

While *M. glaziovii* showed a very limited variation in pole number, cassava (varieties UnB 201 and UnB 122) exhibited variation ranging from 5 to 8 poles. The variation in pole number of cassava cultivars studied here may support the hypothesis of hybrid origin of the cultigen where segregation is likely to produce this notable variation (Nassar, 2002, 2003a,b).

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