Axillary bud and pericycle involved in the thickening process of the rhizophore nodes in *Smilax* species

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(With 25 figures)

Abstract

The species of the genus *Smilax*, popularly known as sarsaparilla, are widely used in folk medicine due to the antirheumatic properties of its underground structures. *Smilax fluminensis* and *S. syphilitica* occur in forested areas and form thickened stems called rhizophores from which adventitious roots grow. To provide information for more accurate identification of the commercialised product and for elucidating the process of stem thickening, a morphology and anatomy study of the underground organs of the two species was conducted. The adventitious roots differ in colour and diameter depending on the stage of development. They are white and have a larger diameter in the early stages of development, but as they grow, the adventitious roots become brown and have a smaller diameter due to the disintegration of the epidermis and virtually the entire cortex. In brown roots, the covering function is then performed by the lignified endodermis and the remaining walls of the cells from the last parenchyma cortical layer. These results are similar to those found in studies of other *Smilax* and suggest that the anatomy of the roots can be useful for identifying fraud in commercialised materials. The thickening process of the nodal regions of the rhizophores in both species involves the activity of axillary buds and pericyclic layers.

Keywords: sarsaparilla, *Smilax*, underground system, rhizophore, anatomy.

Gema axilar e periciclo envolvidos no processo de espessamento nodal do rizóforo em espécies de *Smilax*

Resumo

As espécies de *Smilax*, conhecidas popularmente como salsaparrilha, são amplamente utilizadas na medicina tradicional devido às propriedades antirreumáticas das estruturas subterrâneas. *Smilax fluminensis* e *S. syphilitica* ocorrem em áreas florestais e formam caules espessados denominados rizóforos a partir dos quais são emitidas raízes adventícias. Com o intuito de fornecer informações para a identificação mais precisa do material comercializado e no entendimento do processo de espessamento do caule, foi realizado o estudo morfológico e anatômico dos órgãos subterrâneos das duas espécies. As raízes adventícias apresentam diferenças na coloração e no diâmetro dependendo da fase de desenvolvimento. As raízes nas fases iniciais do desenvolvimento são brancas e possuem diâmetro maior, porém com o desenvolvimento, devido à desintegração da epiderme e de praticamente todo o córtex, as raízes tornam-se marrons e de diâmetro menor. Nas raízes marrons, a função de revestimento passa a ser exercida pela endoderme lignificada e pelas paredes remanescentes das células da penúltima camada cortical. Os resultados são semelhantes aos encontrados nos estudos de outras *Smilax* e sugerem que a anatomia das raízes pode ser útil na identificação de fraudes em materiais comercializados. O processo de espessamento das regiões nodais dos rizóforos nas duas espécies envolve a atividade das gemas axilares e de camadas periciclicas.


1. Introduction

*Smilax* comprises 310 species distributed in temperate and tropical regions (Judd et al., 2009). There are 32 species in Brazil (Andreata, 1997; Andreata, 2009), some of which are popularly known as sarsaparillas and have been used in folk medicine since the 16th century (Medeiros et al., 2007). For many years, sarsaparilla was identified by how their roots were tied in bundles for commercialization and by their external appearance (Farmacopéia, 1929). Studies of the Brazilian *Smilax* have demonstrated that their roots change in thickness and colour according
to the stage of development (Martins and Appezzato-da-Glória, 2006; Martins et al., 2010; Guimarães et al., 2010), i.e., these features are not useful for identifying the sarsaparilla. However, the anatomical features of the root in cross section, such as the relative thicknesses of cortical, vascular, and medullary regions; the presence of starch; the distribution of phenolic and crystalliferous idioblasts; and the phloem arrangement, can provide a larger number of elements for more accurate identification of the drug (Cunha, 1937; Stellfeld, 1938; Soares, 2013). The accurate identification of medicinal plants, including the detailed morphological and anatomical descriptions of its underground organs, is essential for quality control in phytotherapeutic studies (Ming, 1994).

*Smilax fluminensis* and *S. syphilitica* occur in forested areas and form thickened stems called rhizophores, following the classification proposed by Andreata and Menezes (1999) for *S. quinquenervia*. According to them, the rhizophores of this species, unlike the rhizomes, have two stem axes during the early stages of development. An aerial stem system of plumular origin and an underground stem system of cotyledonal bud origin have been described, and the root system is completely adventitious and formed by the underground stem axis (Andreata and Menezes, 1999). The presence of two stem axes was also observed in *Smilax polyantha*, but the underground stem axis originated from axillary buds of the cataphylls at the base of the aerial branching (Martins et al., 2011) and not from the cotyledonal bud as described in *S. rotundifolia* and *S. glauca* (Holm, 1890) and in *S. quinquenervia* (Andreata and Menezes, 1999).

The rhizophores of *Smilax* exhibit nodal thickening from which all adventitious roots of the plant are emitted (Martins et al., 2011). Moreover, the nodes of the stem branches, when stimulated by fragmentation, also thicken and form adventitious roots, thereby increasing the chances for vegetative propagation (Soares et al., 2011).

The primary thickening in monocotyledons has been discussed under a different approach by Menezes et al. (2005, 2012). According to them, in some species, such as *Zingiber officinale* (Zingiberaceae), the observation of Casparian bands in the endodermis allowed researchers to verify that the pericycle is involved in the tuberisation process of the rhizome. However, in *Smilax polyantha*, it was not possible to establish the boundary between the cortex and the vascular cylinder and the meristematic band responsible for tuberisation was named the primary thickening meristem (Martins and Appezzato-da-Glória, 2006).

Therefore, this study presents the morphology and anatomy of the underground organs of *Smilax fluminensis* and *S. syphilitica* to provide information for more accurate identification of the commercialised material and elucidating the rhizophore thickening process.

### 2. Material and Methods

Three individuals were analysed from each species. The *S. syphilitica* Humboldt and Bonpland ex Wilddenow individuals were collected in the Atlantic Forest, in Santa Tereza, Espírito Santo state, and *S. fluminensis* Steud. individuals were collected from the riparian forest area in the municipality of Itirapina, São Paulo state, Brazil. The collected material was identified by Dr. Regina Helena Potsch Andreata, an expert in the genus *Smilax* in Brazil, and the voucher specimens were added to the Herbarium of the School of Agriculture (ESA) under the numbers 107665 (*S. syphilitica*) and 107633 (*S. fluminensis*).

Rhizophores and adventitious roots of different diameters in the underground system were analysed. Samples were fixed in FAA 50 (1:1:8 formaldehyde, glacial acetic acid, and 50% ethanol) (Johansen, 1940). For better fixation, the samples were subjected to a vacuum to remove the air in the tissues. They were then dehydrated in an ethanol series and embedded in hydroxyethyl methacrylate (Leica™ Historesin); the obtained blocks were cut into 5-10 μm thick sections with a rotary microtome. Berberine-aniline blue staining was used for detecting suberin and lignin for identifying Casparian bands (Brundrett et al., 1988). For the remaining analyses, the material was stained with safranin and astra blue (Burger and Richter, 1991) or with 0.05% toluidine blue in phosphate buffer and citric acid at pH 4-6 (Sakai, 1973). The stained slides were mounted in Entellan synthetic resin.

The images were digitally captured with a Leica DMLB microscope (Leica™ - Wetzlar, Germany) using a video camera plugged into a computer utilising the IM50 software for image analysis. The berberine-aniline blue staining was visualised with ultraviolet light.

### 3. Results

Regarding the rhizophores (Figures 1-4), the two species have in common the rigidity, thickening in the nodal regions, and the formation of white (Figure 2) and brown adventitious roots (Figures 3-4). *Smilax fluminensis* individuals due to its elongated internodes (Figure 1) known as “runners”, is distributed in parallel to the soil surface and can reach between 7 and 12 m until a new aerial branch is emitted. The branches are emitted from thickenened nodes (Figures 1-3). There is no horizontal development in *S. syphilitica*, because the internodes are short and the thickened nodes and the aerial branches are too close (Figure 4). In both species, the underground system is located at a depth between 10 and 15 cm.

From the nodal area undergoing the tuberisation process (Figures 5-9), it was possible to observe that the swelling process of the rhizophore occurs through the meristematic activity of the axillary buds (Figures 5-7) and through divisions in the pericyclic cell layers underlying the endodermis, which can be recognised due to the parietal thickening in lipophilic and phenolic substances (Figures 8-9), near the bud.
It is notable that tuberisation is generally bilateral (Figure 7), and an adventitious root is formed at the end of each projection (Figure 10) whose pericyclic origin is evidenced by the continuity of the endodermis with U-shaped thickening (Figure 11).

In both studied species, the adult rhizophore is thickened and exhibits a uniseriate epidermis (Figure 12) with stomata. Phenolic or raphid idioblasts occur between several layers of the cortical parenchyma cells (Figure 12). Peripheral bundles (Figure 13) are commonly found, often surrounded by cells whose parietal thickness is similar to that of the endodermis (compare Figures 12 and 13). The vascular cylinder has a pericycle with several cell layers elongated in the anticlinal direction (Figure 14). There are smaller collateral bundles distributed among the cells of the proliferated pericycle, and the remaining collateral bundles are larger and randomly distributed, as illustrated in Figure 9. There is accumulation of starch grains throughout the vascular cylinder, but with higher concentration in the pericyclic cells (Figure 14) and surrounding the vascular bundles (Figure 15). The endodermis with parietal thickening delimits the cortex and vascular cylinder of the rhizophores (Figure 16).

In both species, the adventitious root system of the rhizophore consists of roots at different stages of development. There are white tender roots of a larger diameter and brown rigid roots of a variable diameter (Figures 2-4). The white roots represent the youngest stage, characterised by root hairs and a uniseriate epidermis (Figure 17). The cortex displays an exodermis with Casparian bands (Figures 18-19), followed by the cortical parenchyma consisting of isodiametric cells. The cells of the last parenchyma cortical layer exhibit thickening in the inner periclinal wall and in the radial walls (Figure 20). The endodermis has more elongated cells in the periclinal direction and walls with Casparian bands (Figure 20). The vascular cylinder is a polyarch, and the pericycle is formed by approximately four cell layers with walls that become gradually thicker.
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(Figures 22-24). Lysis occurs throughout the development and is accompanied by gradual removal of the epidermis and the entire cortical parenchyma (Figure 21). The root becomes brown and rigid at this stage, and the covering function is then performed by periclinal cell walls of the last cortical parenchyma layer and by the endodermis, which at this stage has thick and highly lignified walls (Figures 23-24). These walls have a brown colour due to the accumulation of phenolic compounds, which, in *S. syphilitica*, can be also present within the endodermal cells (Figure 24). The pith contains spherical and polyhedral starch grains (Figure 25).

**Figures 5-9.** Cross-sections of the nodal region of *Smilax fluminensis* Steud (5-7) and *S. syphilitica* Humboldt and Bonpland ex Willdenow (8-9). 5-7 Note the different stages of thickening with no change in the original vascular cylinder of the nodal region (Vc), which demonstrates the meristematic activity underlying the axillary bud. 7. Note the bilateral thickening (arrows). 8. Tuberisation initiates in the region below the axillary bud and divisions of the pericyclic cell layers underlying the endodermis (En) with thickened walls. Ab = axillary bud.

**Figures 10-11.** *Smilax fluminensis* Steud. 10. Cross-section of the end of a nodal tuberisation displaying the emission of an adventitious root. 11. Detail of the section indicated in the previous figure demonstrating the endoderm with thickened walls (arrow).
4. Discussion

The white and brown adventitious roots of the root system, as well as their ontogenetic processes, are extremely similar to those described for *S. polyantha* (Martins and Appezzato-da-Glória, 2006), *S. subsessifolia* (Guimarães et al., 2010), *S. brasiliensis*, *S. campestris*, and *S. cissoides* (Martins et al., 2010). The distribution of the phenolic and crystalliferous idioblasts, the circular arrangement of the primary phloem, and the absence of metaxylem elements in the centre of the structure are also common among other species already described in the...
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literature and have allowed identification of the adulteration of products being sold as sarsaparilla (Soares, 2013).

The elongated internodes in the underground system were only verified in *S. fluminensis*, similar to that described in *S. rufescens* (Martins et al., 2010).

The epidermis with stomata in rhizophores, most likely resulting from the derivation from aerial structures (Andreata and Menezes, 1999; Appezzato-da-Glória, 2003), is a pattern already recorded in other *Smilax* species (Andreata and Menezes, 1999; Palhares and Silveira, 2003).

Figsures 17-25. Cross-sections of the root of *Smilax fluminensis* Steud (17-23) and *S. syphilitica* Humboldt and Bonpland ex Wilddenow (24-25). 17. White root covered by the epidermis. 18. White root stained with berberine-aniline blue, observed under ultraviolet light. 19. Detail of the exodermis with Casparian bands. 20. Cells of the last cortical parenchyma layer with thick walls and tall endodermal cells. 21. Beginning of the lysigenous process (arrows) of the cortical cells. 22. Detail of the parietal thickening of the endodermis with a U-shaped thickening and pericycle. 23. Brown root covered by the cell wall of the last cortical parenchyma layer and endodermis. 24. Detail of the brown root lining with phenolic compounds evidenced by the ferric chloride on the wall and lumen (arrows) of the endodermal cells. 25. Spherical and polyhedral starch grains located in the medulla, observed under polarised light.
The data presented here suggest an investment in an anti-herbivore strategy, as observed in the parenchyma of the tuberous roots of *Asphodelus aestivus* (Xanthorrhoeaceae) (Sawidis et al., 2005). In addition, because the rhizosphere is an organ capable of expanding the rhizosphere and conferring resistance and vegetative propagation (Andreatta and Menezes, 1999), it is understandable that the rhizosphere accumulates large amounts of starch grains, such as the ones regularly found in the cortex and throughout the vascular cylinder of the structure in both studied species. 

In this study, unlike reports in other species of *Smilax* (Martins and Appezzato-da-Glória, 2006; Martins et al., 2010), it was possible to observe the endodermis with parietal thickening delimiting the cortex and vascular cylinder of the rhizophores. This analysis made it possible to determine that the cell proliferation leading to nodal thickening occurs in the proximity of the axillary bud in the layers underlying the endodermis, i.e., there is a collaboration from the pericycle in this process. Although it was not possible to establish the exact boundary between the cortex and vascular cylinder in *S. campestris*, *S. goyazana*, and *S. oblongifolia* (Martins et al., 2010), the cell proliferation is extremely similar to that observed in our study. Furthermore, in the analysed species and in *S. campestris*, *S. goyazana*, and *S. oblongifolia* (Martins et al., 2010) there is no establishment of a meristematic area similar to that described in *S. polyantha* by Martins and Appezzato-da-Glória (2006). The data presented here support the proposal by Menezes et al. (2012) that, in monocotyledons, the pericycle itself is involved in the thickening process with the formation of new vascular bundles and parenchyma. Moreover, the continuity of the endodermis of the rhizophores and adventitious roots, observed on their emission sites, corroborates the statements that the endodermis is a continuous tissue present in roots and stems (Menezes et al., 2005). The involvement of the axillary bud in the nodal thickening process and consequent rooting in the studied species had already been observed in stem cuttings of *S. fluminensis* (Soares et al., 2011). The authors found that adventitious root formation did not involve callus formation and neither occur in the basal region of the cuttings, but instead occurred in the basal region of the swollen bud.

Regarding the peripheral bundles of the rhizophores, only a few cells had parietal thickening similar to that of the endodermis, but in *Smilax goyazana*, the endodermis was described as surrounding all the vascular bundles (Palhares and Silveira, 2005).

In conclusion, the axillary bud and the pericycle participate in the process of rhizophore nodal thickening. The adventitious roots have common features among the *Smilax* species, which may facilitate the diagnosis of adulteration in drugs commercialised as sarsaparilla.

References


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